



(19) **United States**

(12) **Patent Application Publication**

**Shelton, IV et al.**

(10) **Pub. No.: US 2021/0315579 A1**

(43) **Pub. Date: Oct. 14, 2021**

(54) **METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS**

*A61B 5/026* (2006.01)  
*A61B 6/00* (2006.01)  
*A61B 17/068* (2006.01)  
*A61B 17/072* (2006.01)  
*A61B 17/11* (2006.01)  
*A61B 17/128* (2006.01)  
*A61B 17/32* (2006.01)

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(52) **U.S. Cl.**

CPC ..... *A61B 17/1155* (2013.01); *A61B 34/30* (2016.02); *A61M 1/79* (2021.05); *A61B 34/20* (2016.02); *A61B 34/32* (2016.02); *A61B 34/71* (2016.02); *A61B 90/35* (2016.02); *A61B 90/361* (2016.02); *G16H 40/67* (2018.01); *G16H 10/60* (2018.01); *G16H 50/20* (2018.01); *G16H 40/63* (2018.01); *G16H 70/20* (2018.01); *A61B 1/00009* (2013.01); *A61B 1/00045* (2013.01); *A61B 1/051* (2013.01); *A61B 1/0661* (2013.01); *A61B 5/0066* (2013.01); *A61B 5/0075* (2013.01); *A61B 5/0261* (2013.01); *A61B 6/5247* (2013.01); *A61B 17/0682* (2013.01); *A61B 17/072* (2013.01); *A61B 17/1114* (2013.01); *A61B 17/1285* (2013.01); *A61B 17/320092* (2013.01); *A61B 18/1442* (2013.01); *A61B 18/1445* (2013.01); *B25J 9/1697* (2013.01); *B25J 13/006* (2013.01); *G06K 7/10316* (2013.01); *G06K 19/07749* (2013.01); *H01Q 1/22* (2013.01); *H04L 63/1416* (2013.01); *H04L 67/10* (2013.01); *H04L 67/12* (2013.01); *H04N 5/272* (2013.01); *H04N 7/183* (2013.01); *H05K 1/028* (2013.01); *H05K 1/189* (2013.01); *A61M 1/80* (2021.05); *A61B 2034/2055* (2016.02); *A61B 2034/2057* (2016.02); *A61M 1/73* (2021.05)

(Continued)

(21) Appl. No.: **17/358,239**

(22) Filed: **Jun. 25, 2021**

**Related U.S. Application Data**

(63) Continuation of application No. 16/209,416, filed on Dec. 4, 2018.

(Continued)

**Publication Classification**

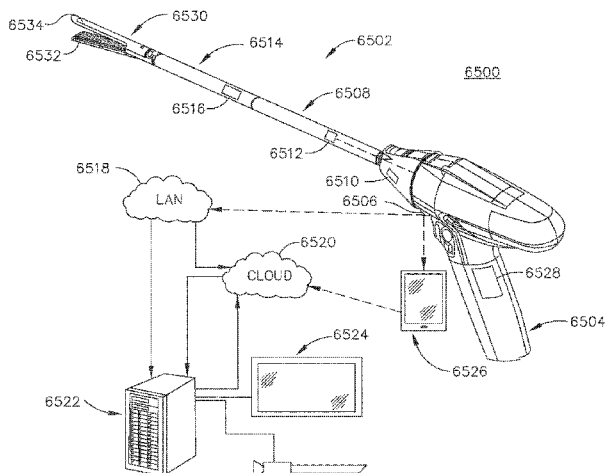
(51) **Int. Cl.**

*A61B 17/115* (2006.01)  
*A61M 1/00* (2006.01)  
*A61B 34/20* (2006.01)  
*A61B 34/32* (2006.01)  
*A61B 34/00* (2006.01)  
*A61B 90/35* (2006.01)  
*A61B 90/00* (2006.01)  
*G16H 40/67* (2006.01)  
*G16H 10/60* (2006.01)  
*G16H 50/20* (2006.01)  
*G16H 40/63* (2006.01)  
*G16H 70/20* (2006.01)  
*A61B 1/00* (2006.01)  
*A61B 1/05* (2006.01)  
*A61B 1/06* (2006.01)  
*A61B 5/00* (2006.01)

(57)

**ABSTRACT**

A method of displaying an operational parameter of a surgical system is disclosed. The method includes receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system; analyzing, by the cloud computing system, the first and the second usage data to correlate the first and the second usage data with surgical outcome data; determining, by the cloud computing system, based on the correlation, a recommended medical resource usage configuration; and displaying, on respective displays on the first and the second subset of surgical hubs, indications of the recommended medical resource usage configuration.



**Related U.S. Application Data**

(60) Provisional application No. 62/773,778, filed on Nov. 30, 2018, provisional application No. 62/773,728, filed on Nov. 30, 2018, provisional application No. 62/773,741, filed on Nov. 30, 2018, provisional application No. 62/773,742, filed on Nov. 30, 2018, provisional application No. 62/750,529, filed on Oct. 25, 2018, provisional application No. 62/750,539, filed on Oct. 25, 2018, provisional application No. 62/750,555, filed on Oct. 25, 2018, provisional application No. 62/729,183, filed on Sep. 10, 2018, provisional application No. 62/729,177, filed on Sep. 10, 2018, provisional application No. 62/729,176, filed on Sep. 10, 2018, provisional application No. 62/729,185, filed on Sep. 10, 2018, provisional application No. 62/729,184, filed on Sep. 10, 2018, provisional application No. 62/729,182, filed on Sep. 10, 2018, provisional application No. 62/729,191, filed on Sep. 10, 2018, provisional application No. 62/729,195, filed on Sep. 10, 2018, provisional application No. 62/729,186, filed on Sep. 10, 2018, provisional application No. 62/721,995, filed on Aug. 23, 2018, provisional application No. 62/721,998, filed on Aug. 23, 2018, provisional application No. 62/721,999, filed on Aug. 23, 2018, provisional application No. 62/721,994, filed on Aug. 23, 2018, provisional application No. 62/721,996, filed on Aug. 23, 2018, provisional application No. 62/692,747, filed on Jun. 30, 2018, provisional application No. 62/692,748, filed on Jun. 30, 2018, provisional application No. 62/691,228, filed on Jun. 28, 2018, provisional application No. 62/691,227, filed on Jun. 28, 2018, provisional application No. 62/691,230, filed on Jun. 28, 2018, provisional application No. 62/691,219, filed on Jun. 28, 2018, provisional application No. 62/691,257, filed on Jun. 28, 2018, provisional application No. 62/691,262, filed on Jun. 28, 2018, provisional application No. 62/691,251, filed on Jun. 28, 2018, provisional application No. 62/665,129, filed on May 1, 2018, provisional application No. 62/665,139, filed on May 1, 2018, provisional application No. 62/665,177, filed on May 1, 2018, provisional application No. 62/665,128, filed

on May 1, 2018, provisional application No. 62/665,192, filed on May 1, 2018, provisional application No. 62/665,134, filed on May 1, 2018, provisional application No. 62/659,900, filed on Apr. 19, 2018, provisional application No. 62/650,898, filed on Mar. 30, 2018, provisional application No. 62/650,887, filed on Mar. 30, 2018, provisional application No. 62/650,882, filed on Mar. 30, 2018, provisional application No. 62/650,877, filed on Mar. 30, 2018, provisional application No. 62/649,302, filed on Mar. 28, 2018, provisional application No. 62/649,294, filed on Mar. 28, 2018, provisional application No. 62/649,300, filed on Mar. 28, 2018, provisional application No. 62/649,309, filed on Mar. 28, 2018, provisional application No. 62/649,310, filed on Mar. 28, 2018, provisional application No. 62/649,291, filed on Mar. 28, 2018, provisional application No. 62/649,296, filed on Mar. 28, 2018, provisional application No. 62/649,333, filed on Mar. 28, 2018, provisional application No. 62/649,327, filed on Mar. 28, 2018, provisional application No. 62/649,315, filed on Mar. 28, 2018, provisional application No. 62/649,313, filed on Mar. 28, 2018, provisional application No. 62/649,320, filed on Mar. 28, 2018, provisional application No. 62/649,307, filed on Mar. 28, 2018, provisional application No. 62/649,323, filed on Mar. 28, 2018, provisional application No. 62/611,341, filed on Dec. 28, 2017, provisional application No. 62/611,340, filed on Dec. 28, 2017, provisional application No. 62/611,339, filed on Dec. 28, 2017.

**Publication Classification**

(51) **Int. Cl.**  
*A61B 18/14* (2006.01)  
*B25J 9/16* (2006.01)  
*B25J 13/00* (2006.01)  
*G06K 7/10* (2006.01)  
*G06K 19/077* (2006.01)  
*H01Q 1/22* (2006.01)  
*H04L 29/06* (2006.01)  
*H04L 29/08* (2006.01)  
*H04N 5/272* (2006.01)  
*H04N 7/18* (2006.01)  
*H05K 1/02* (2006.01)  
*H05K 1/18* (2006.01)

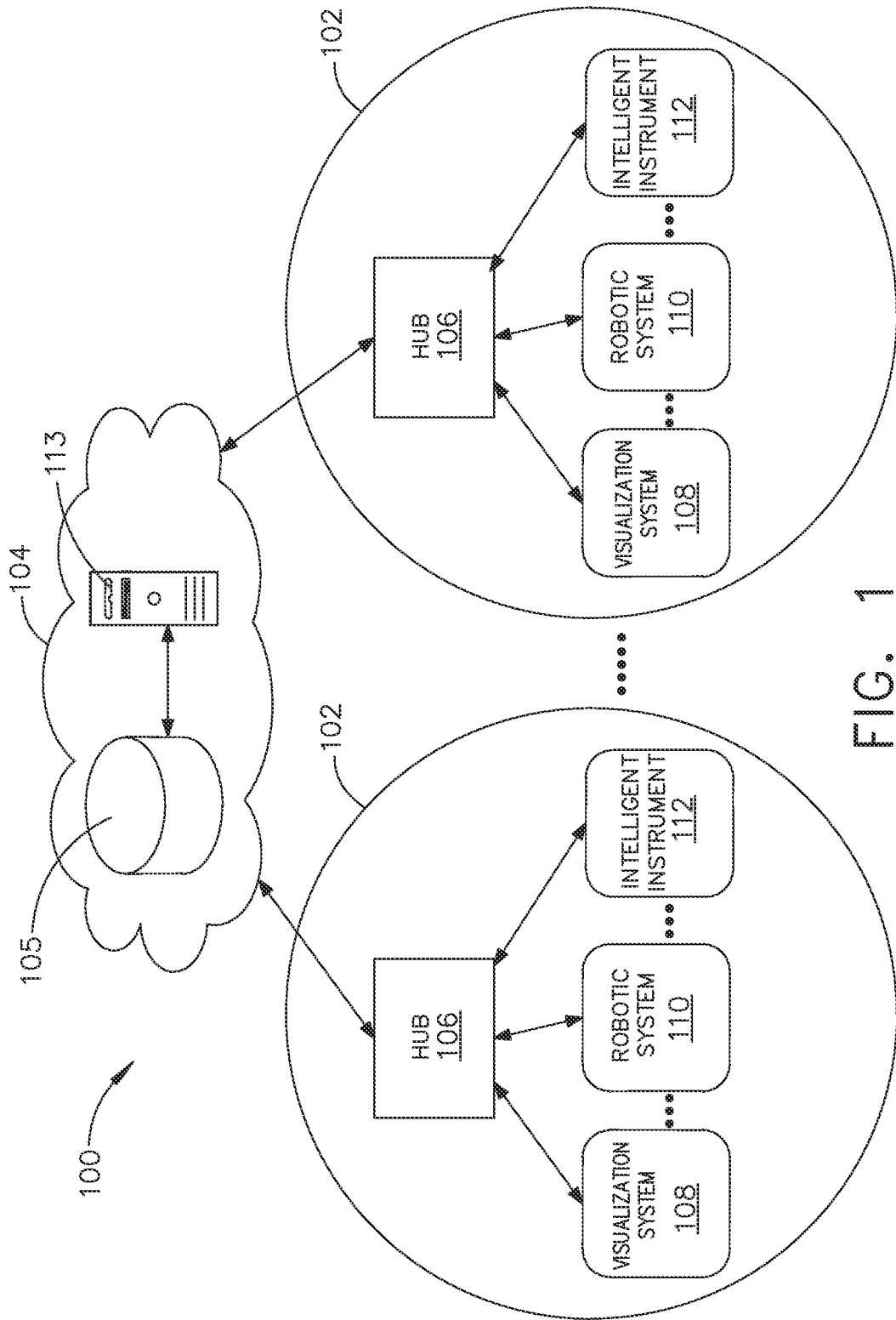


FIG. 1

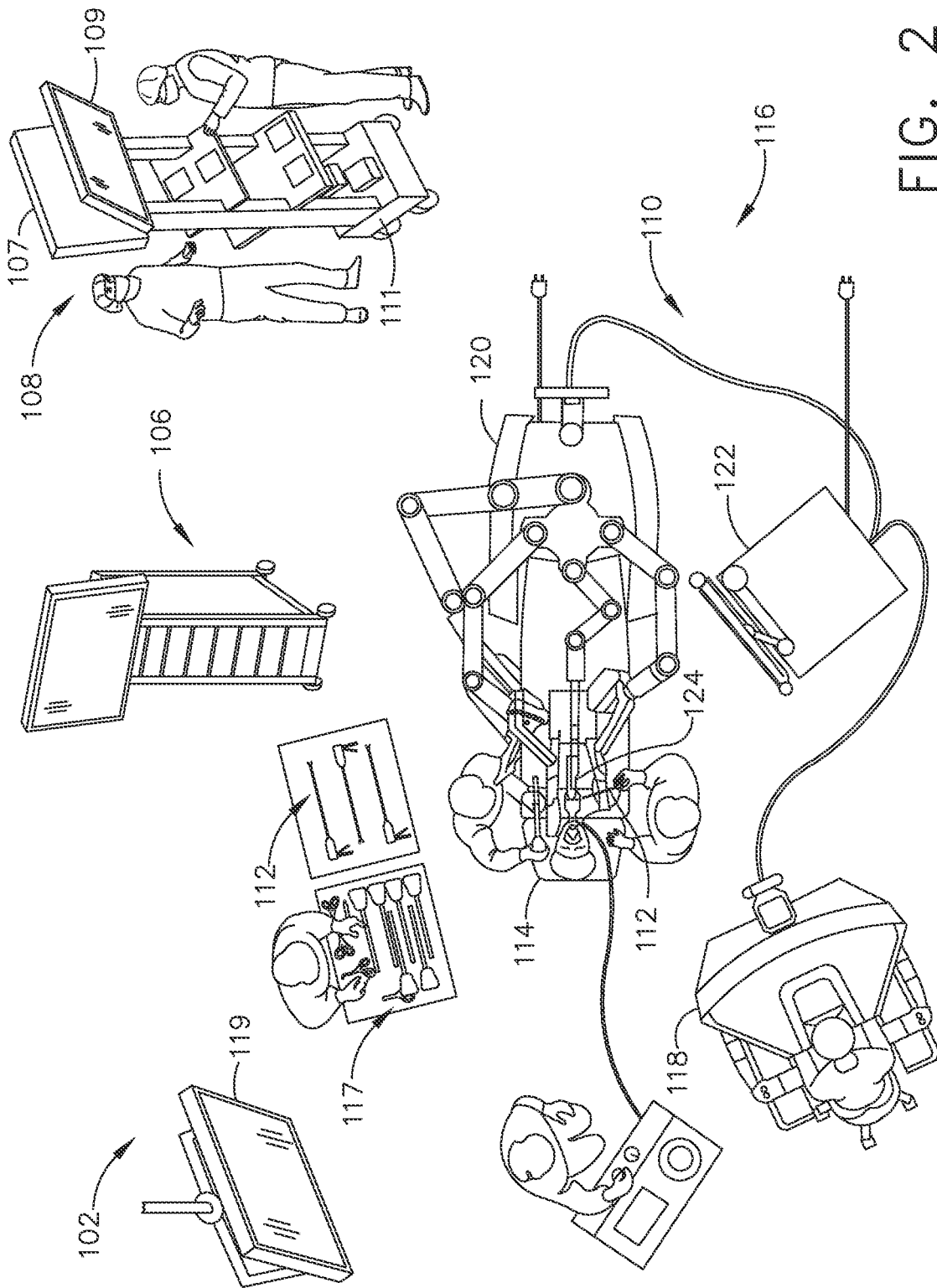


FIG. 2

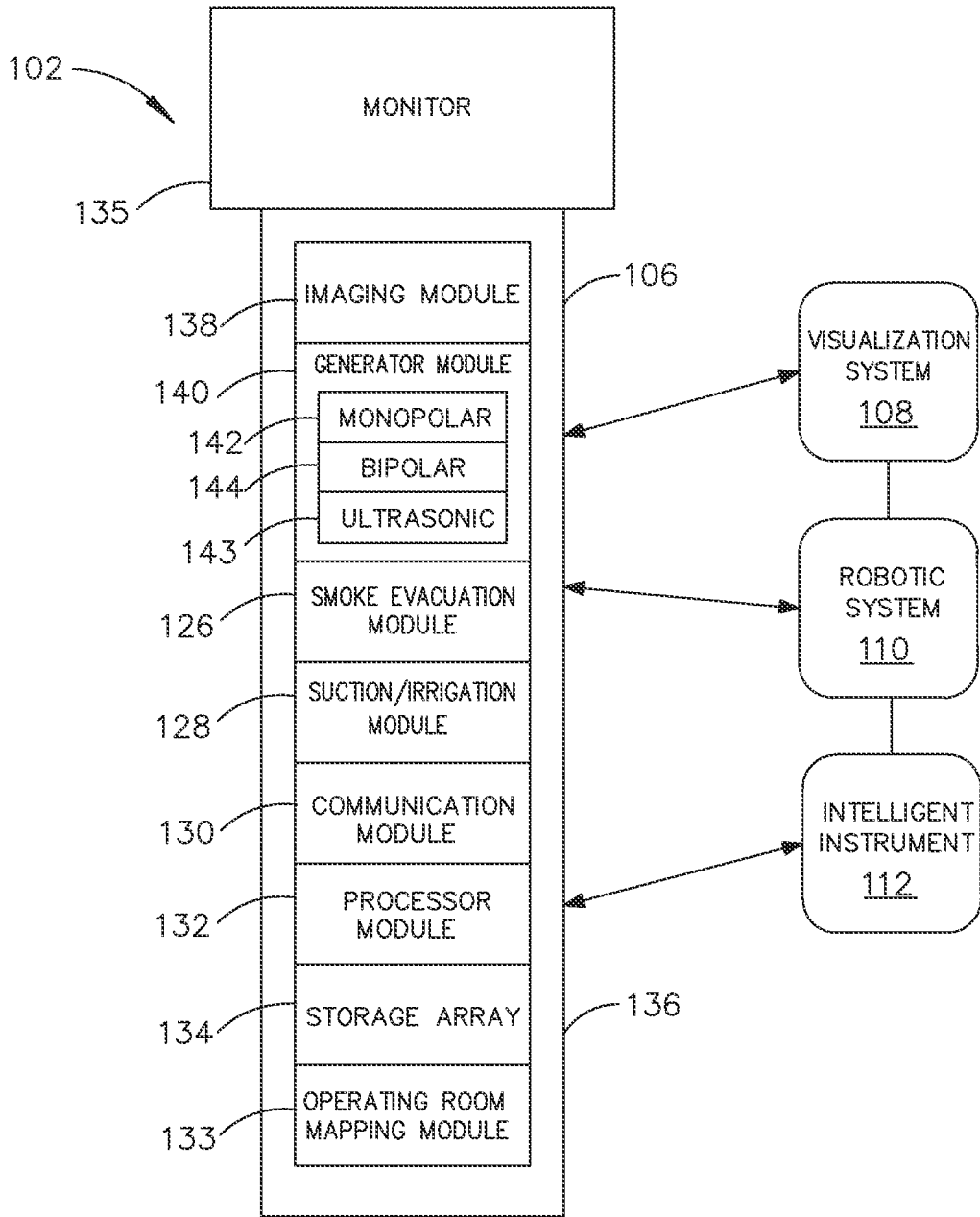


FIG. 3

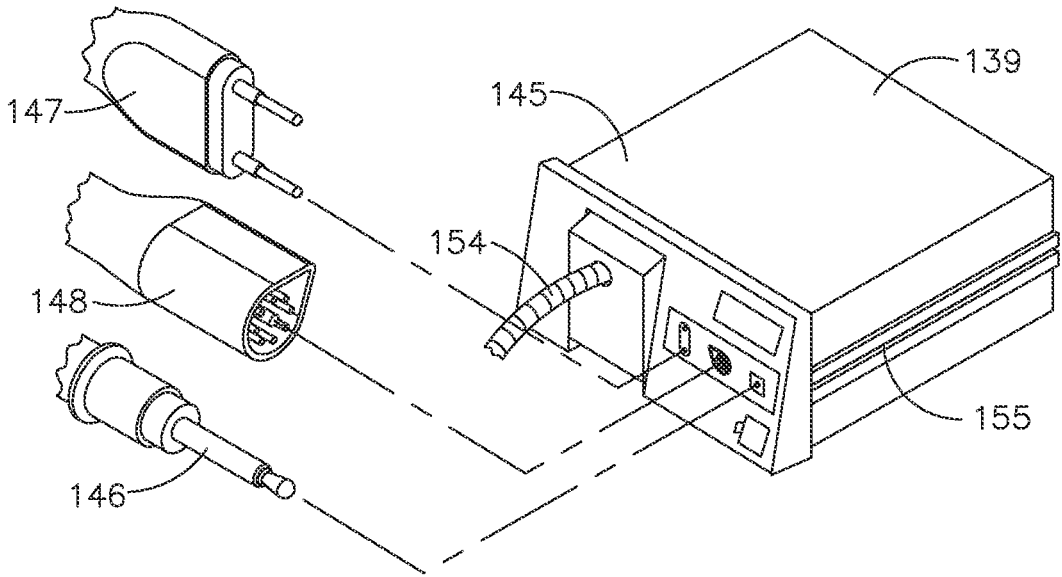


FIG. 5

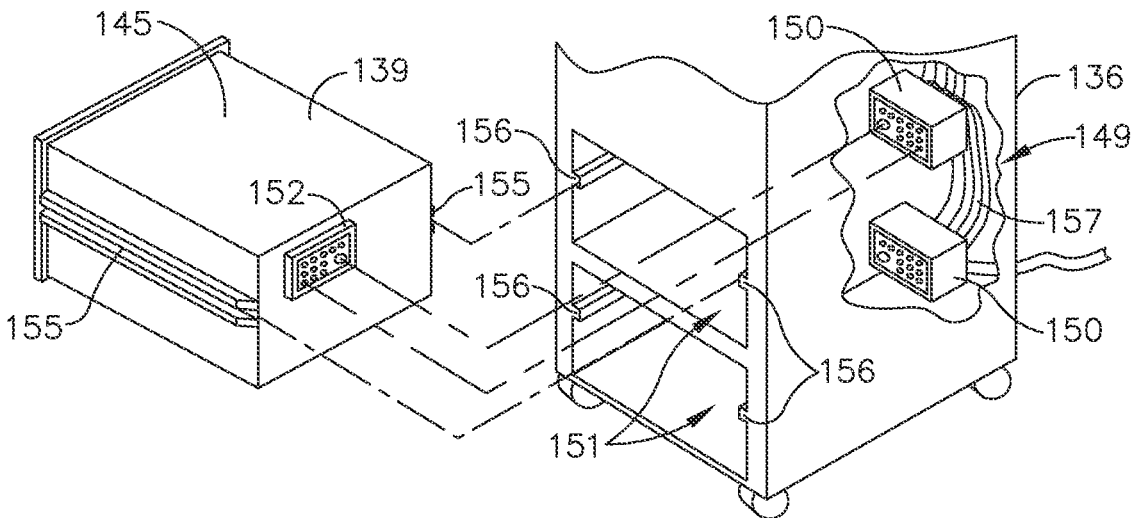


FIG. 4

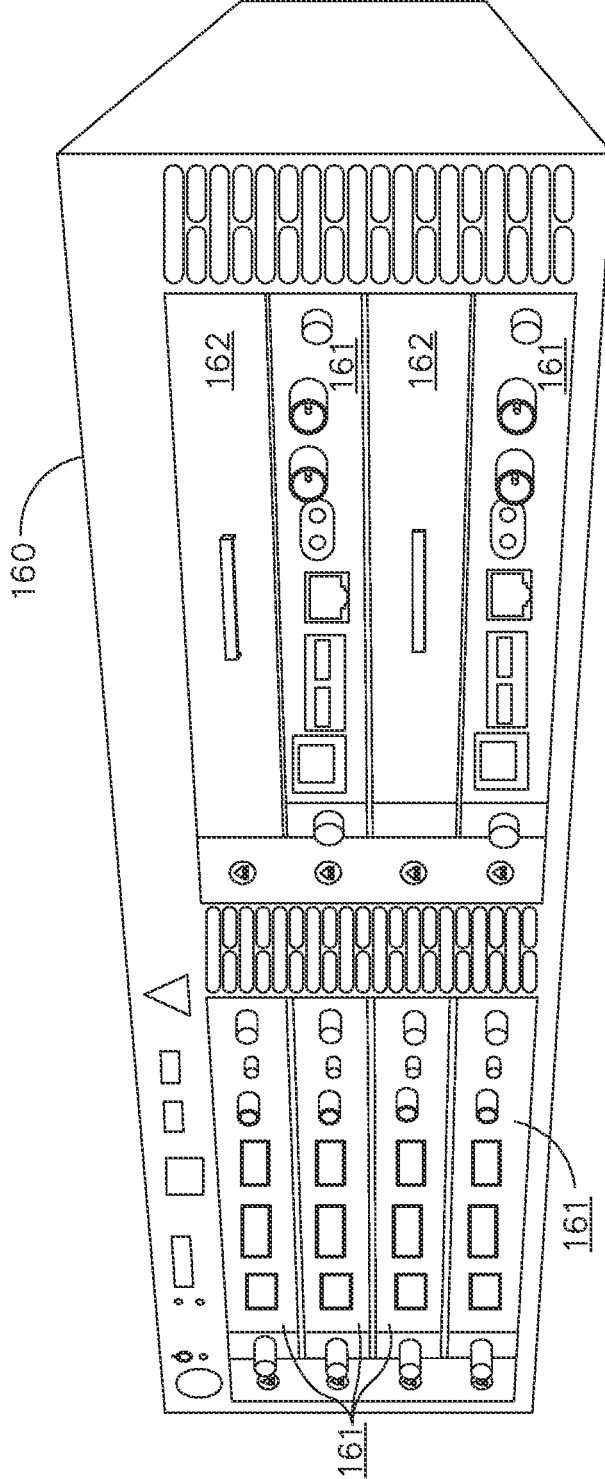


FIG. 6

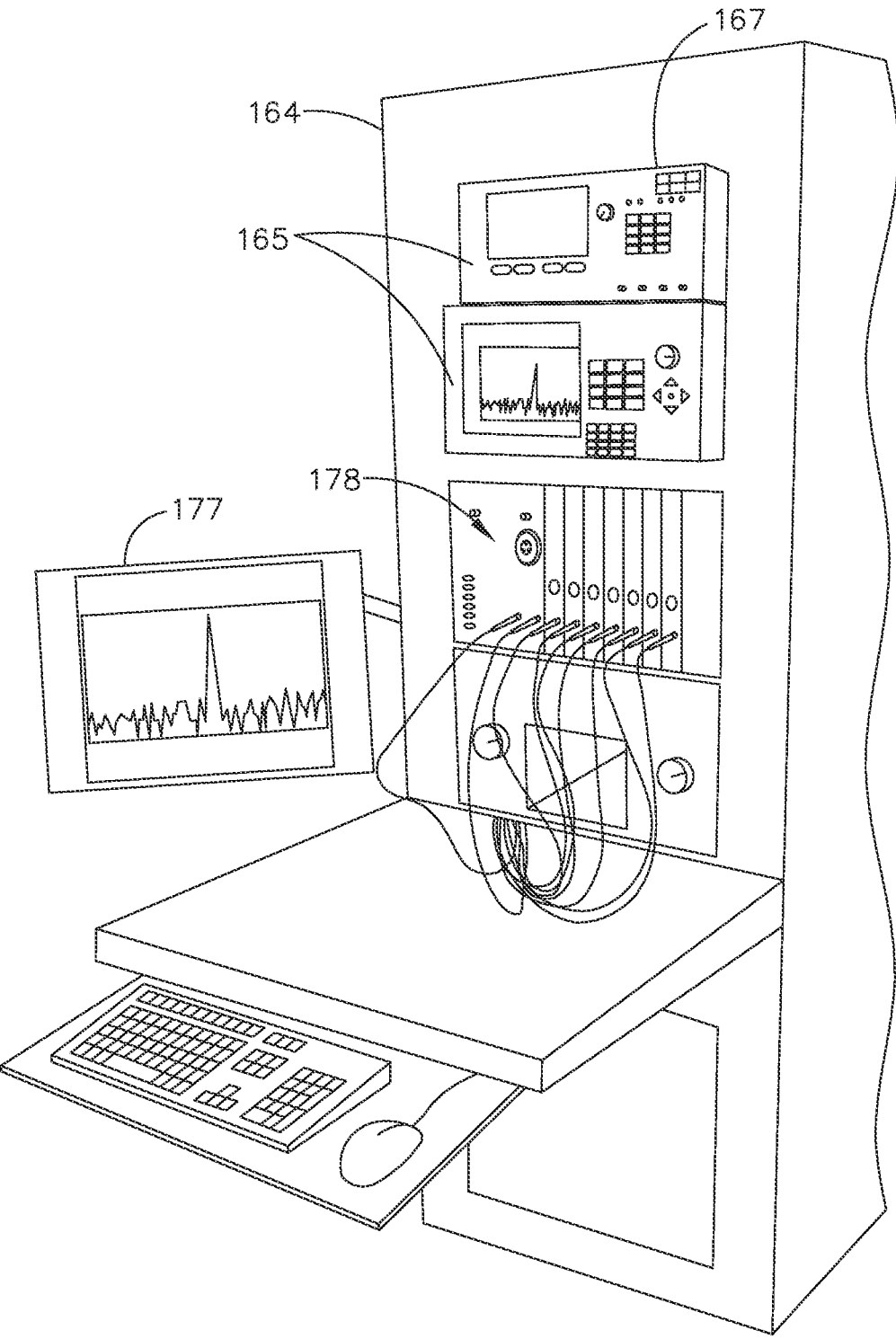


FIG. 7



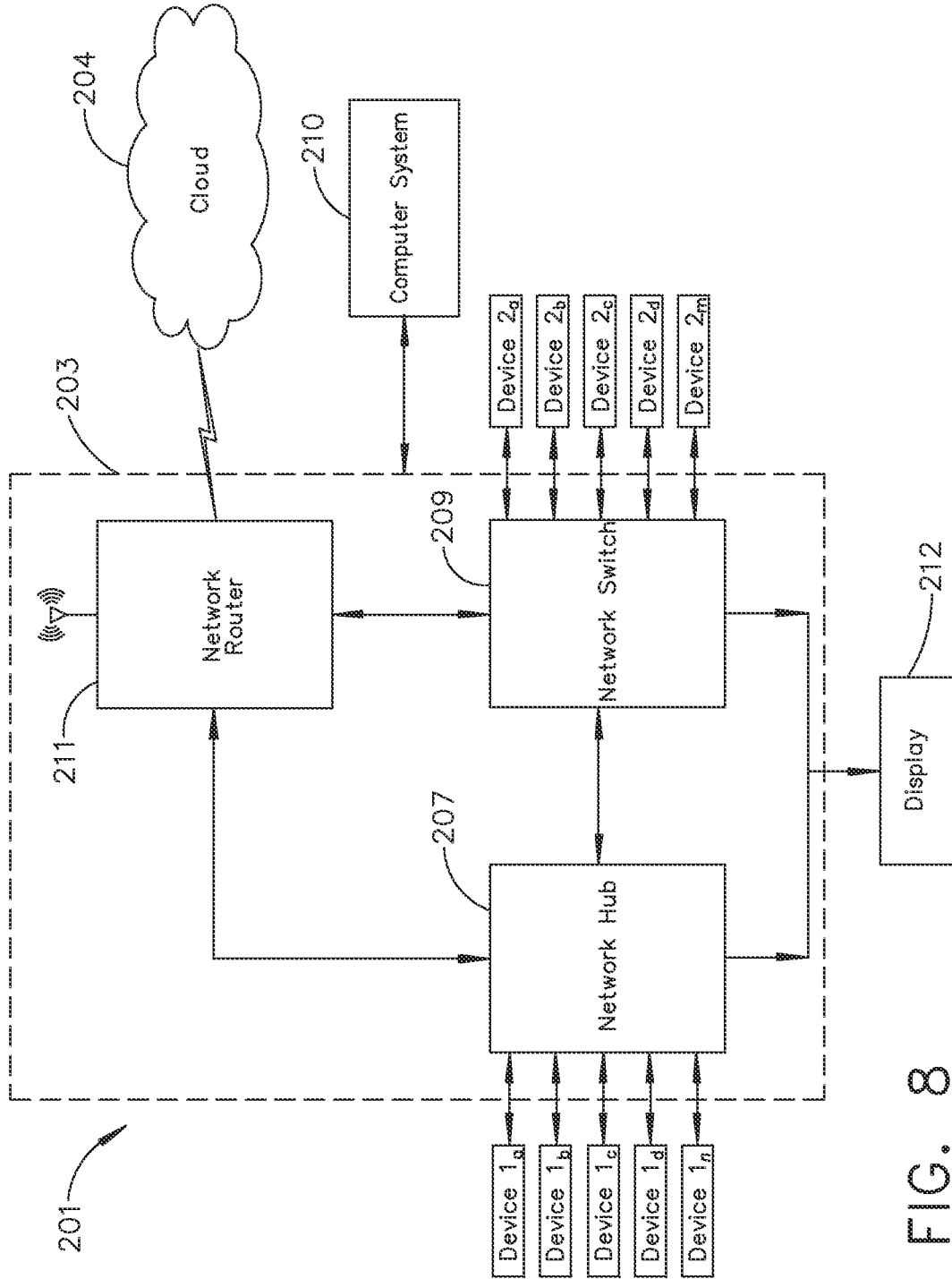


FIG. 8

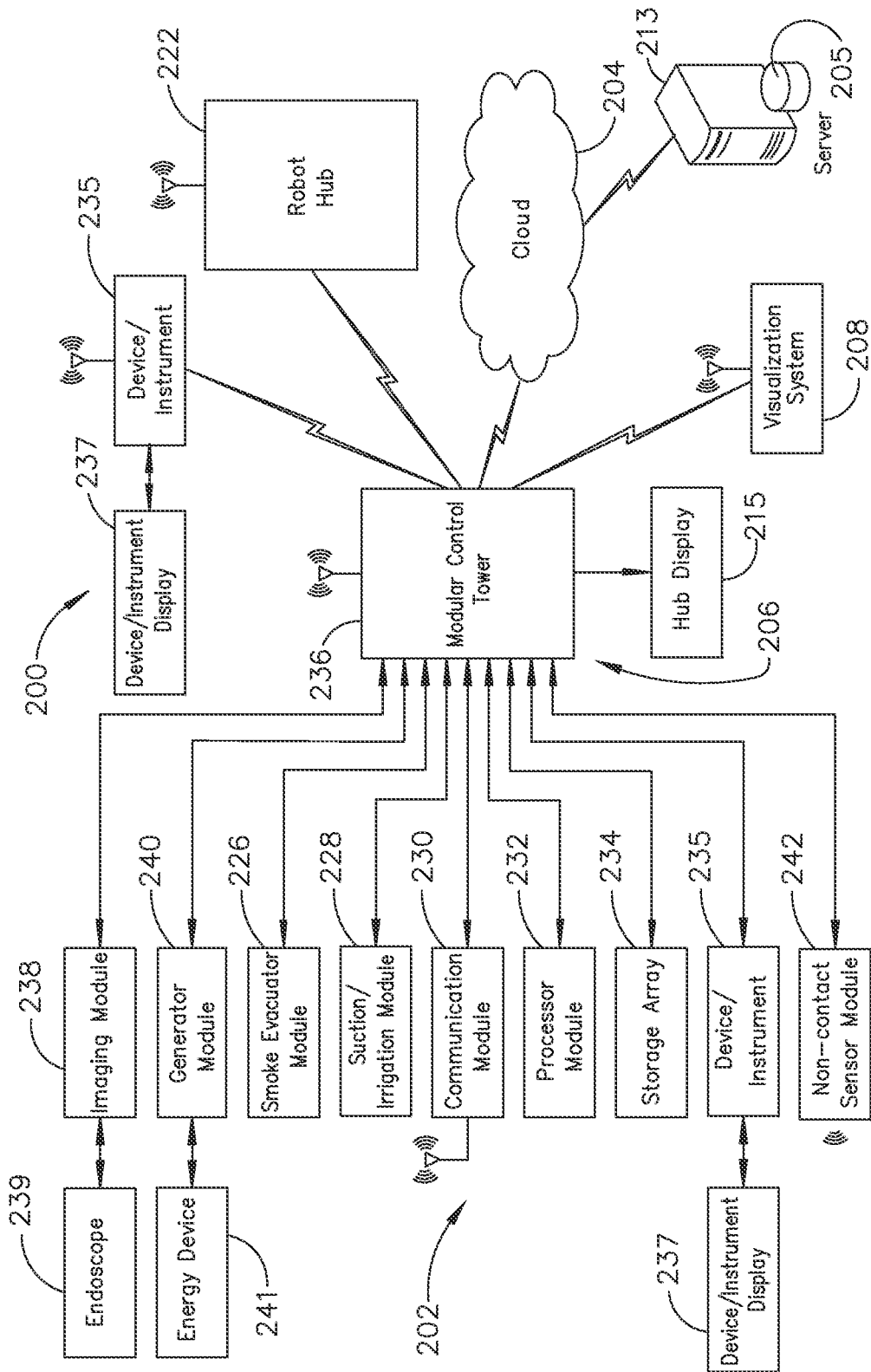


FIG. 9

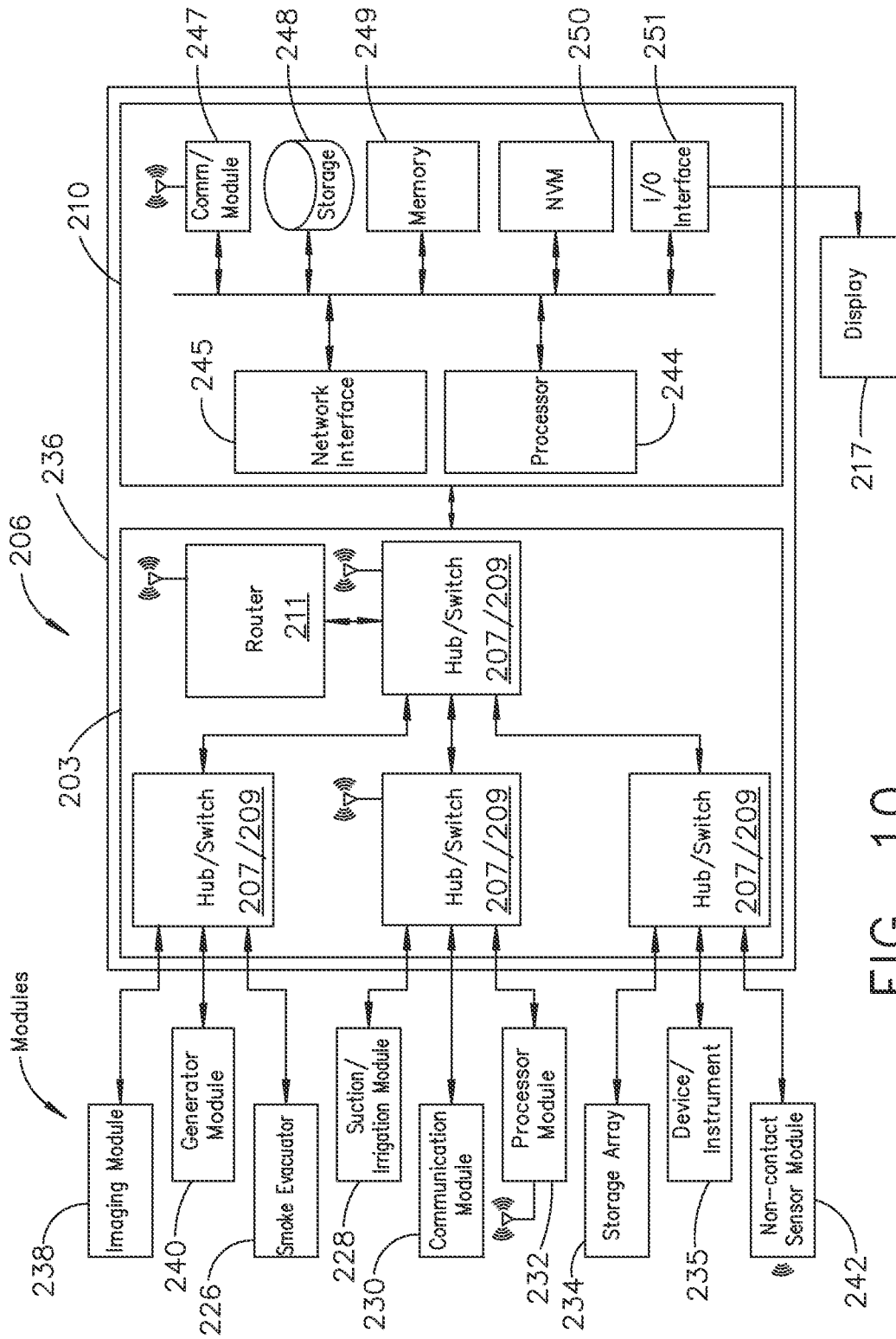


FIG. 10

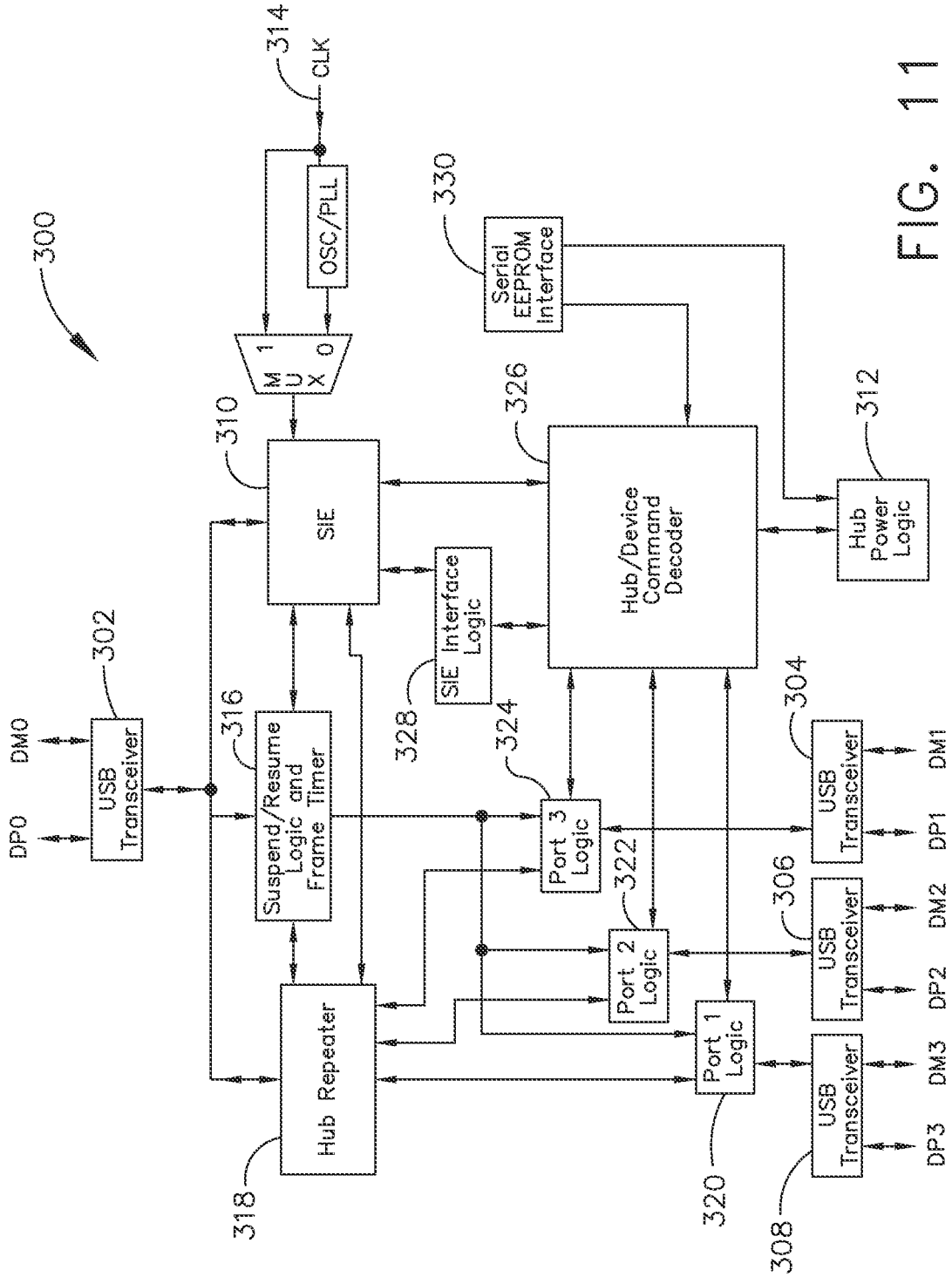


FIG. 11

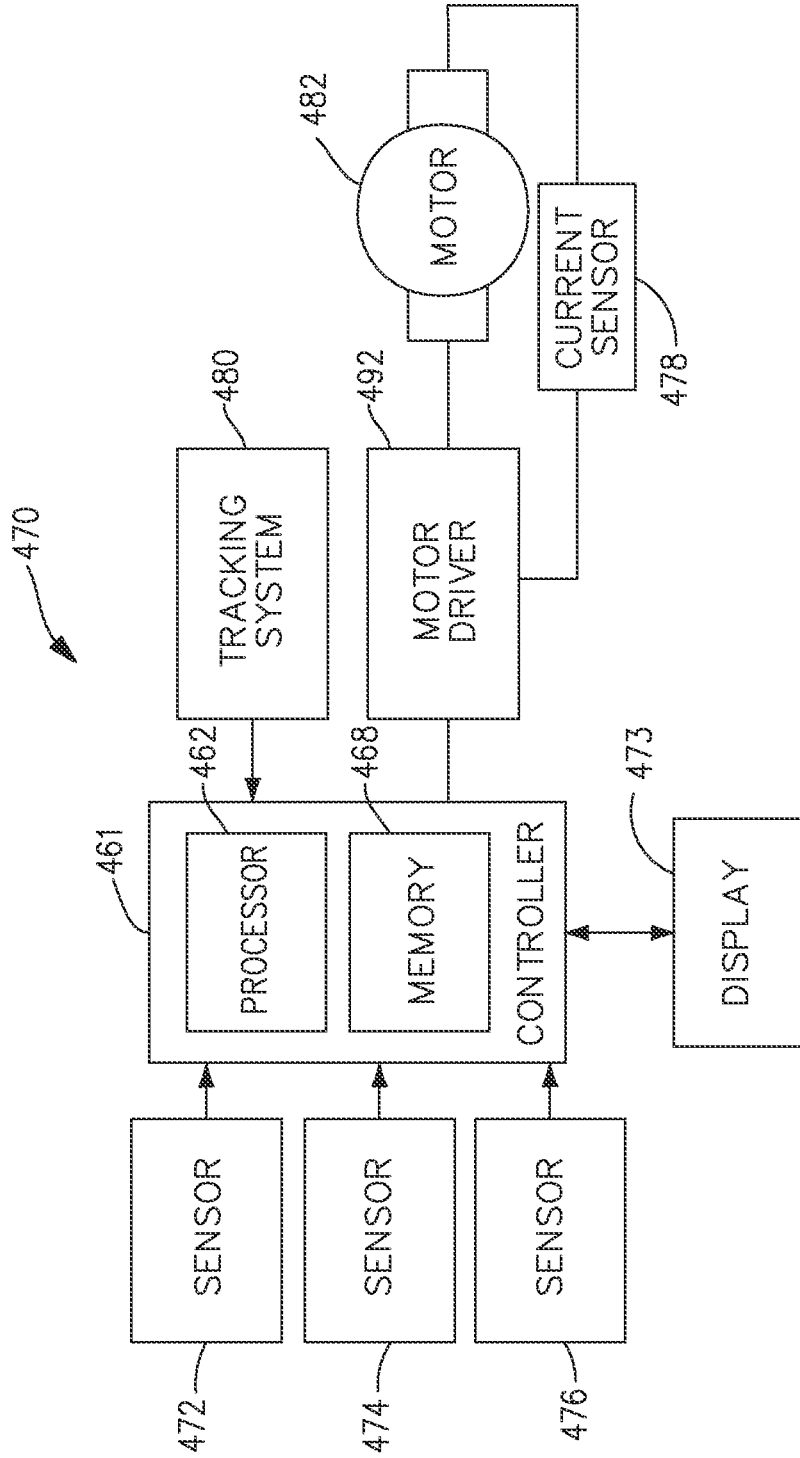


FIG. 12

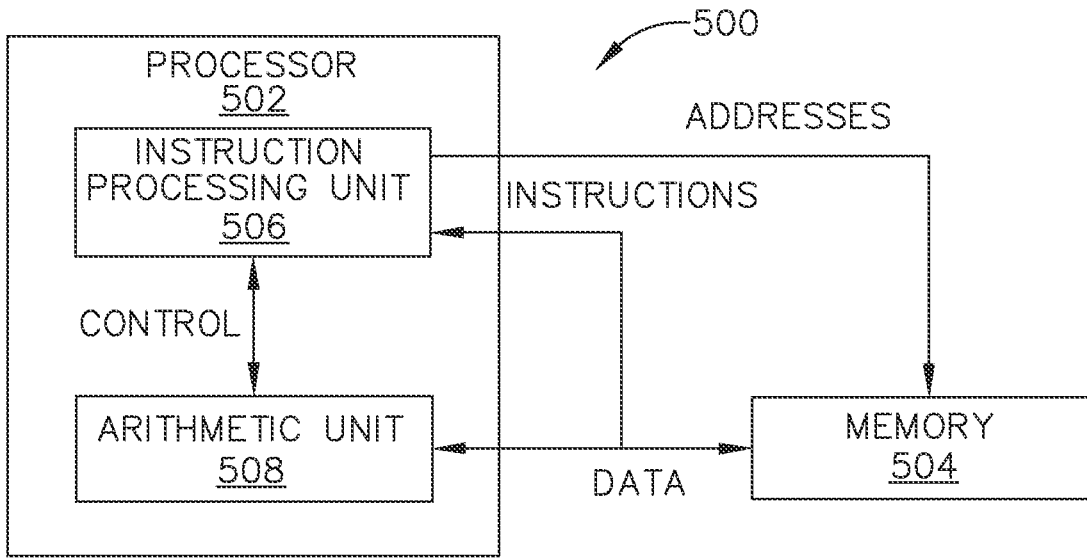


FIG. 13

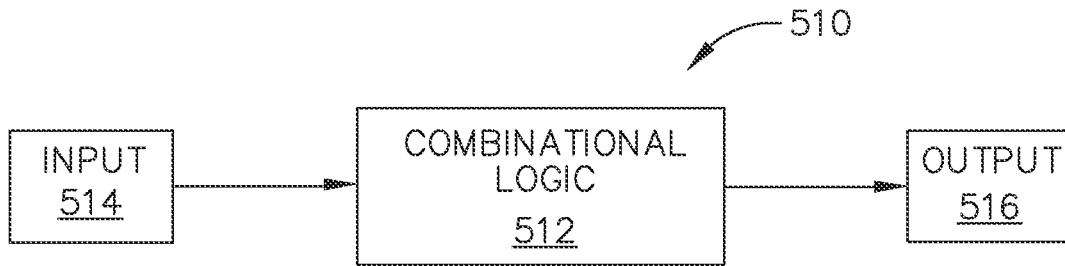


FIG. 14

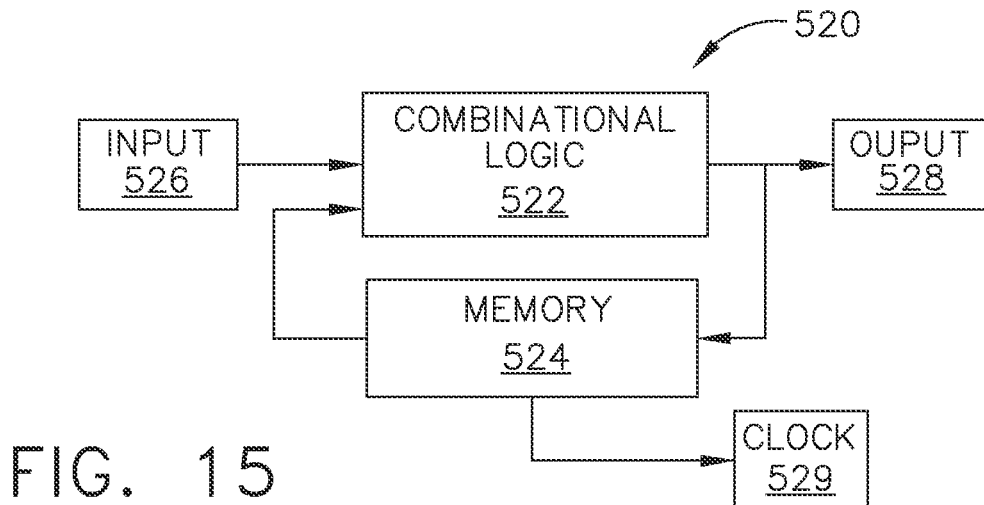


FIG. 15

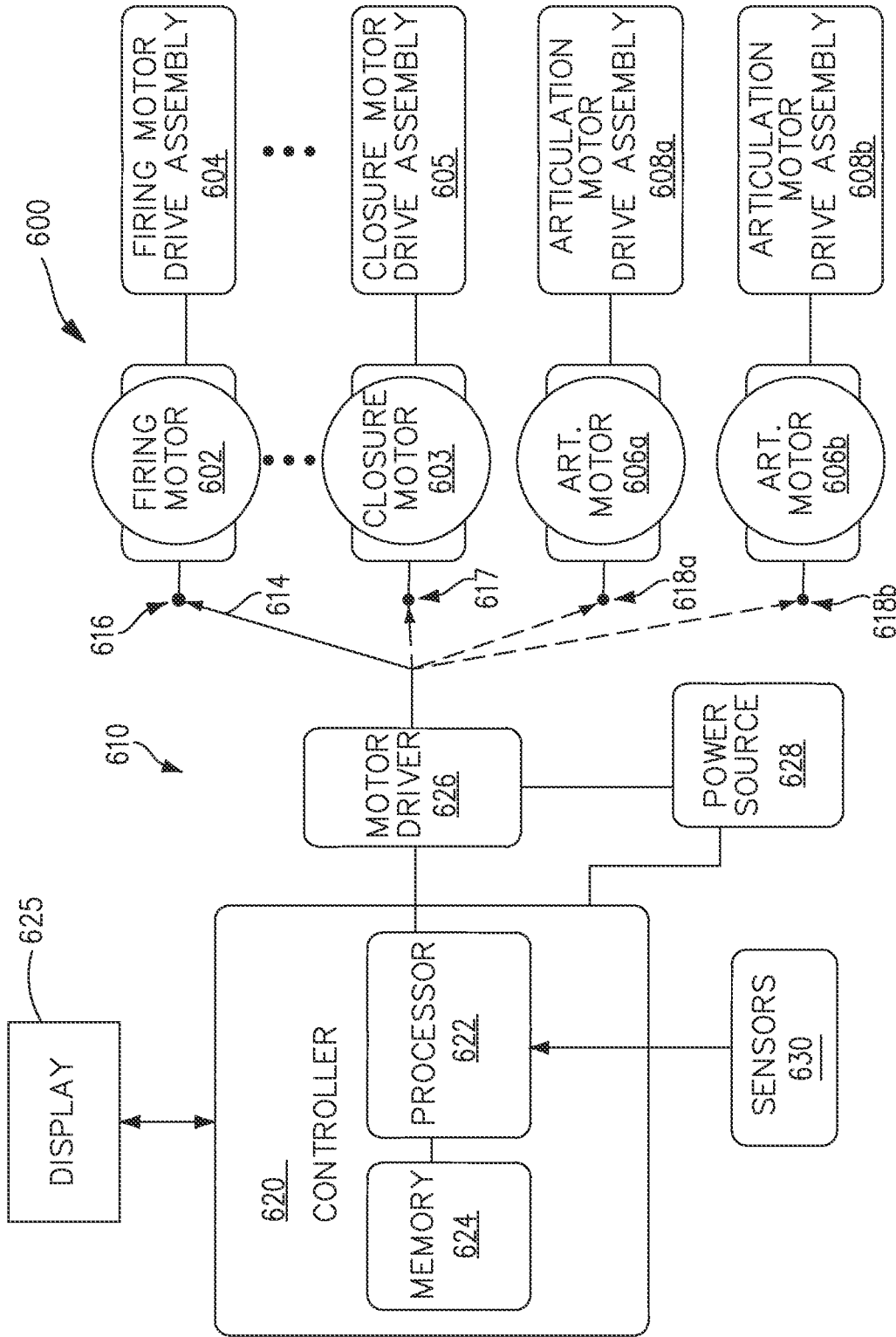


FIG. 16

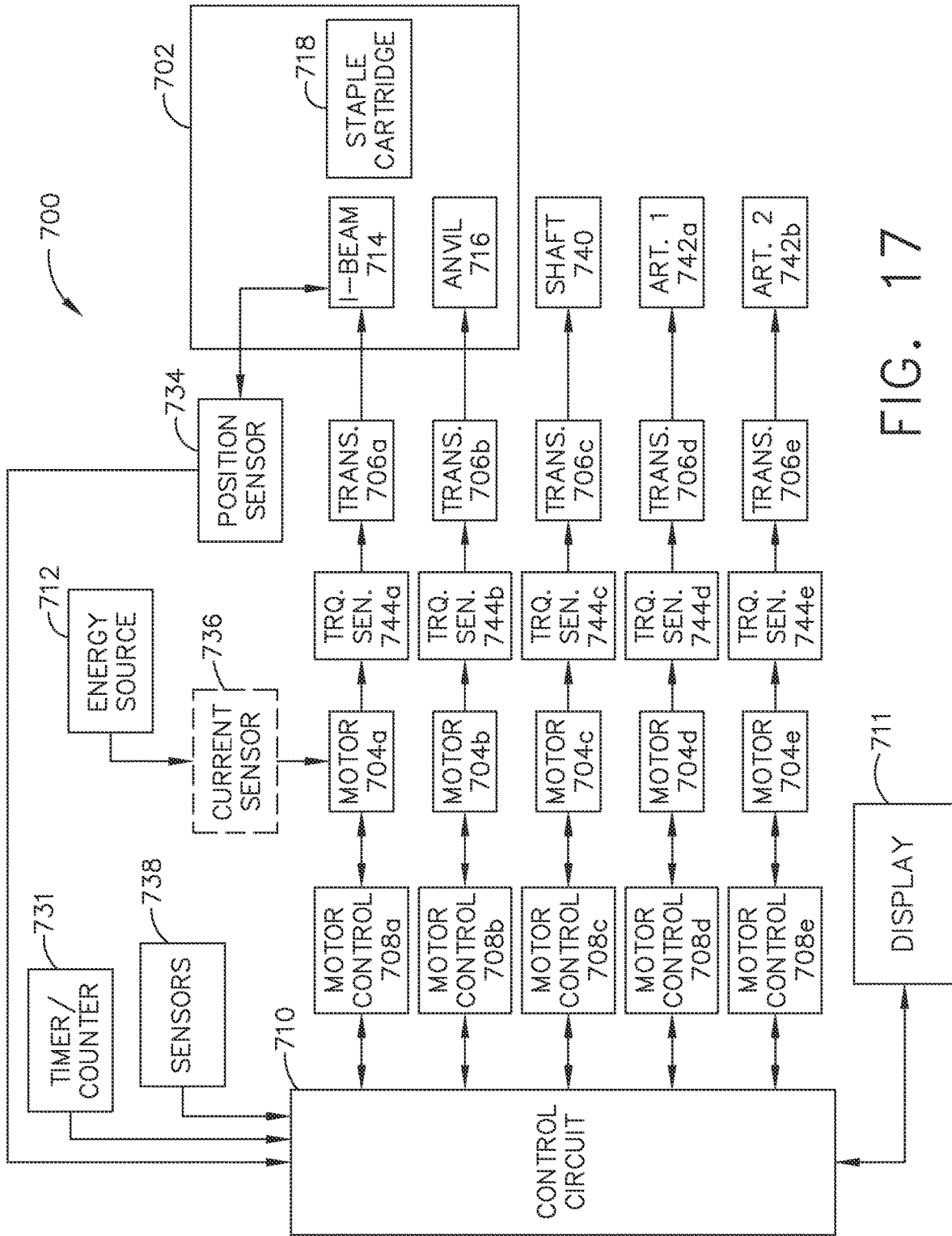


FIG. 17



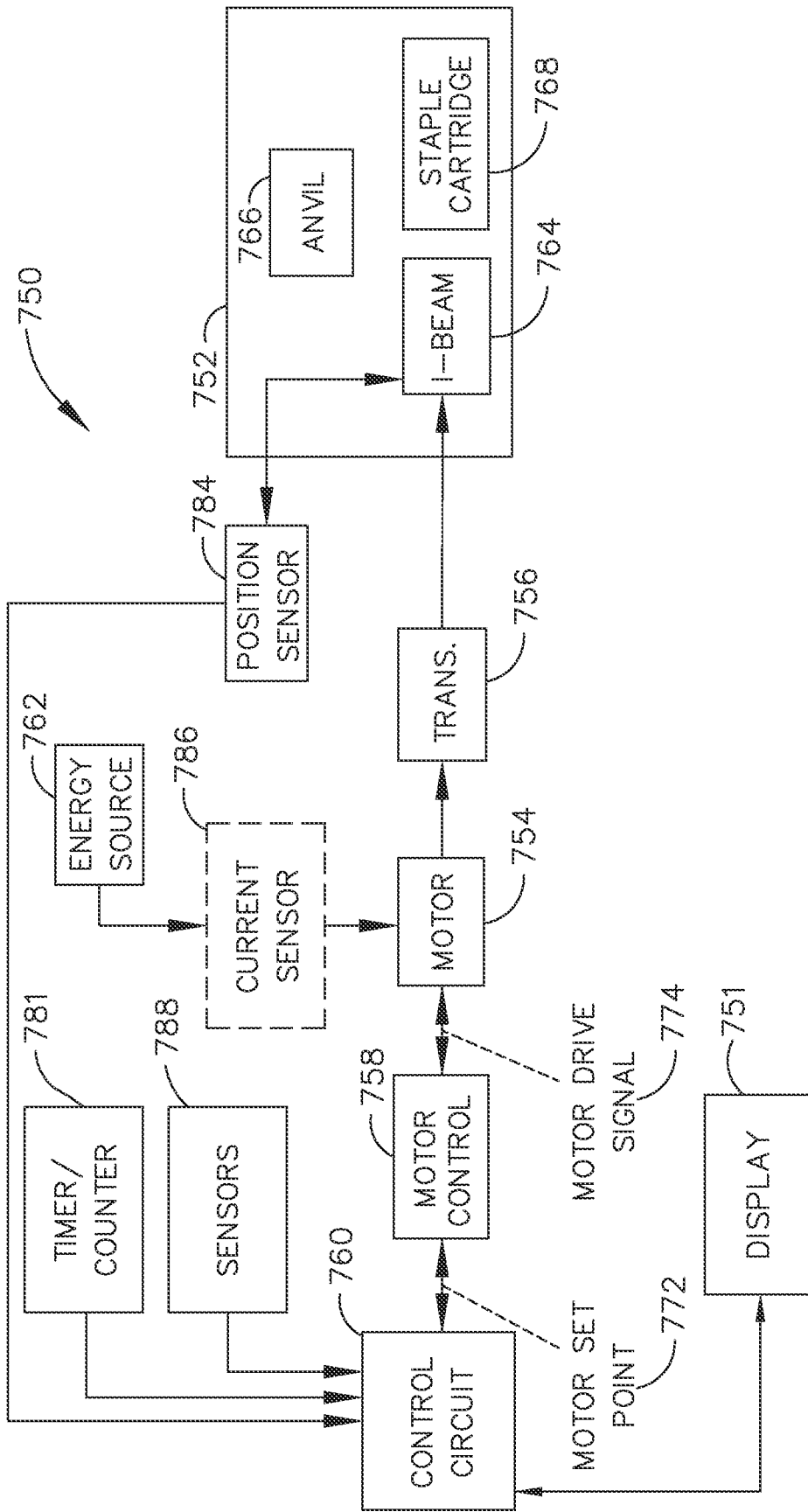


FIG. 18

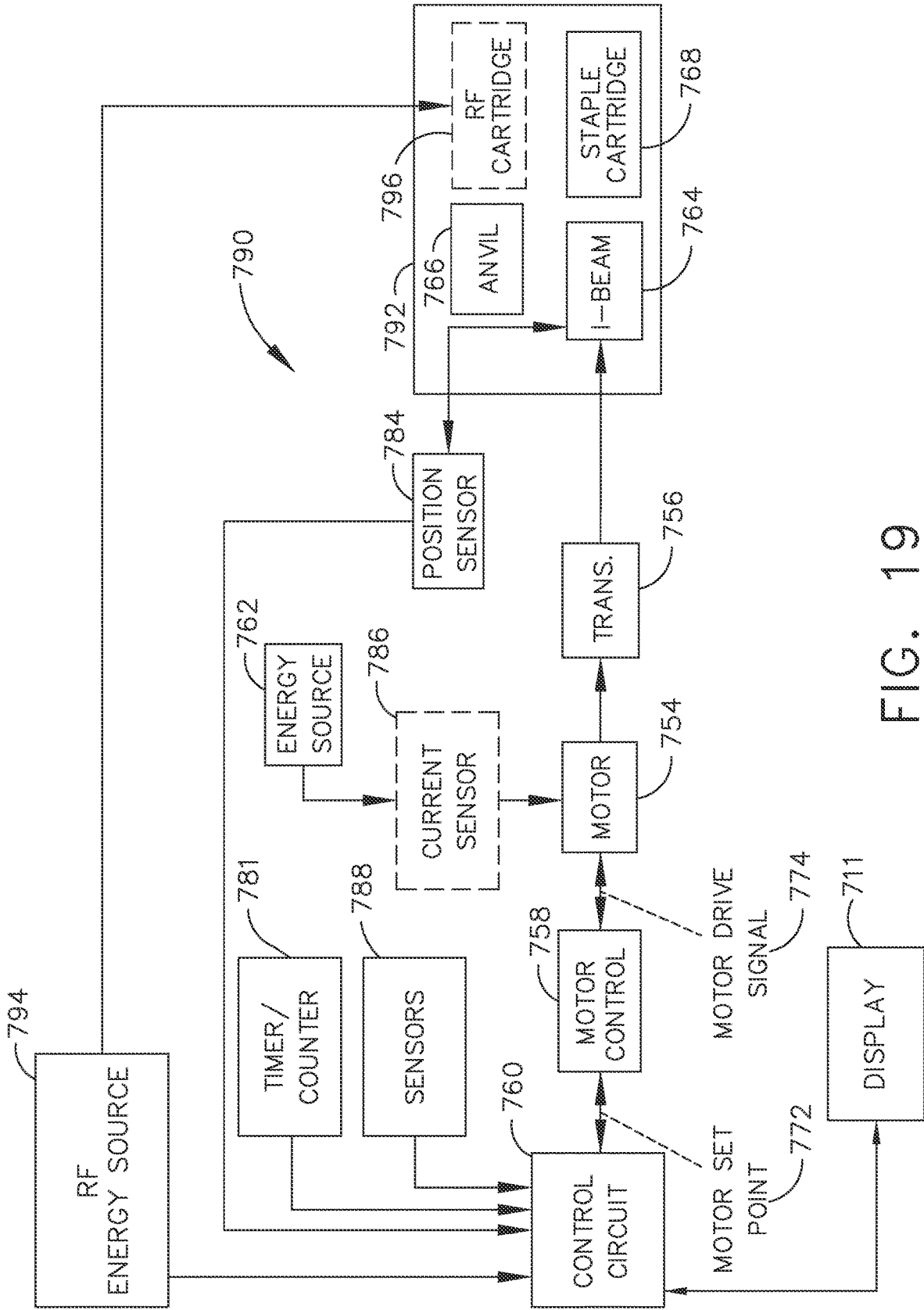


FIG. 19

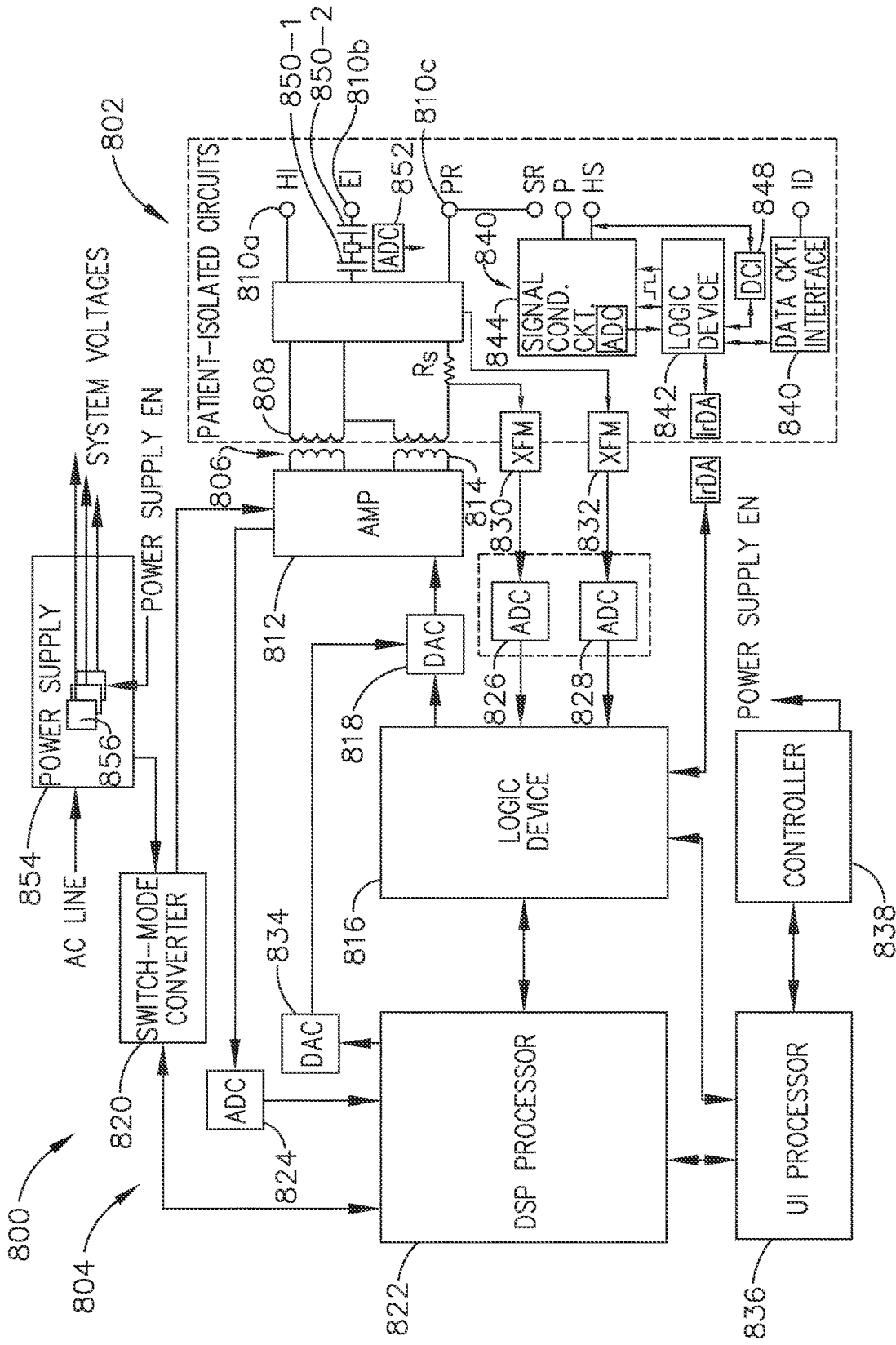


FIG. 20

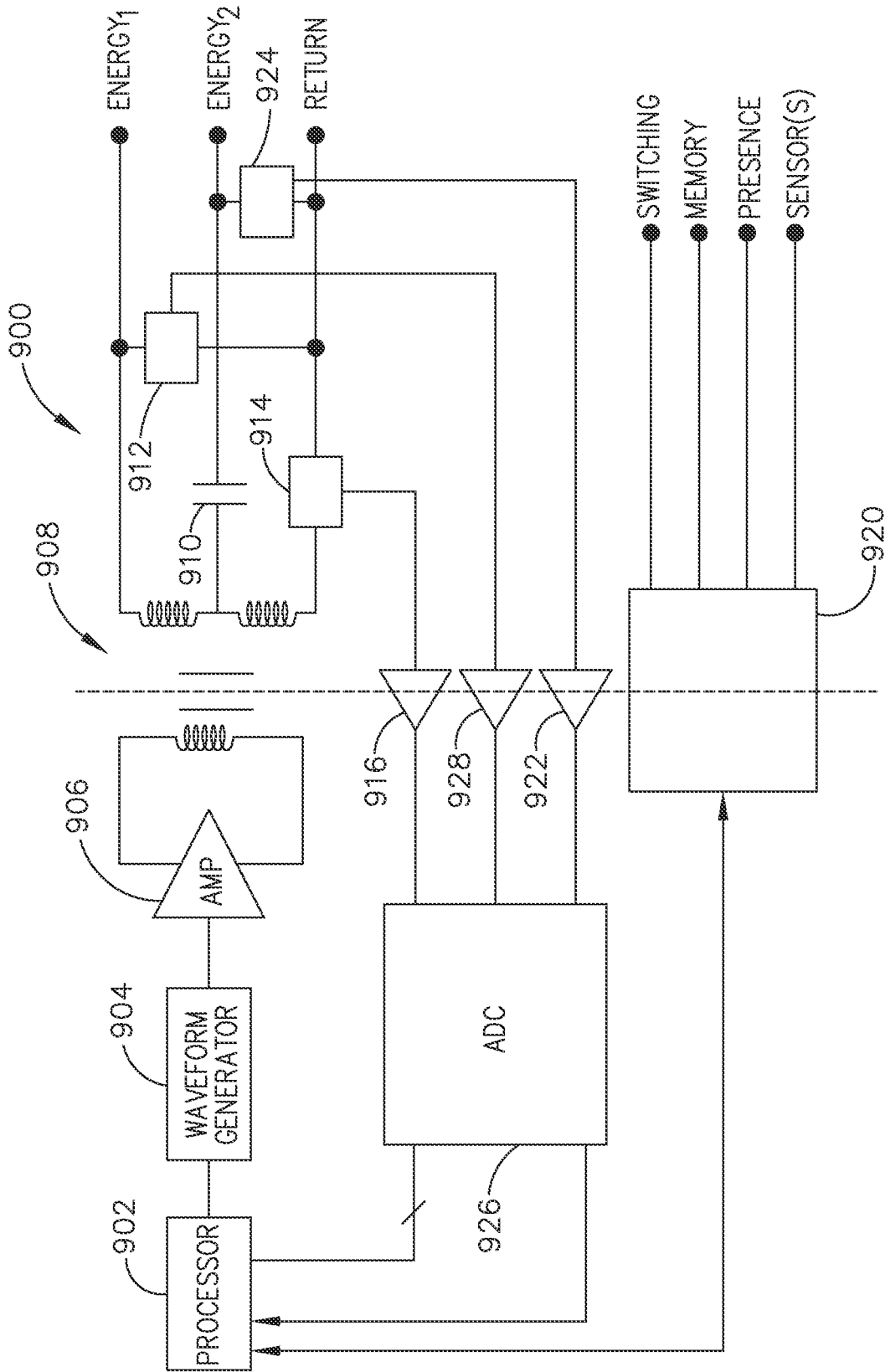


FIG. 21

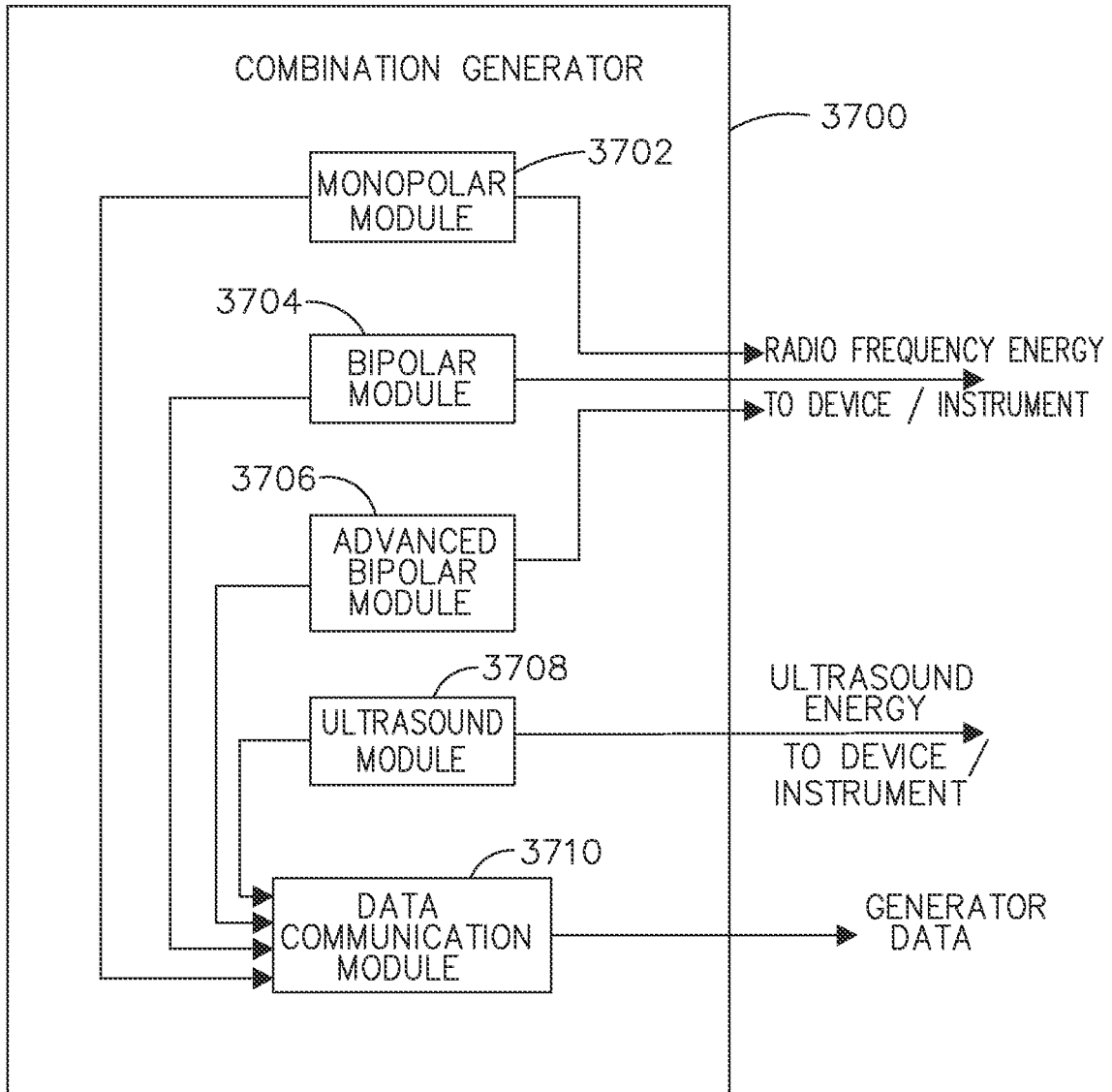


FIG. 22

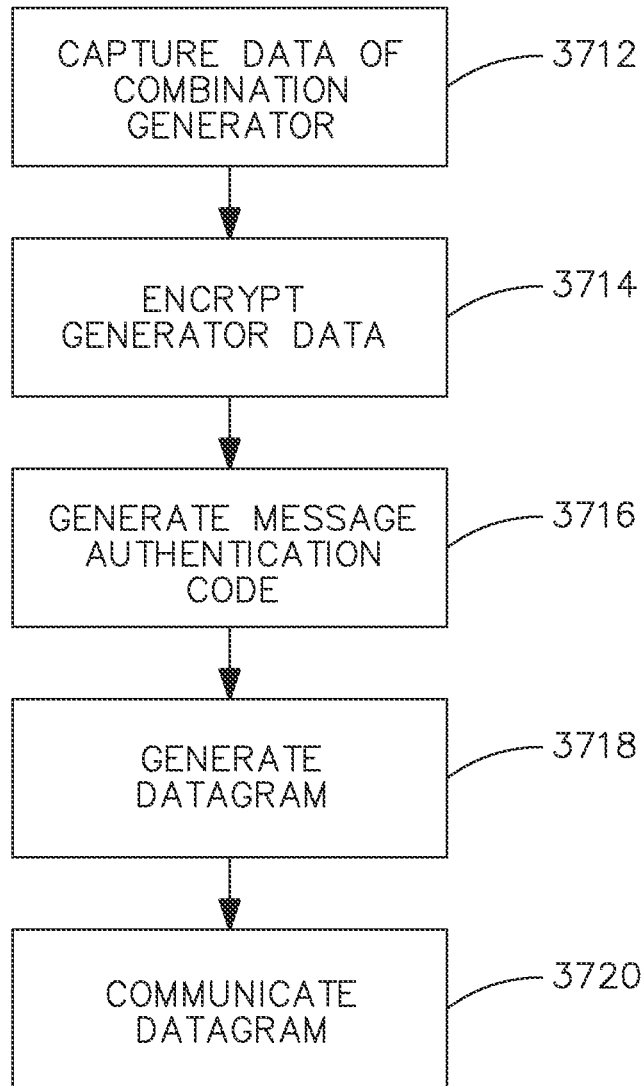


FIG. 23

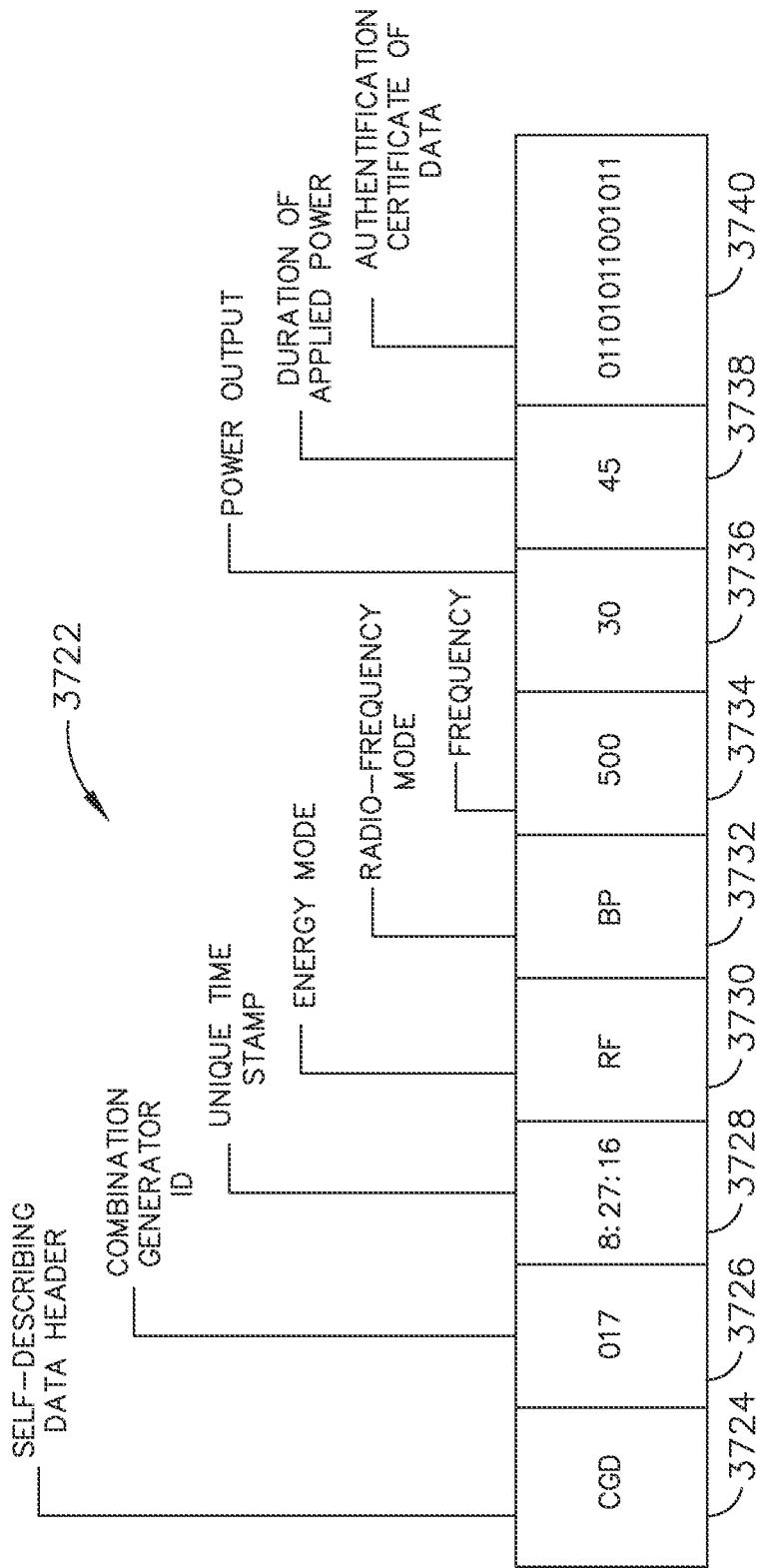


FIG. 24

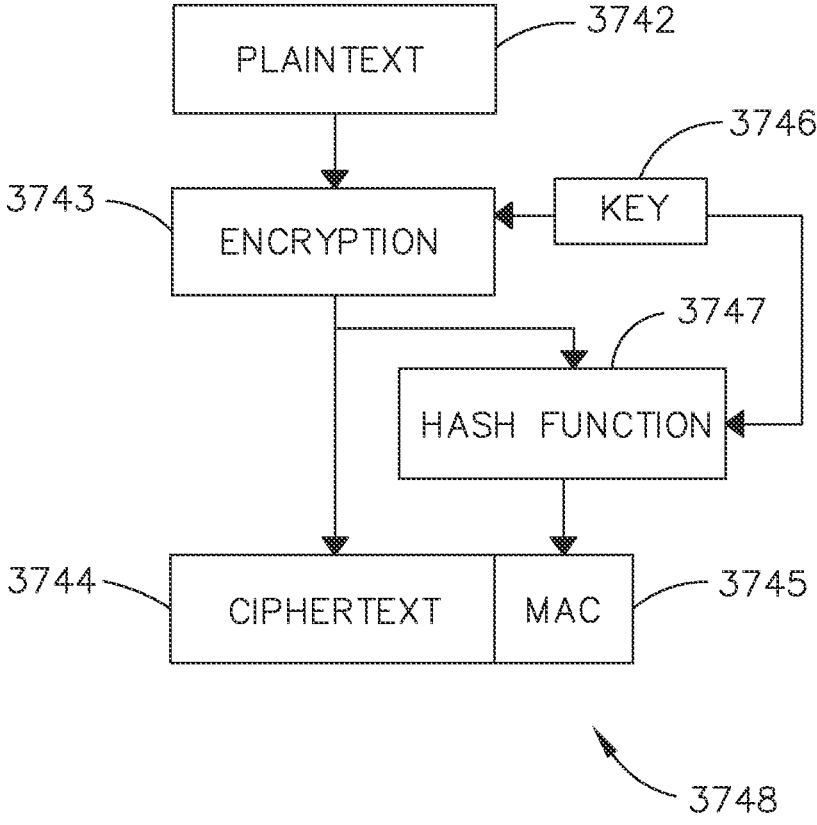


FIG. 25



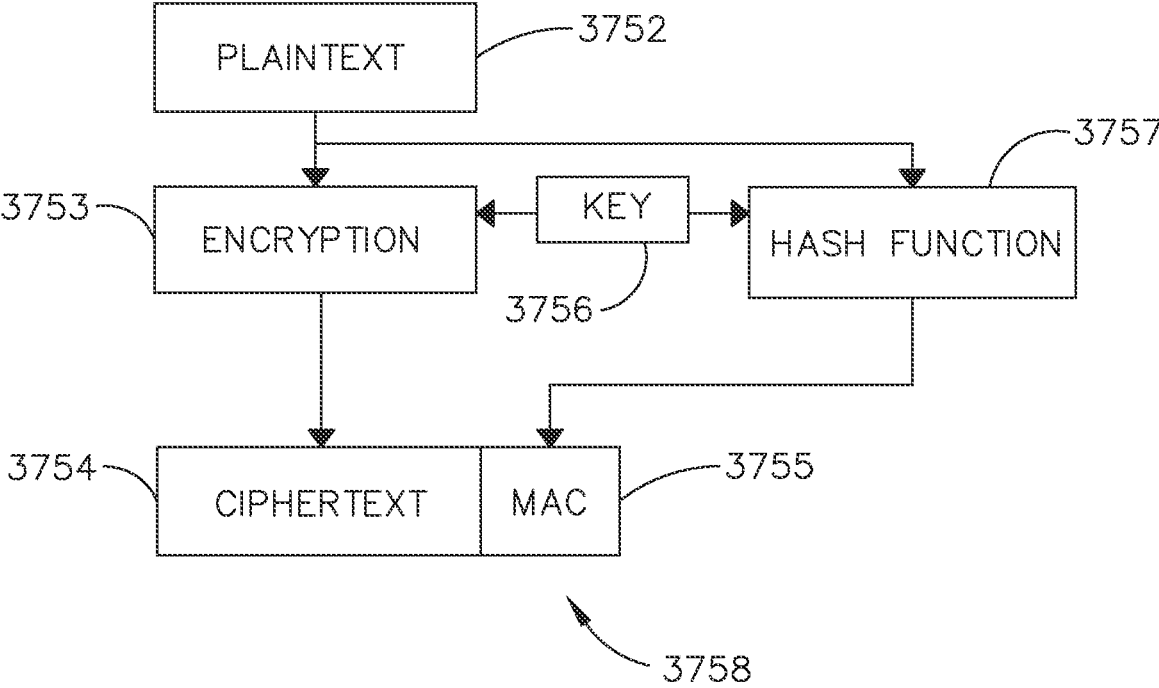


FIG. 26

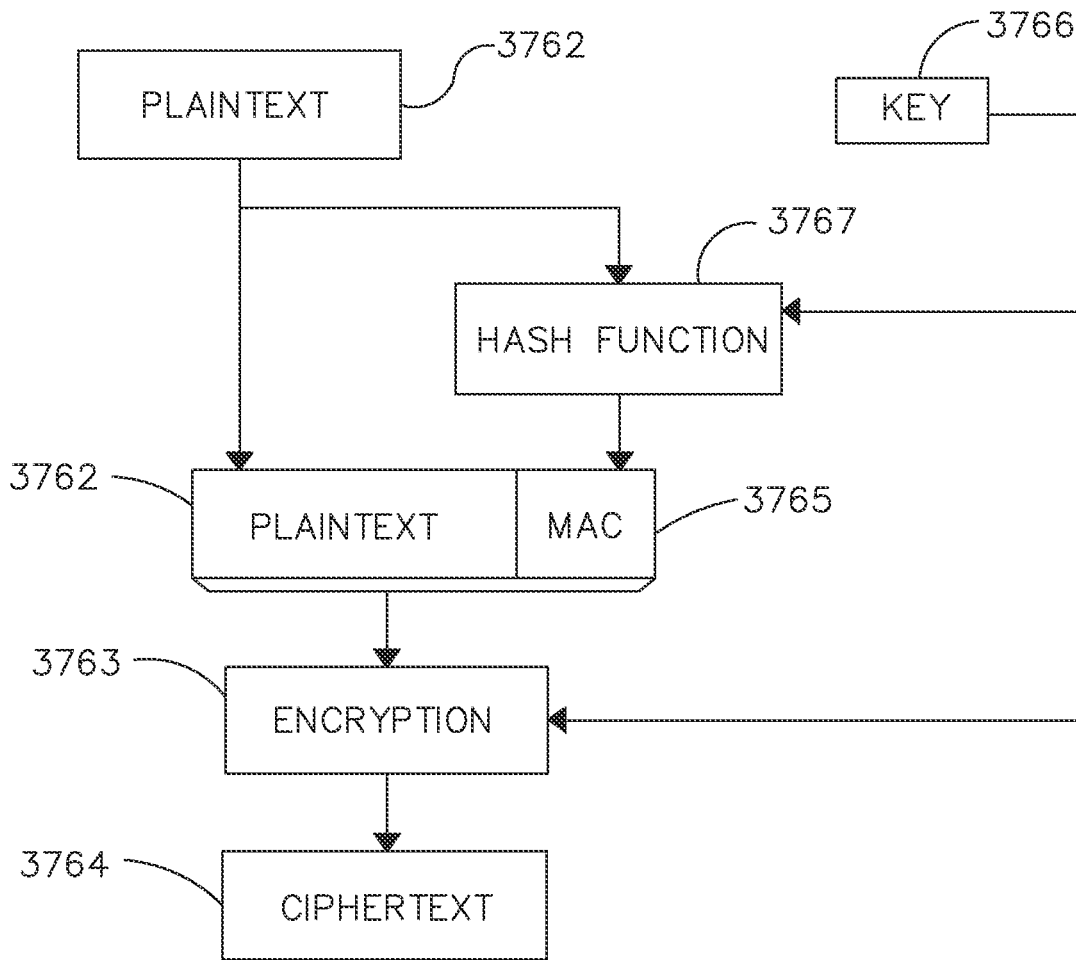


FIG. 27

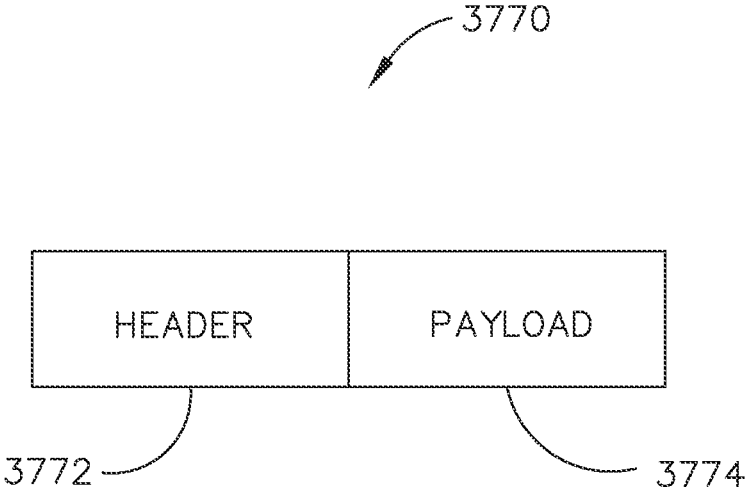


FIG. 28

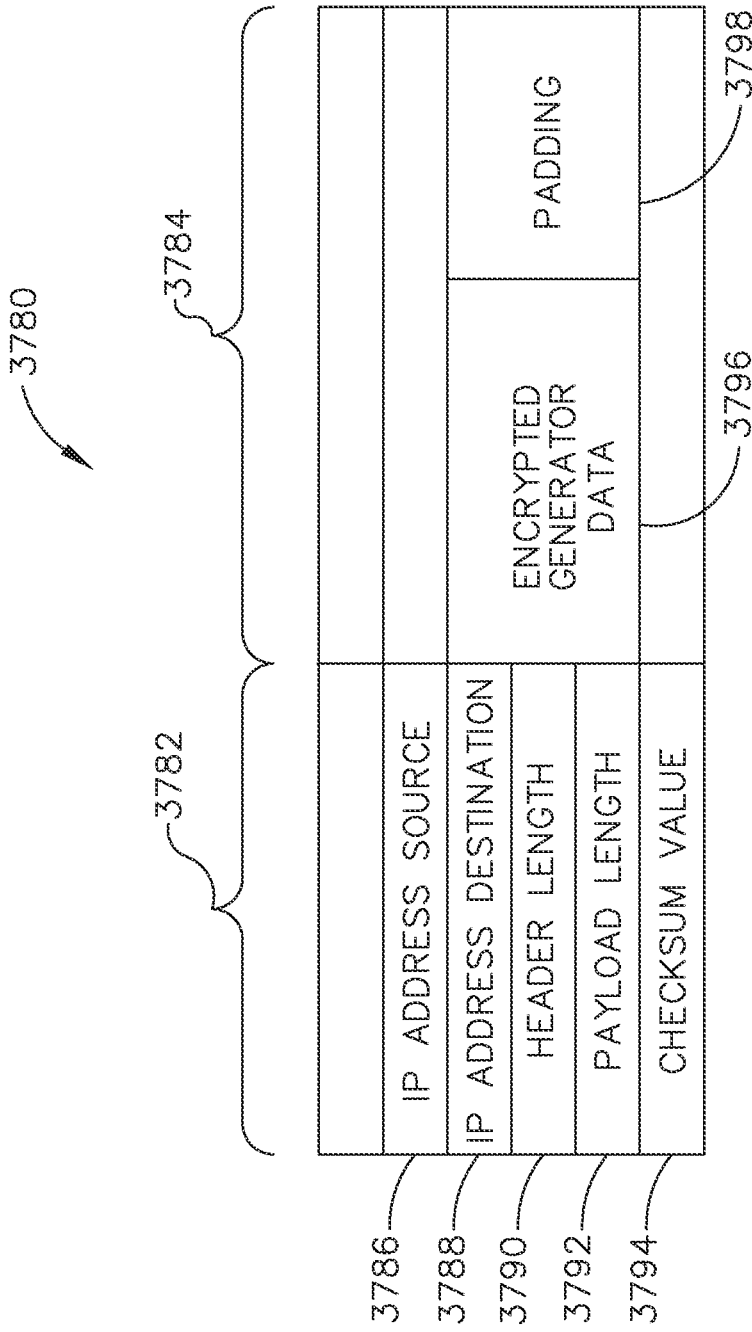


FIG. 29

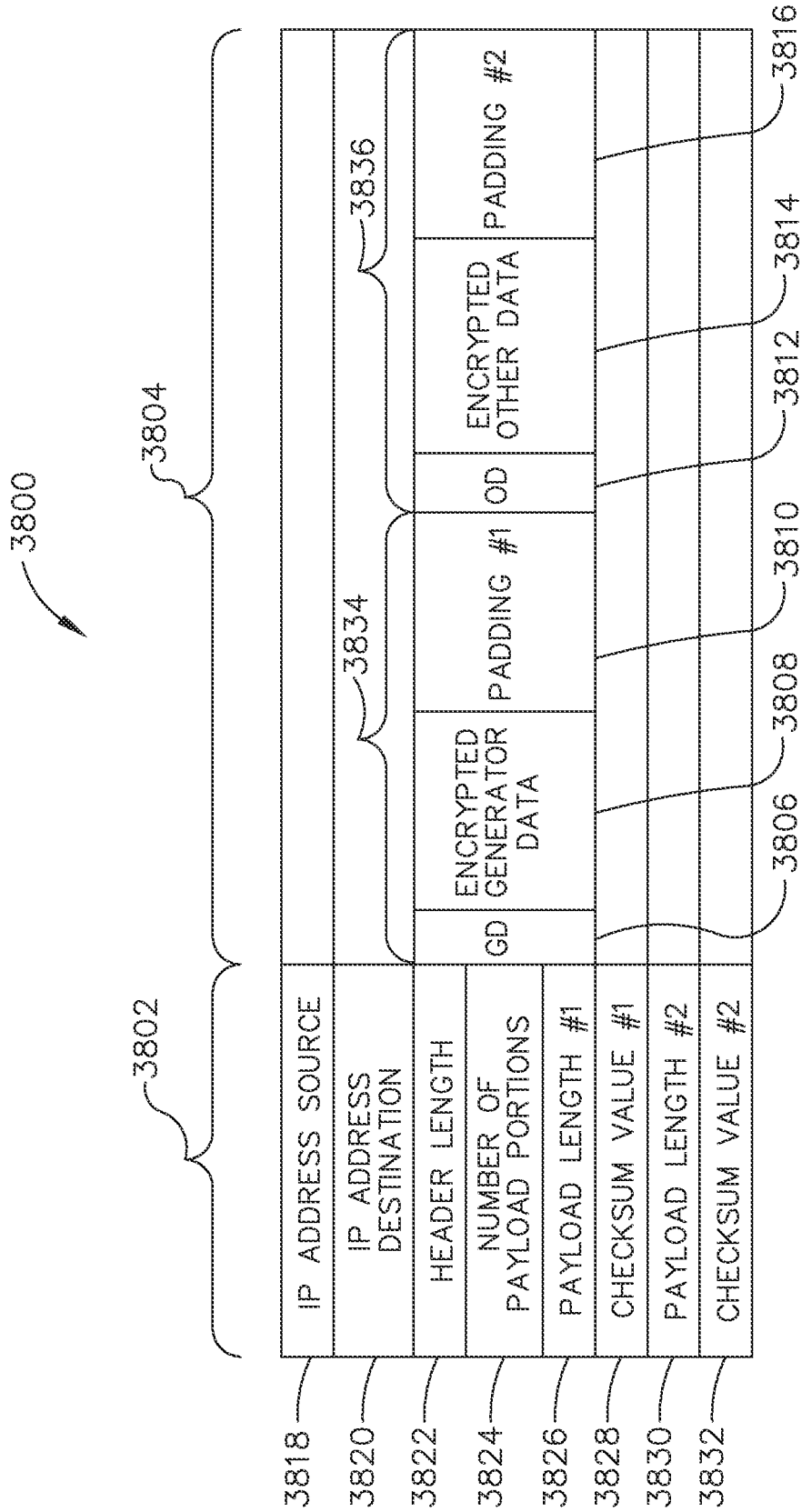


FIG. 30

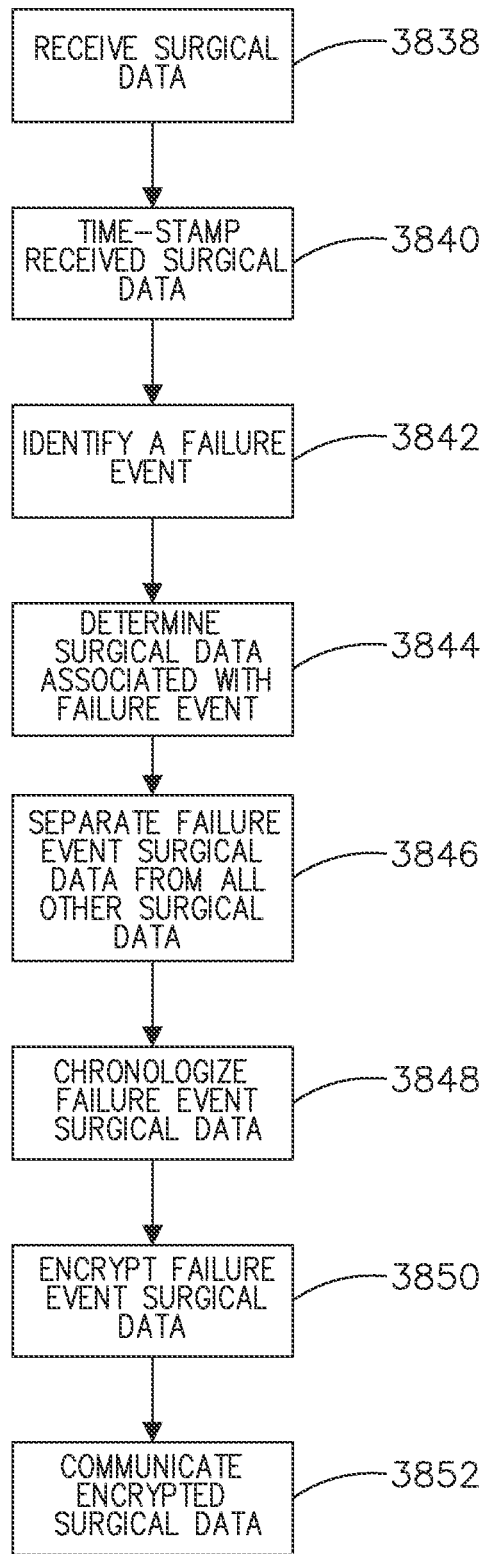


FIG. 31

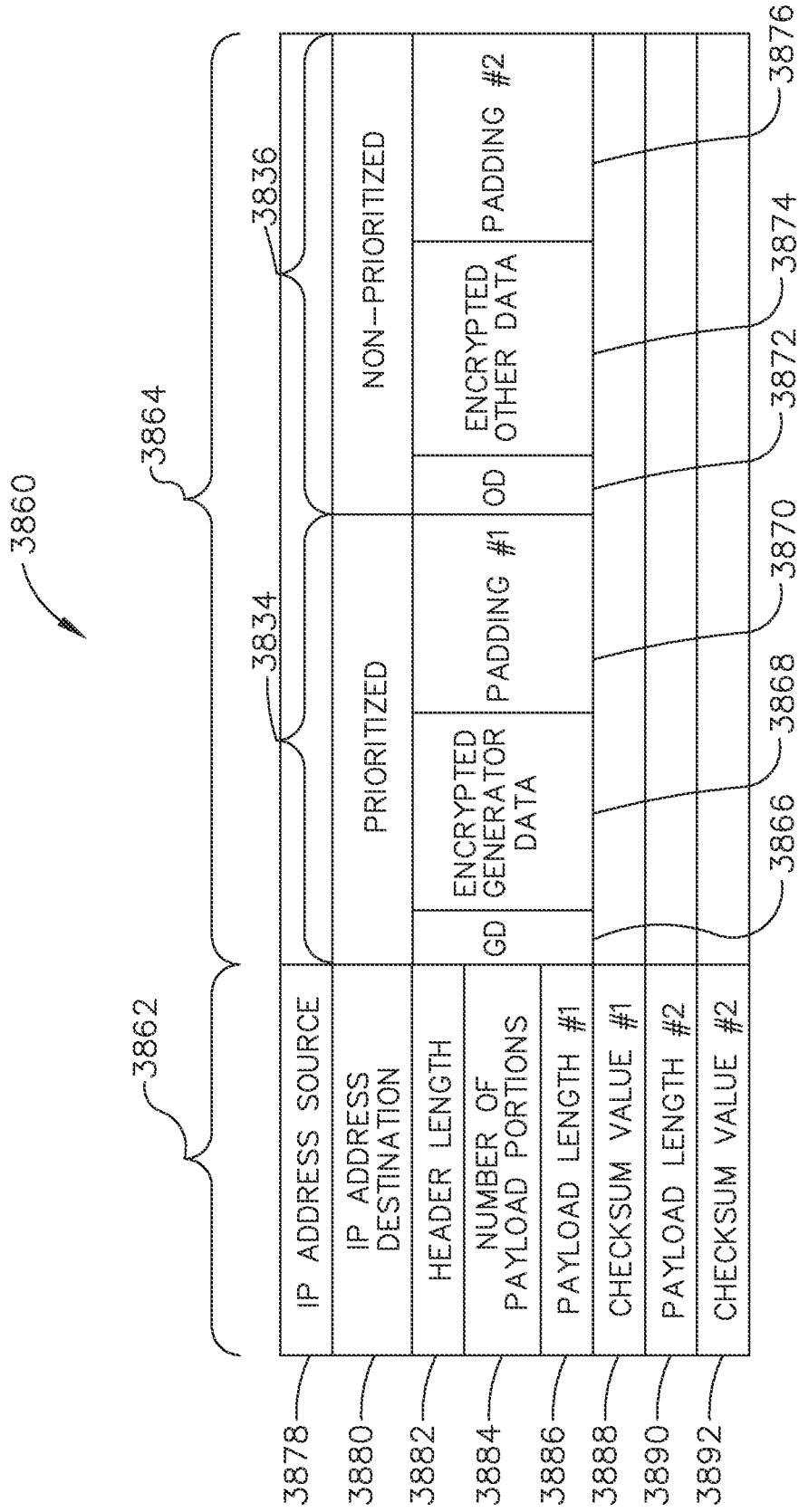
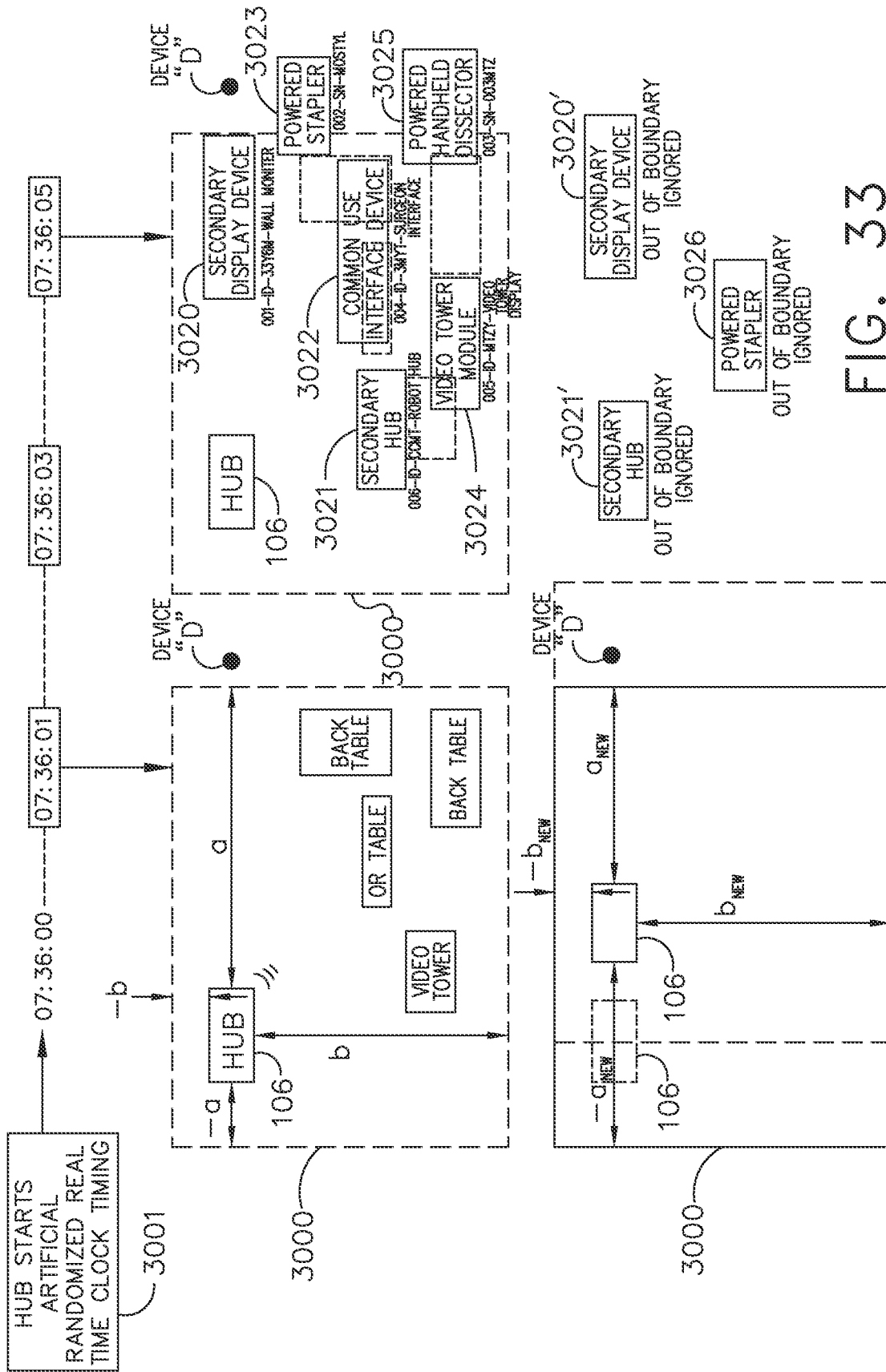


FIG. 32





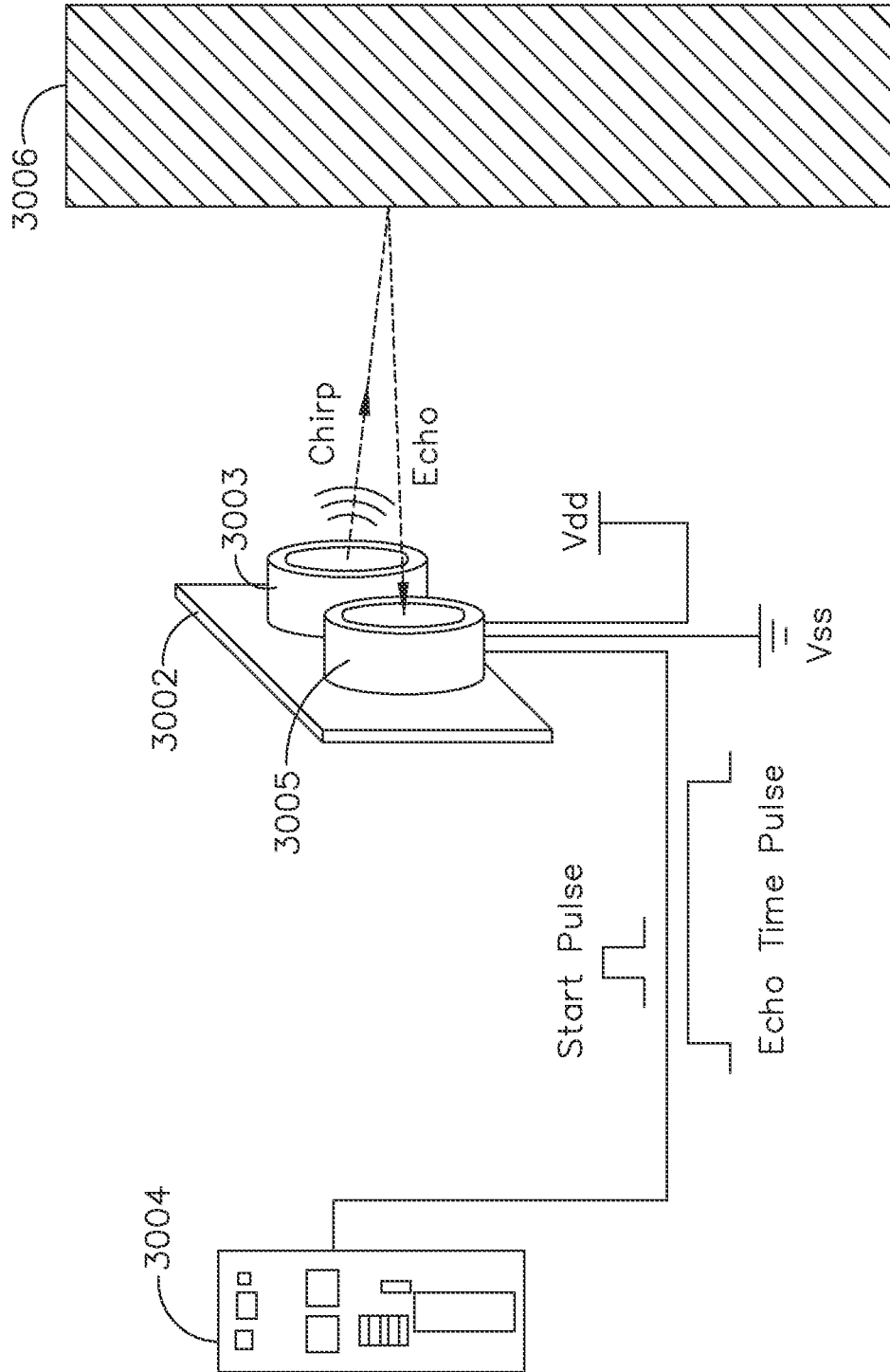


FIG. 34

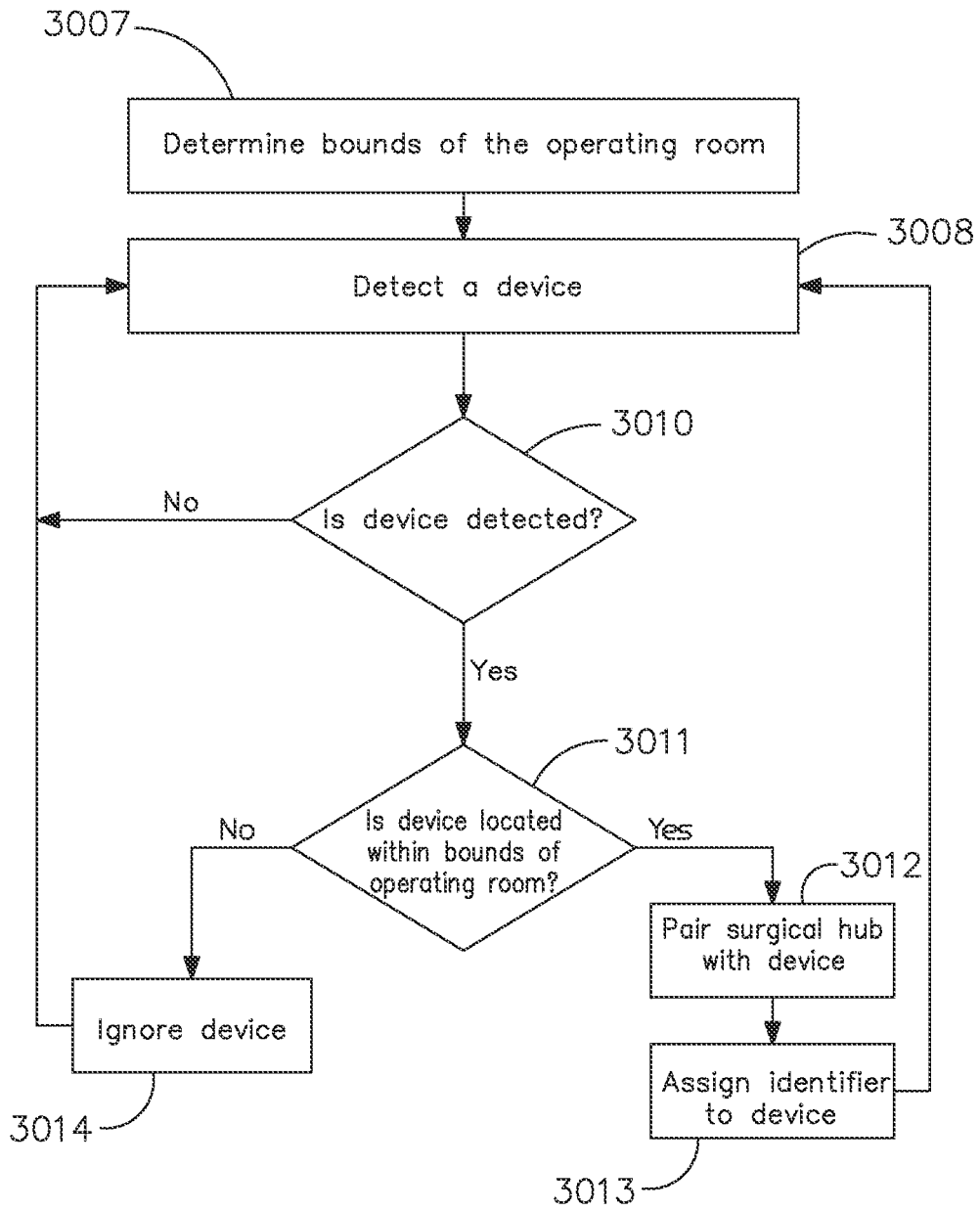


FIG. 35

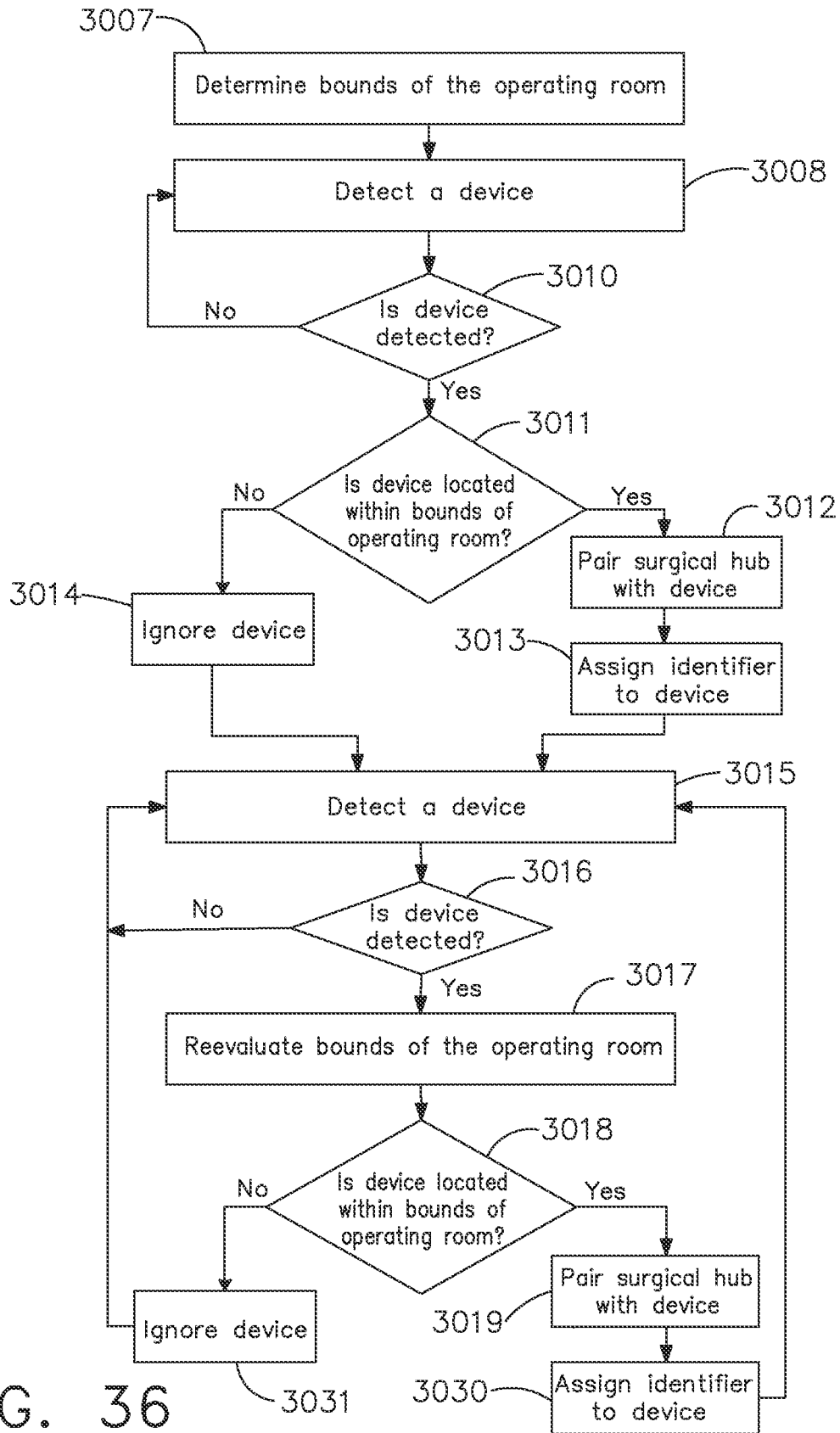


FIG. 36

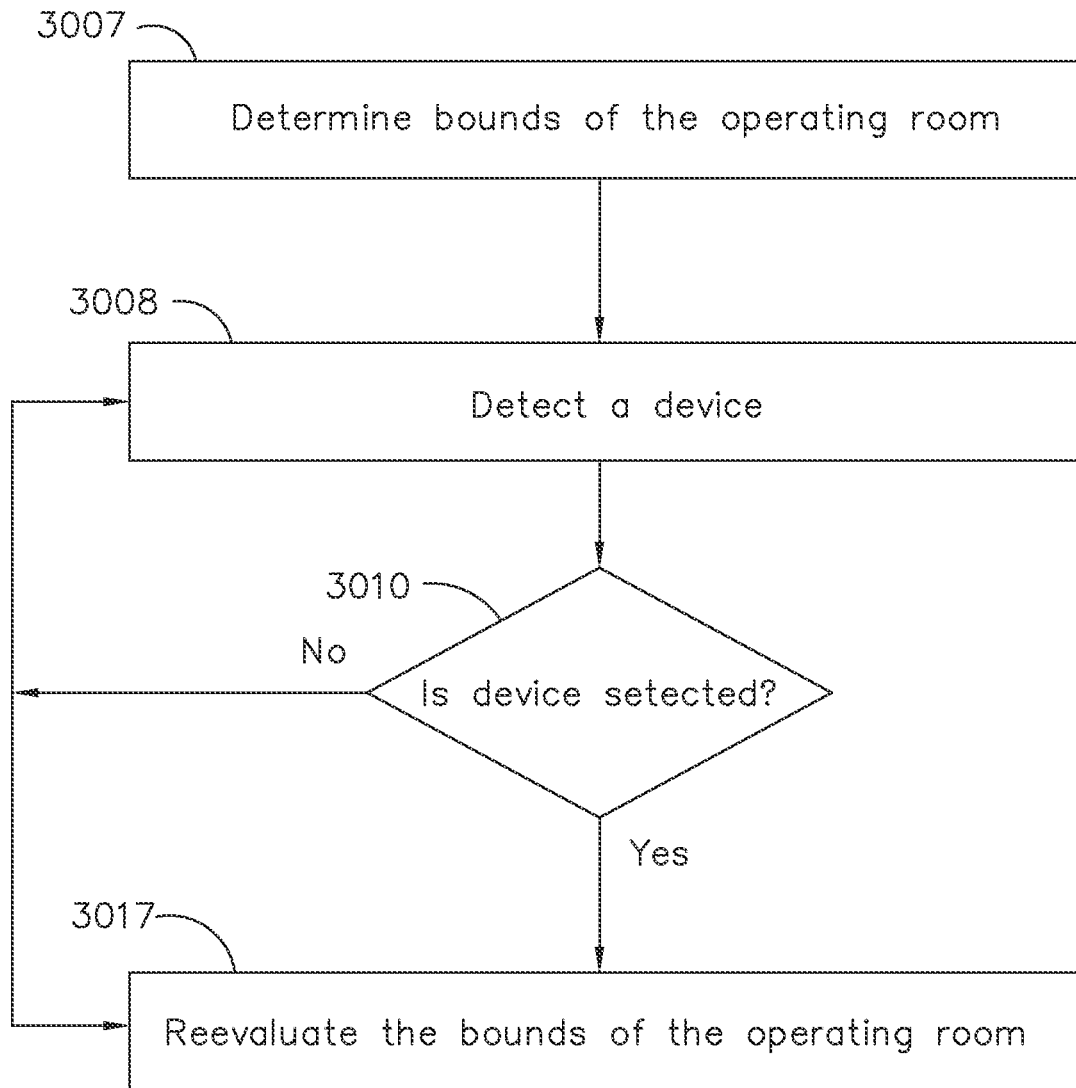


FIG. 37

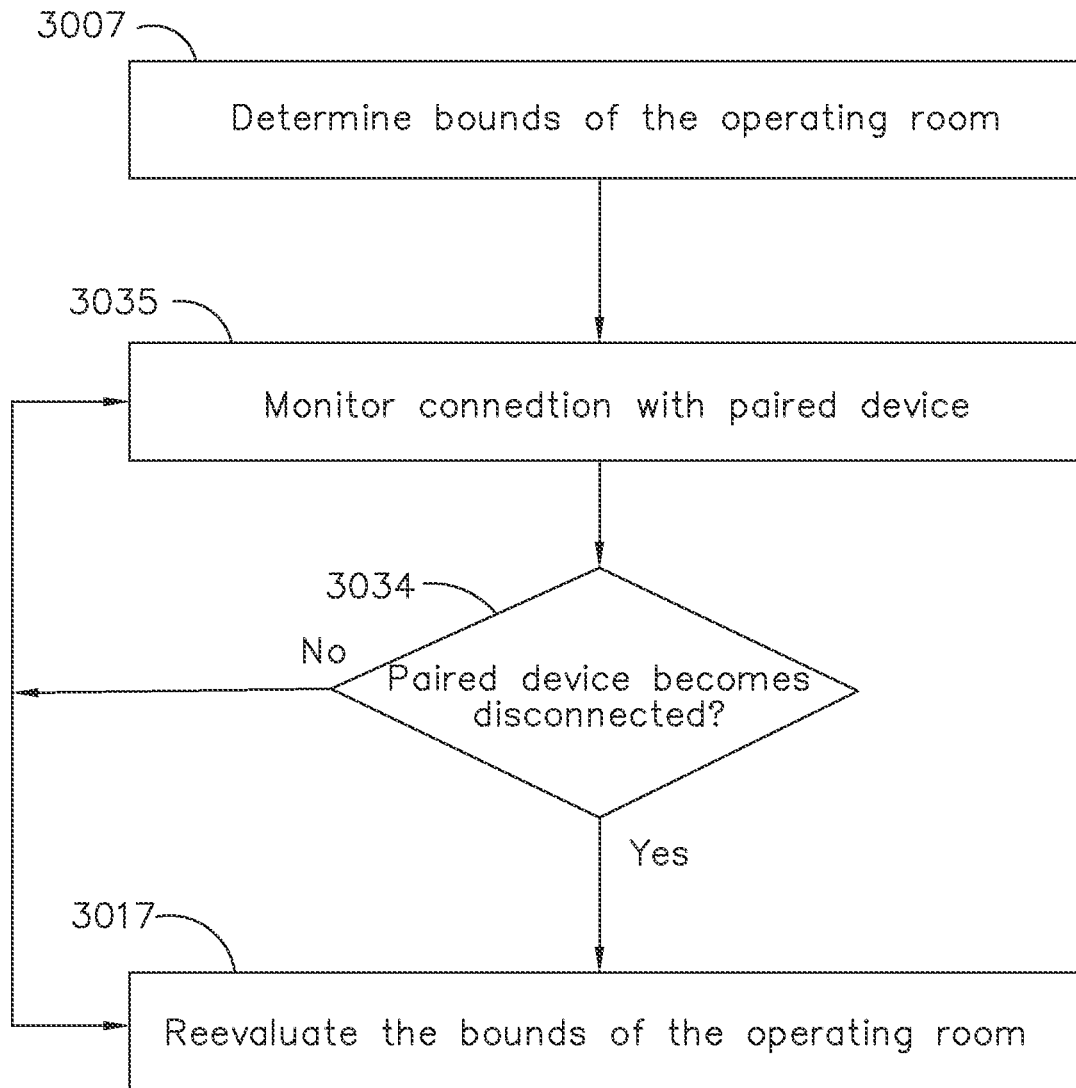


FIG. 38

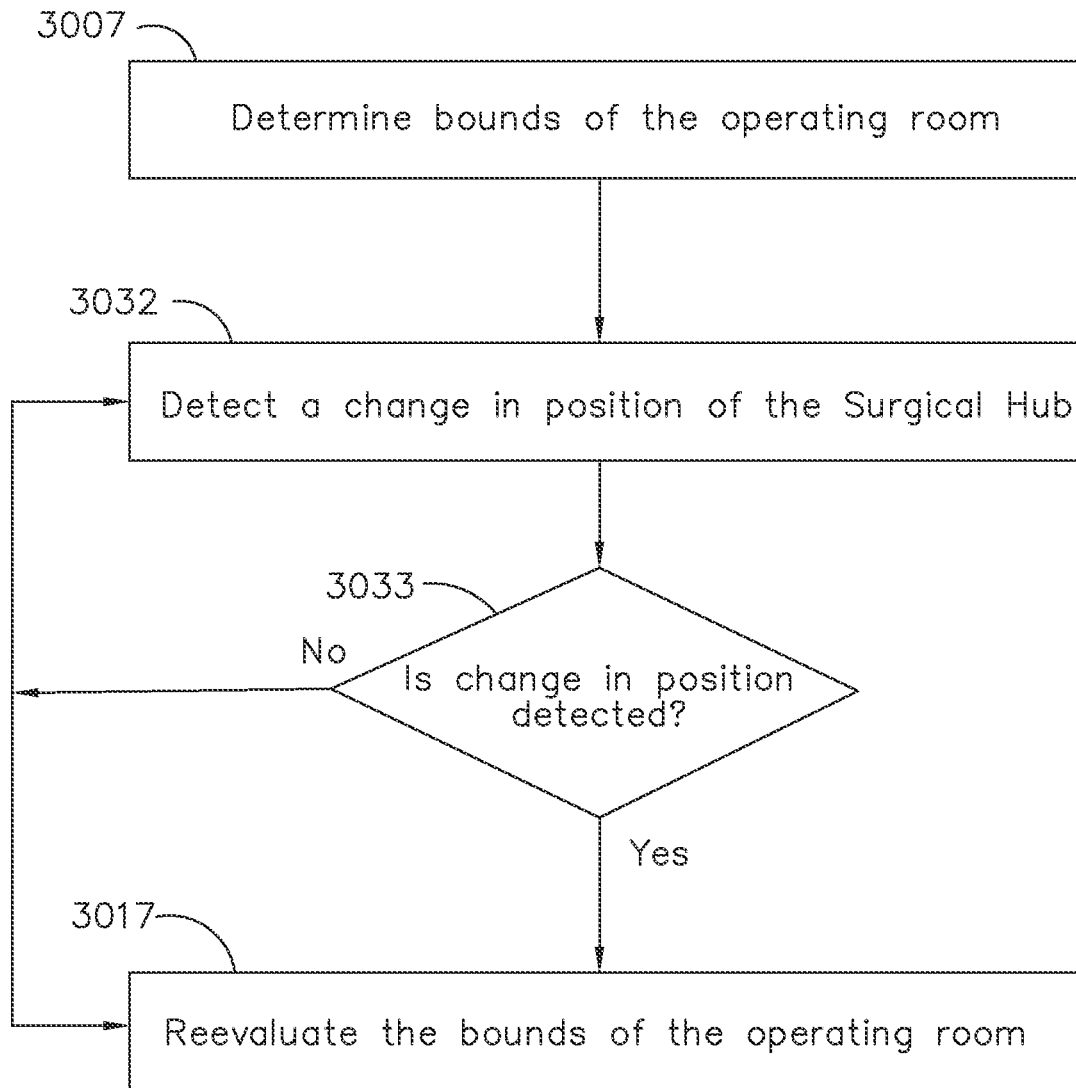


FIG. 39

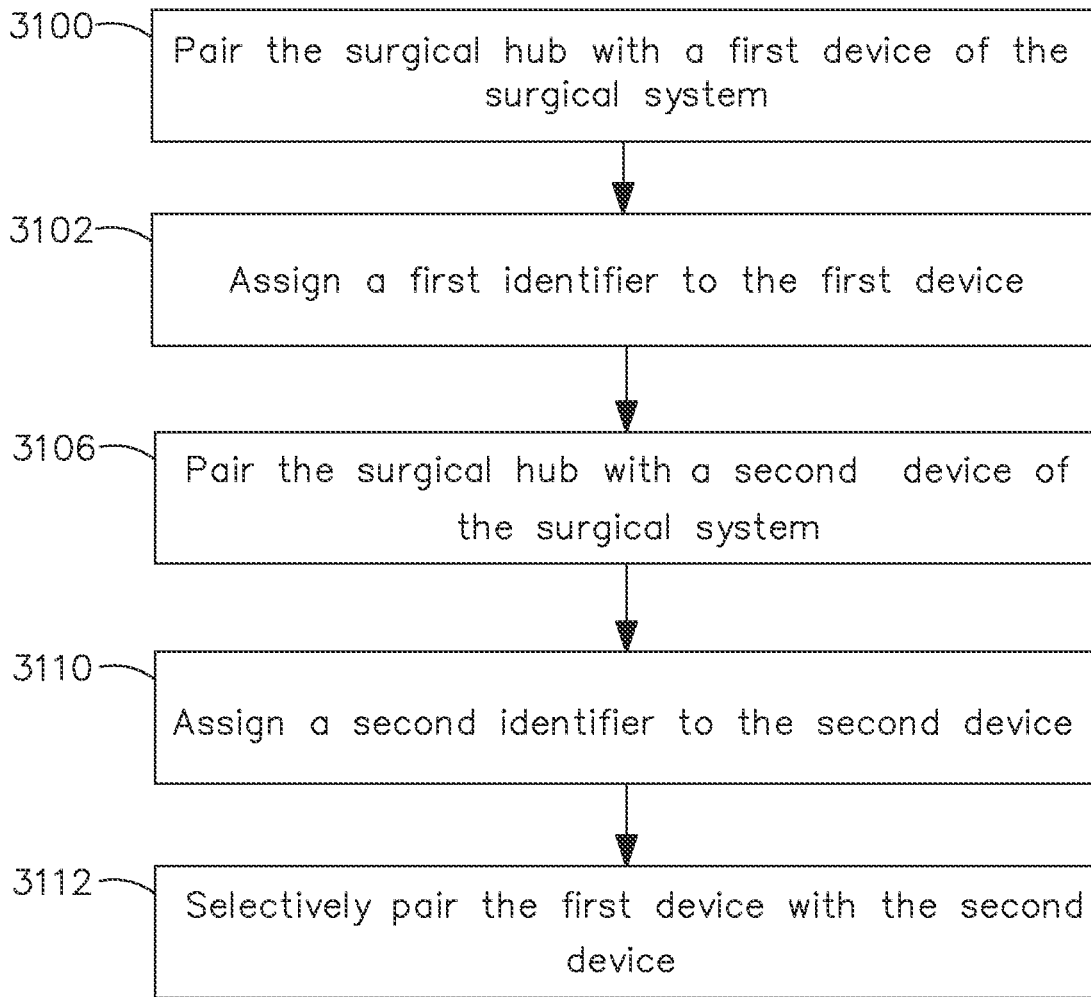


FIG. 40

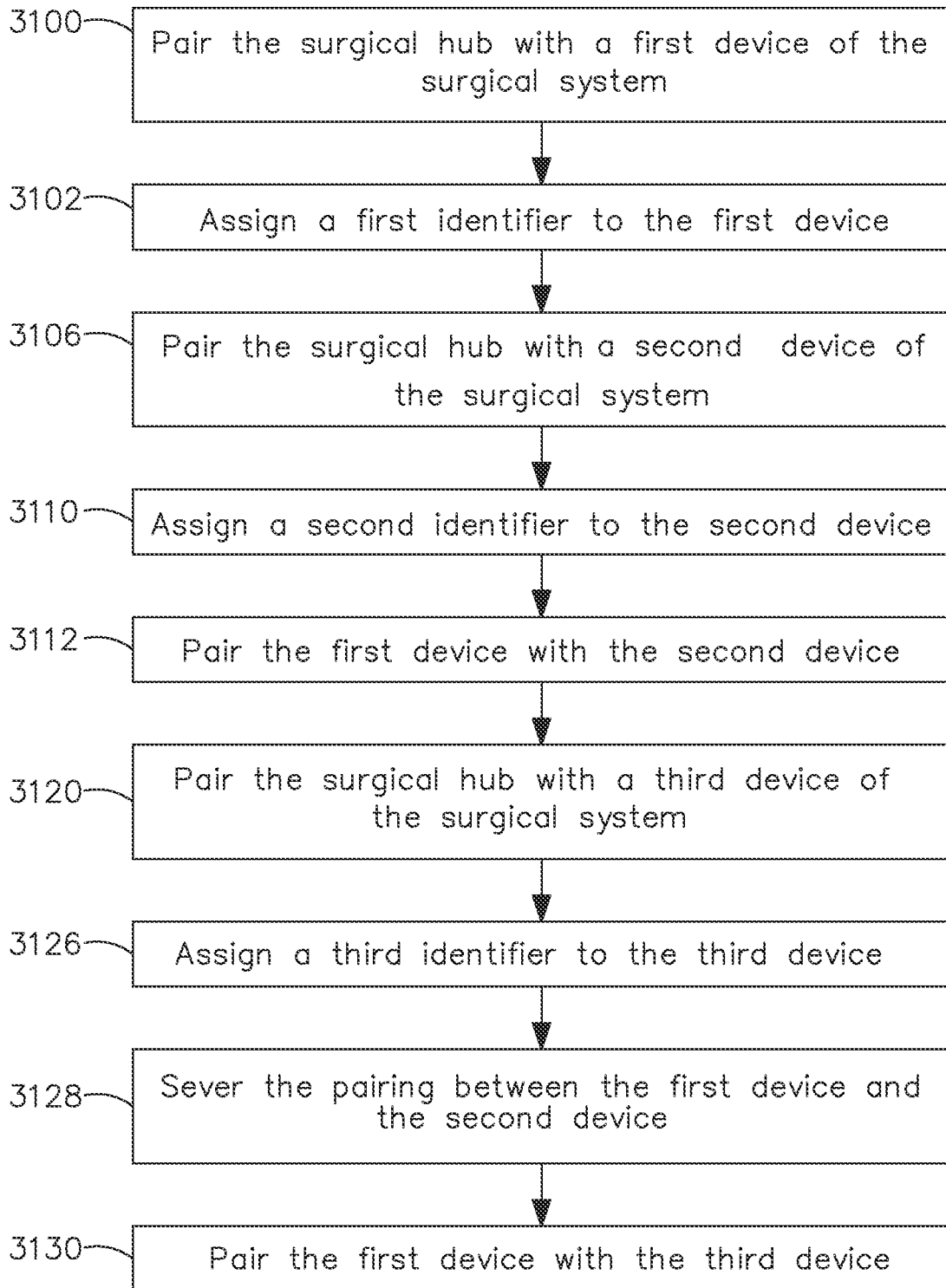


FIG. 41



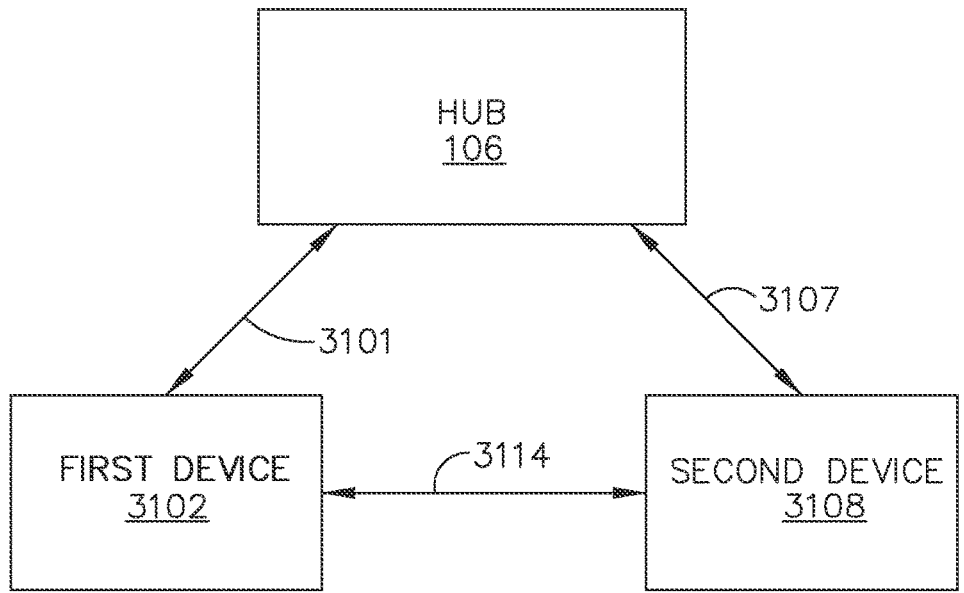


FIG. 42

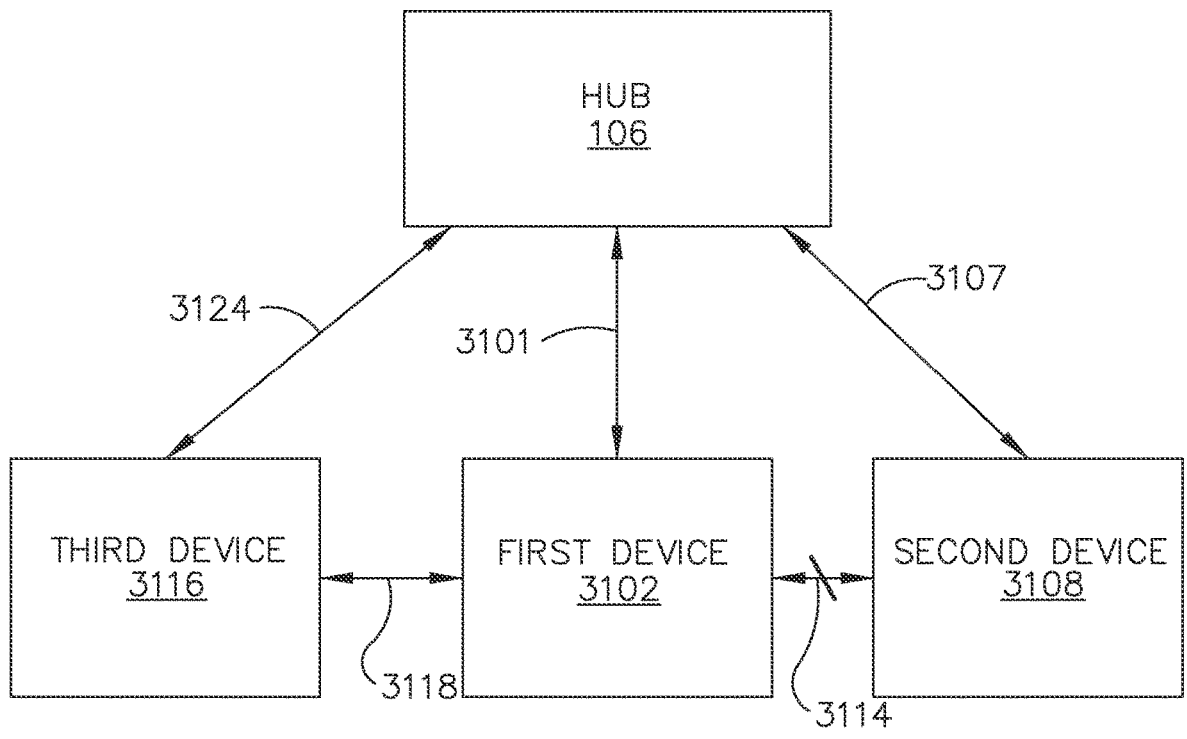


FIG. 43

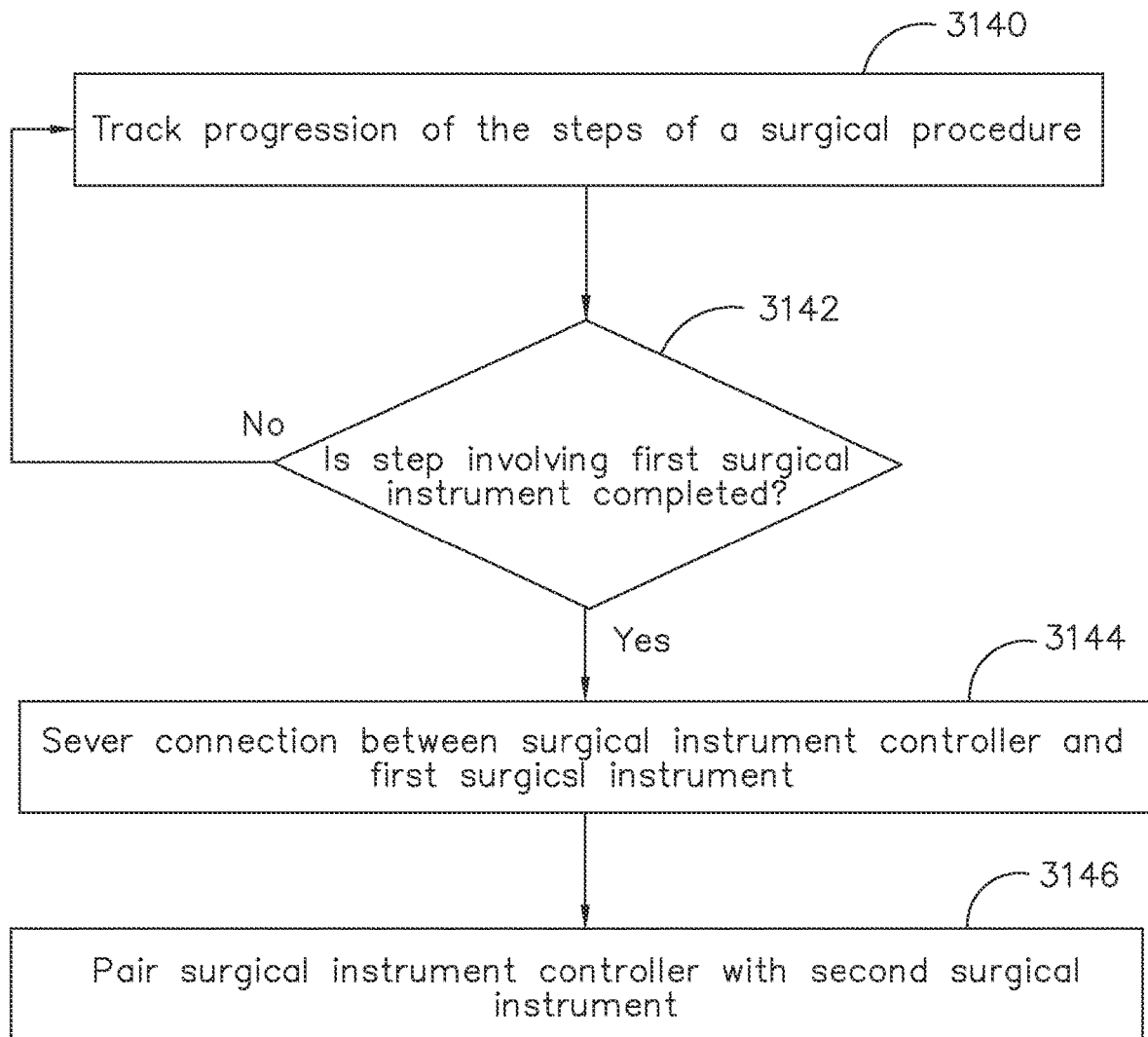


FIG. 44

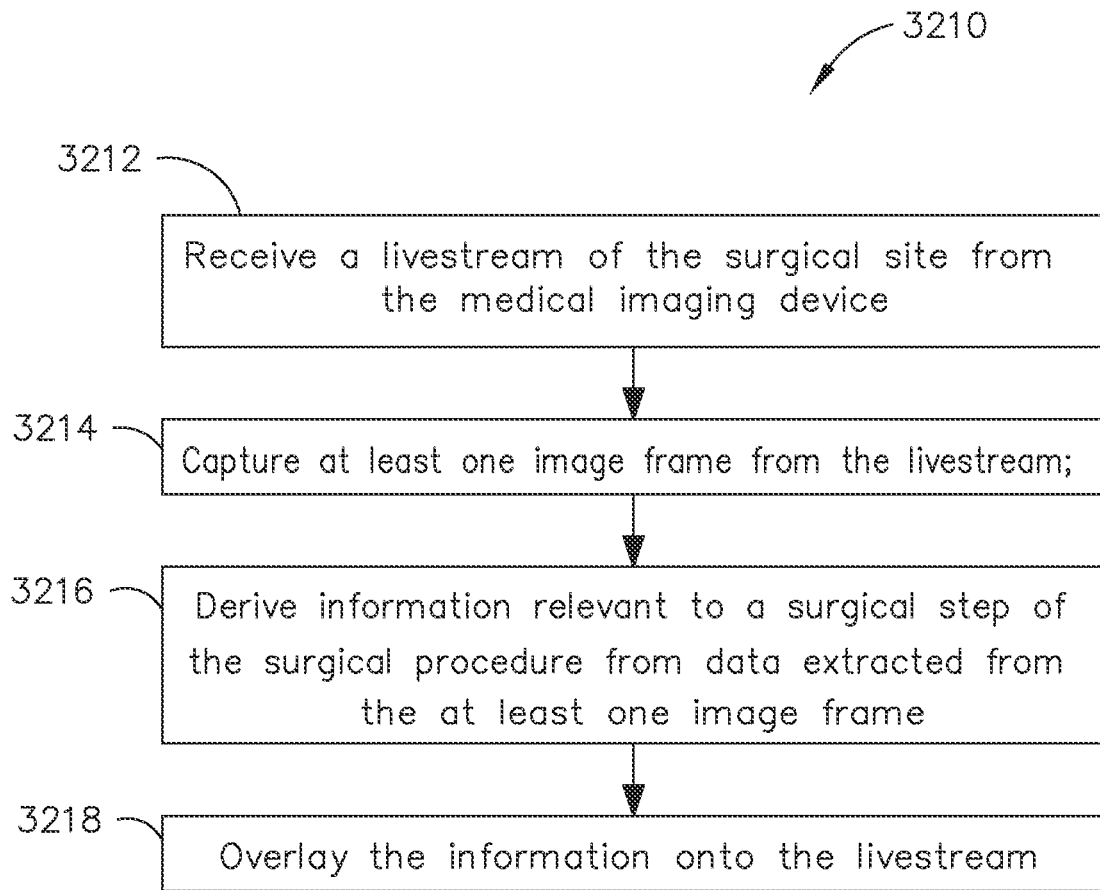


FIG. 45

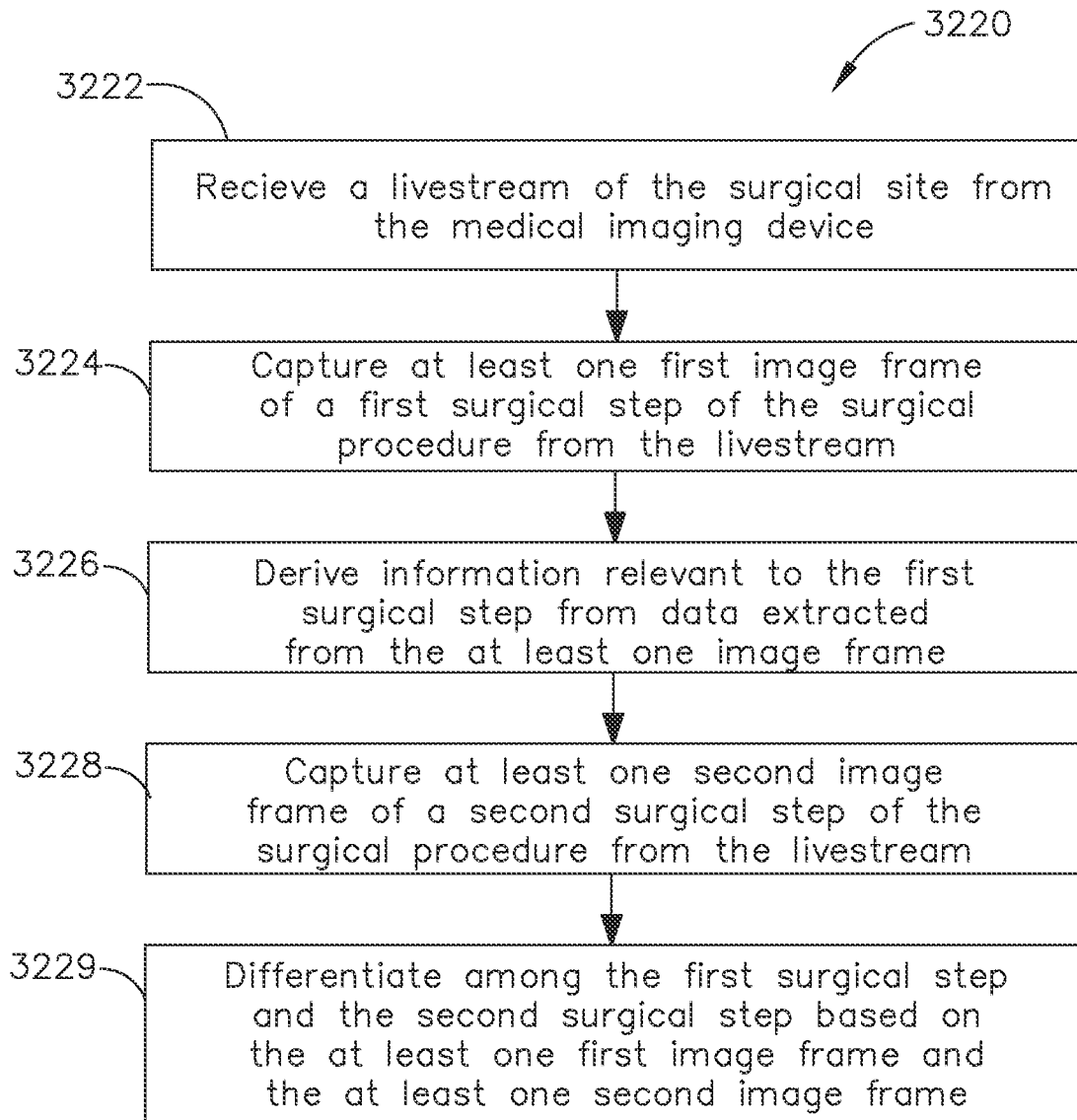


FIG. 46

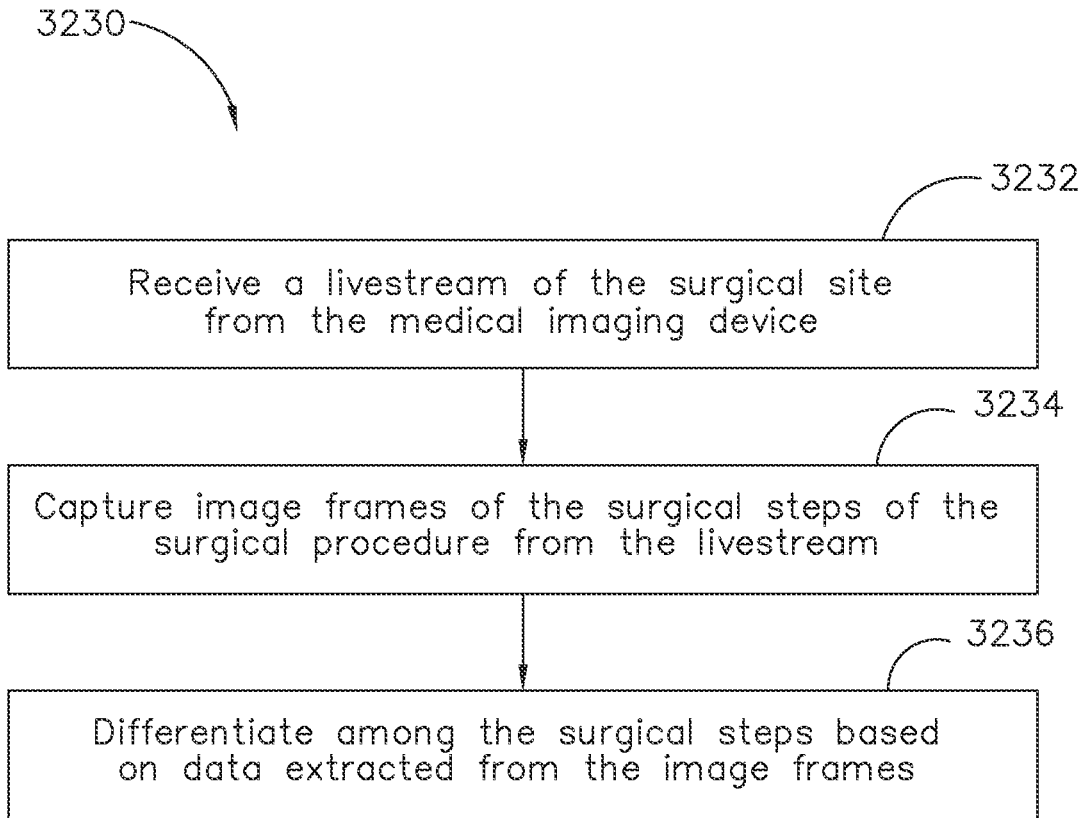


FIG. 47

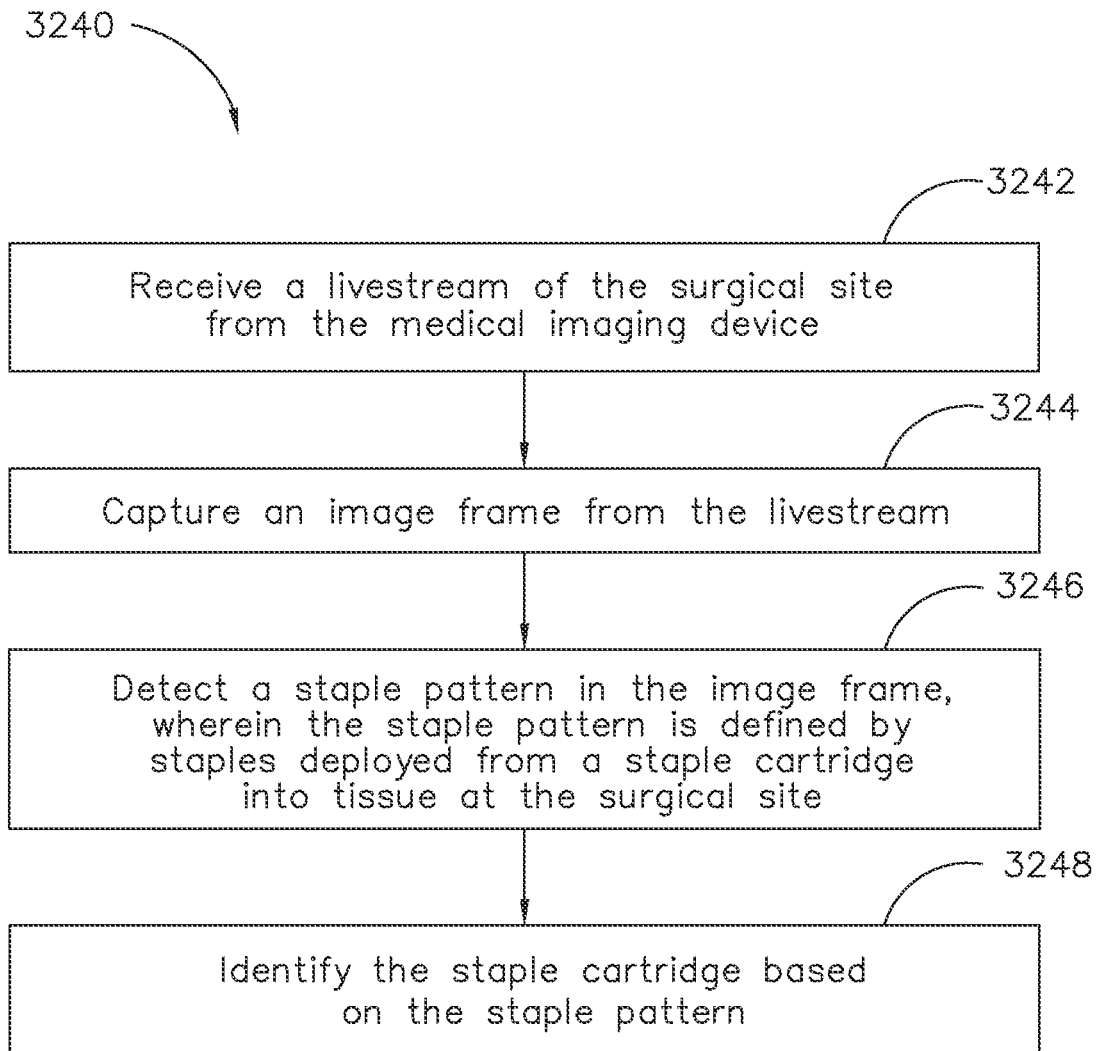


FIG. 48

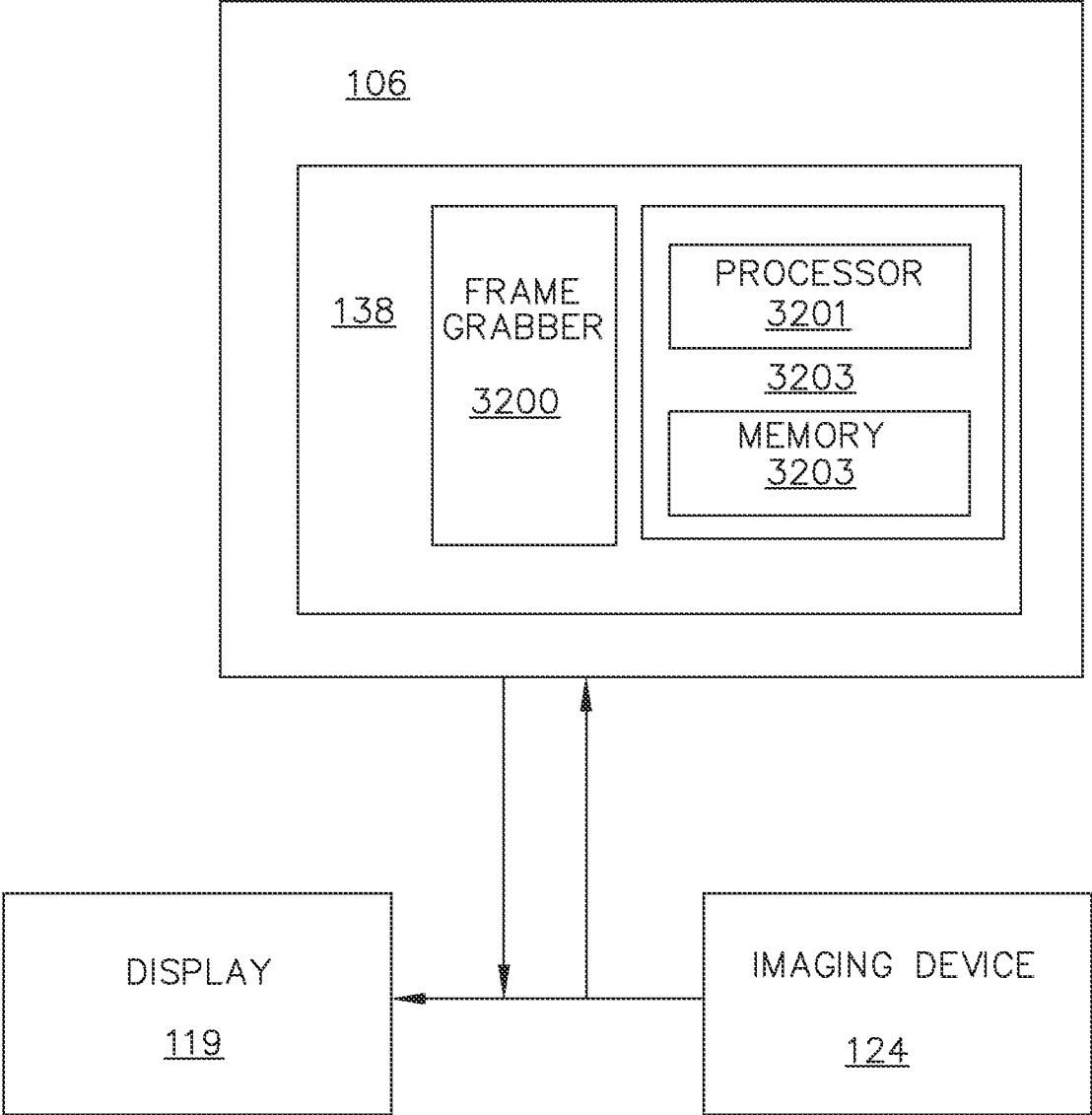


FIG. 49

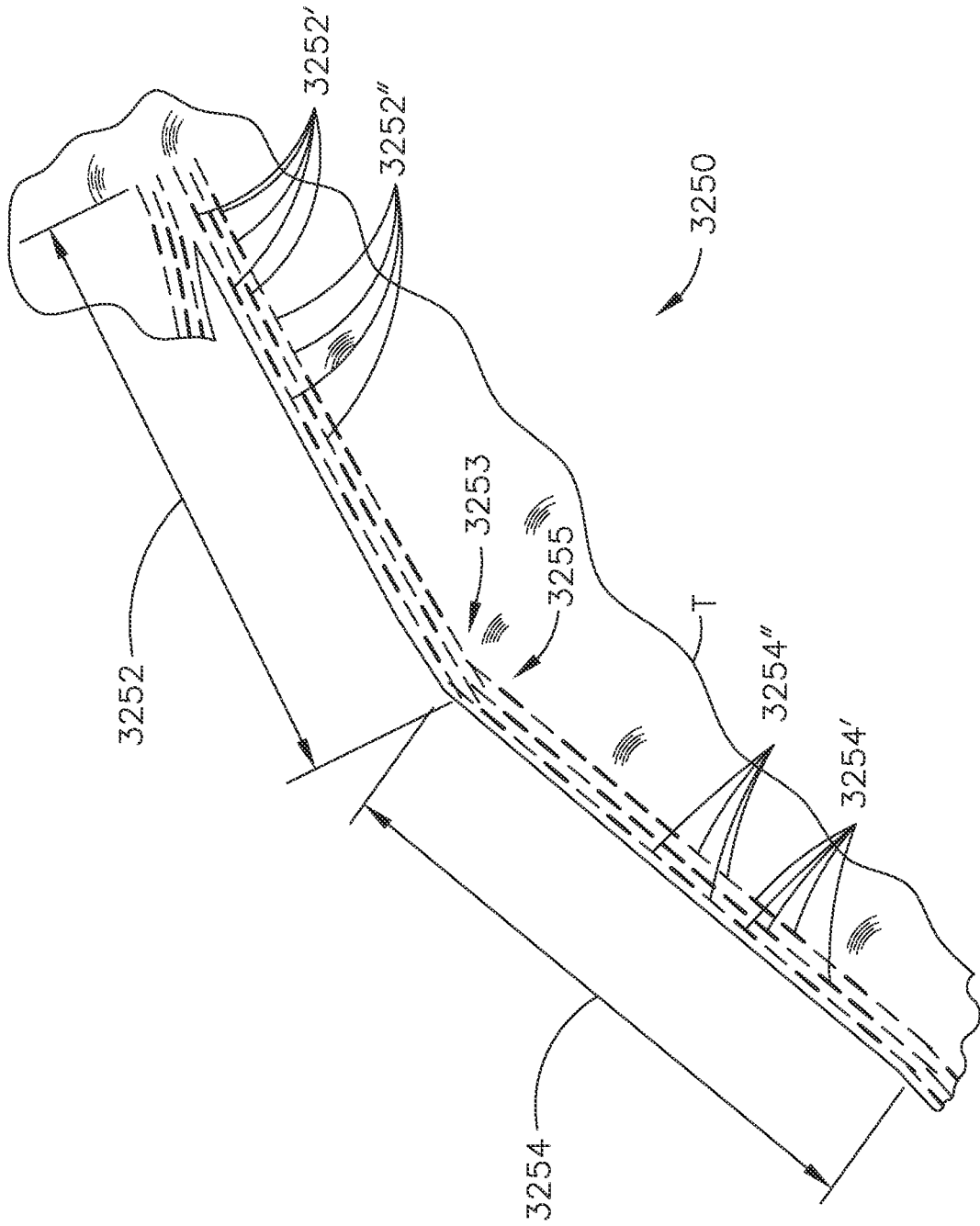


FIG. 50



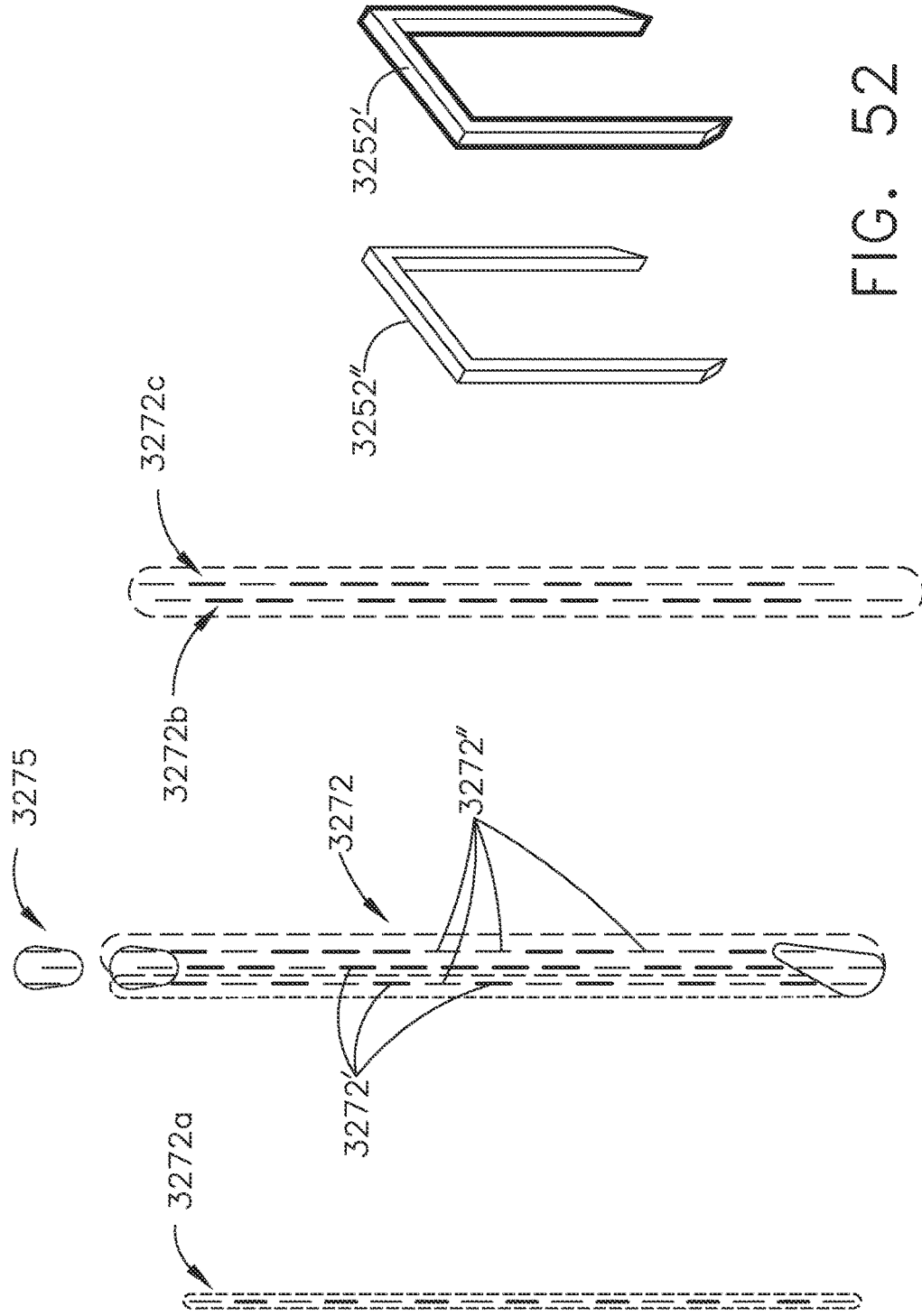


FIG. 51

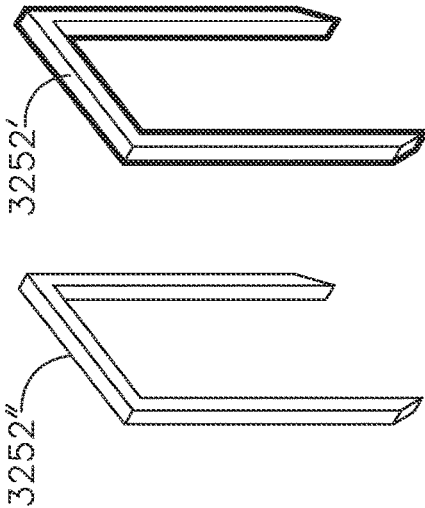


FIG. 52

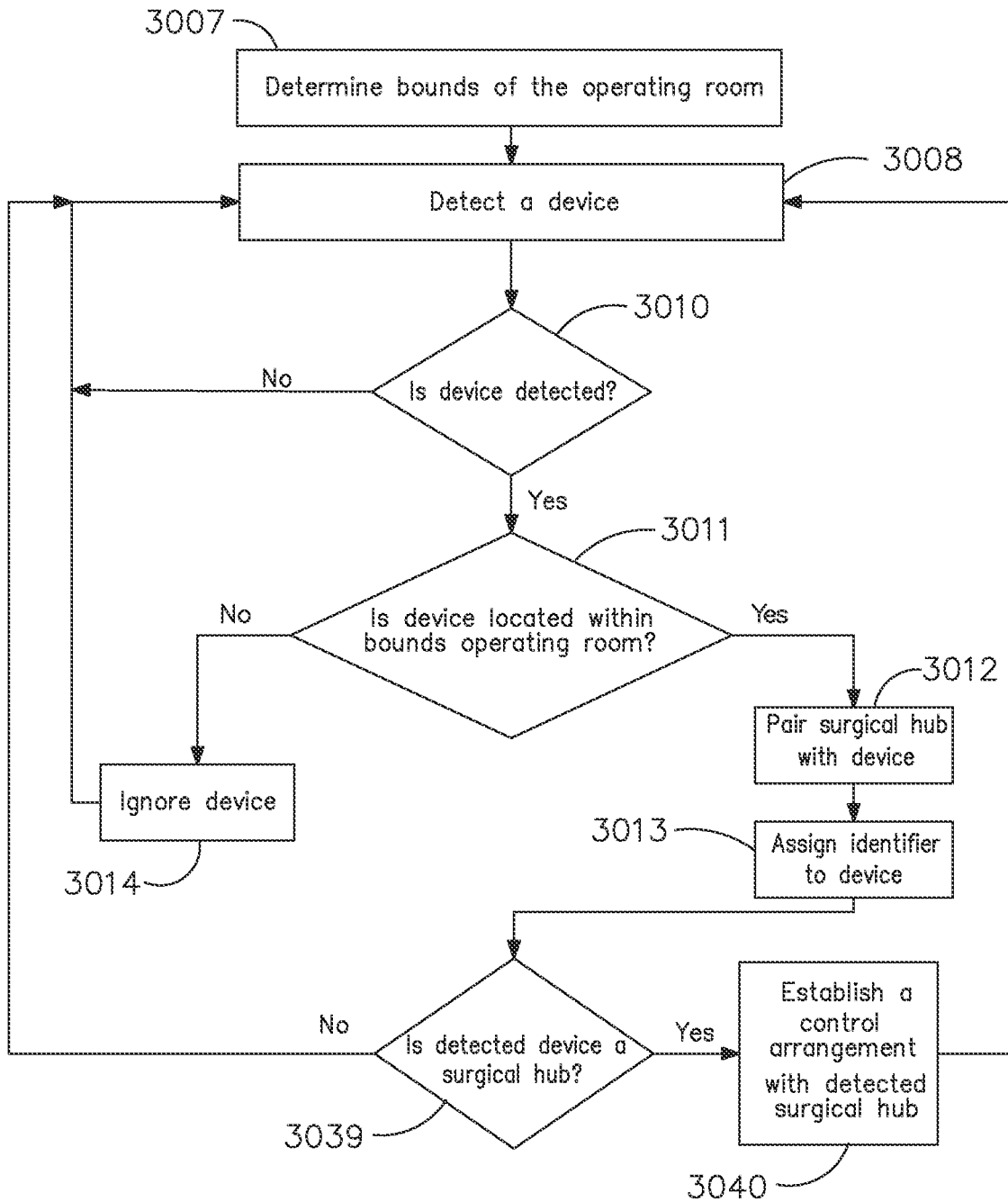


FIG. 53

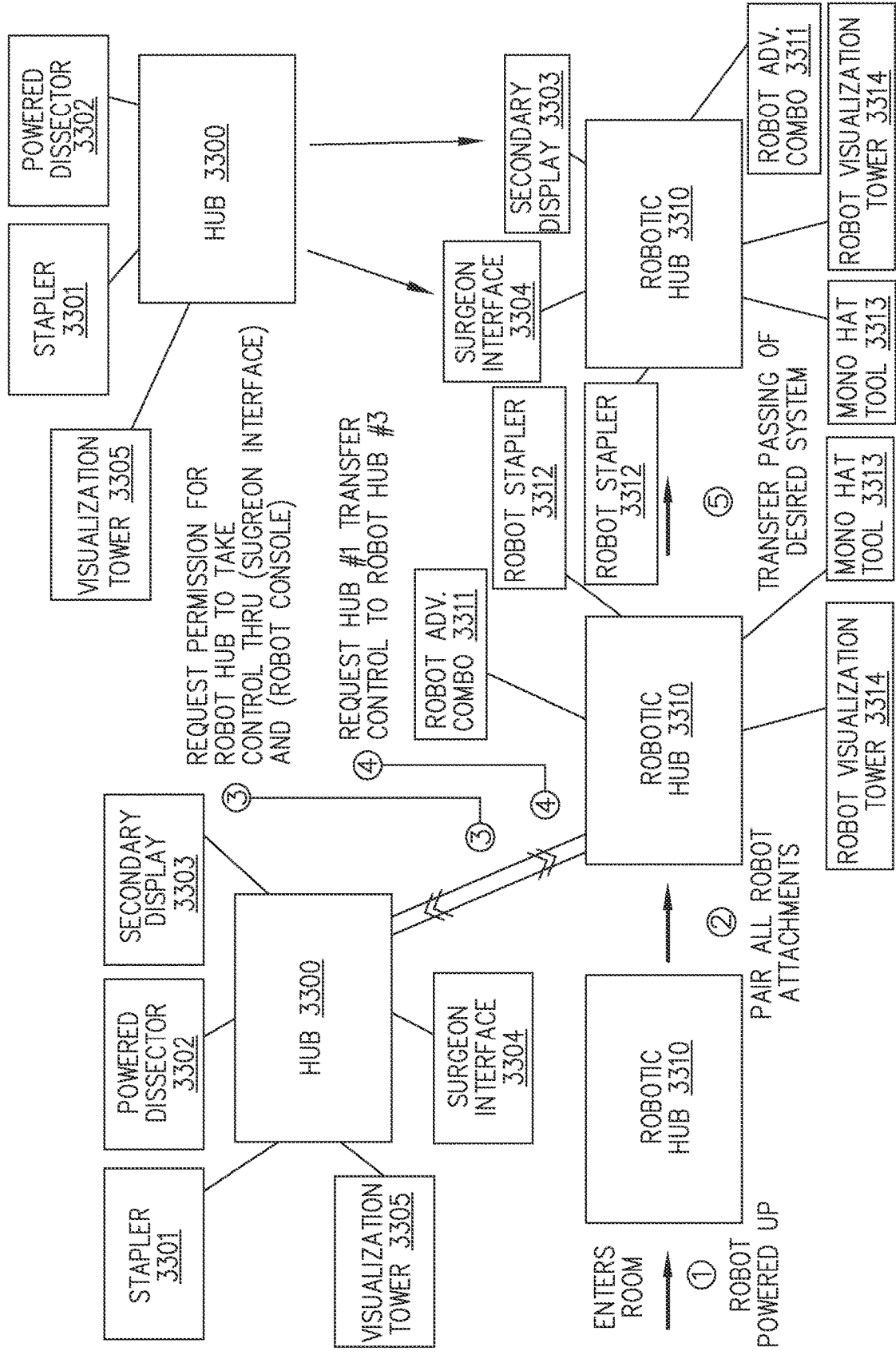


FIG. 54

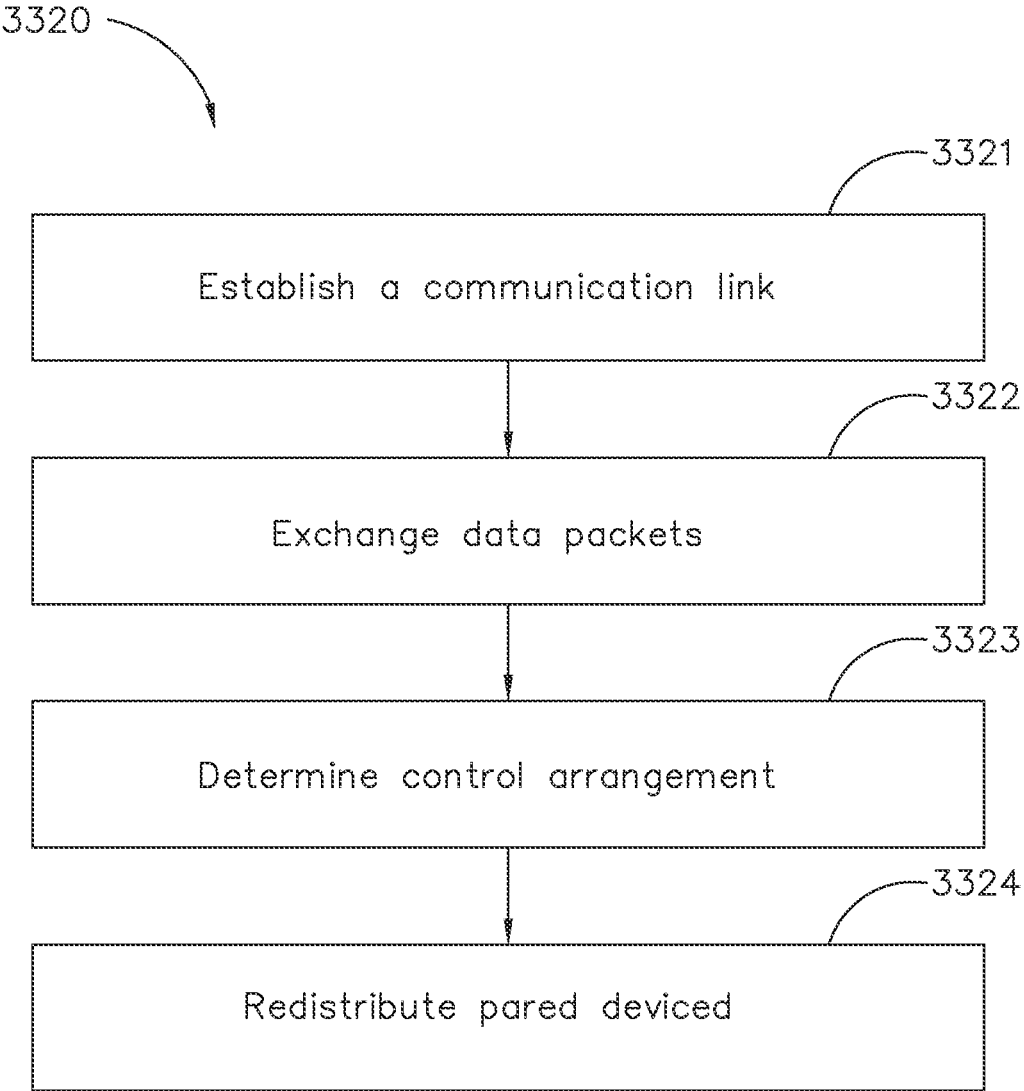
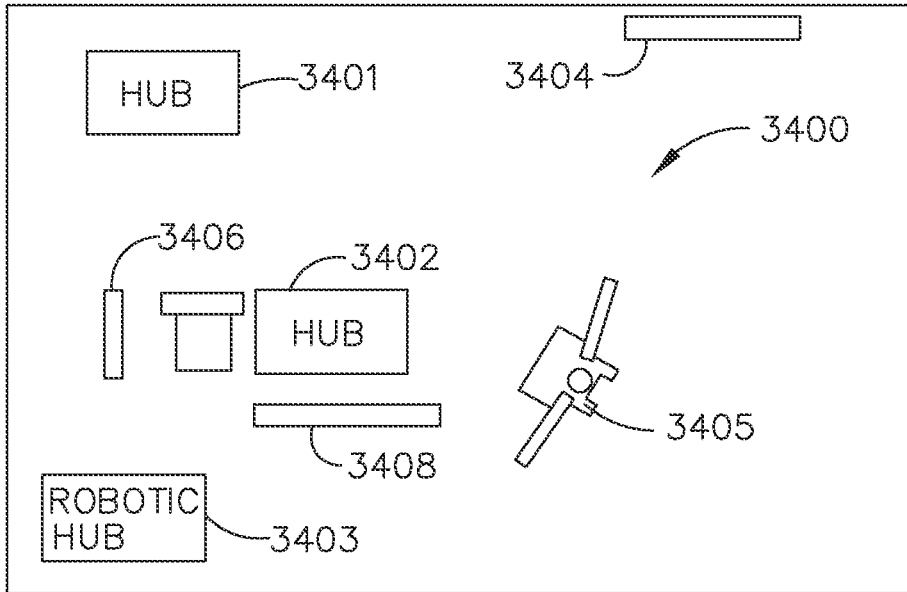
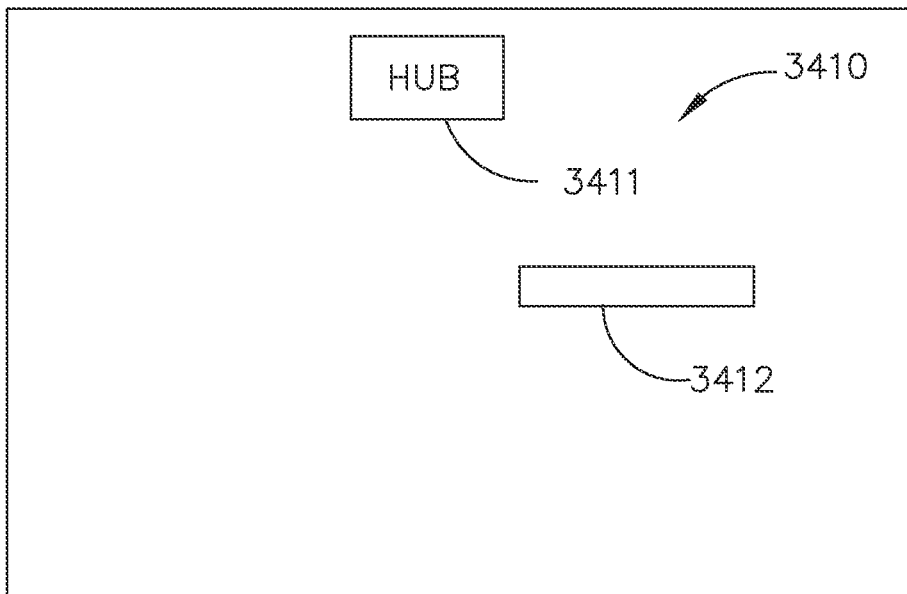


FIG. 55



OR 1 THORACIC SEGMENTECTOMY



OR 3 COLORECTAL

FIG. 56

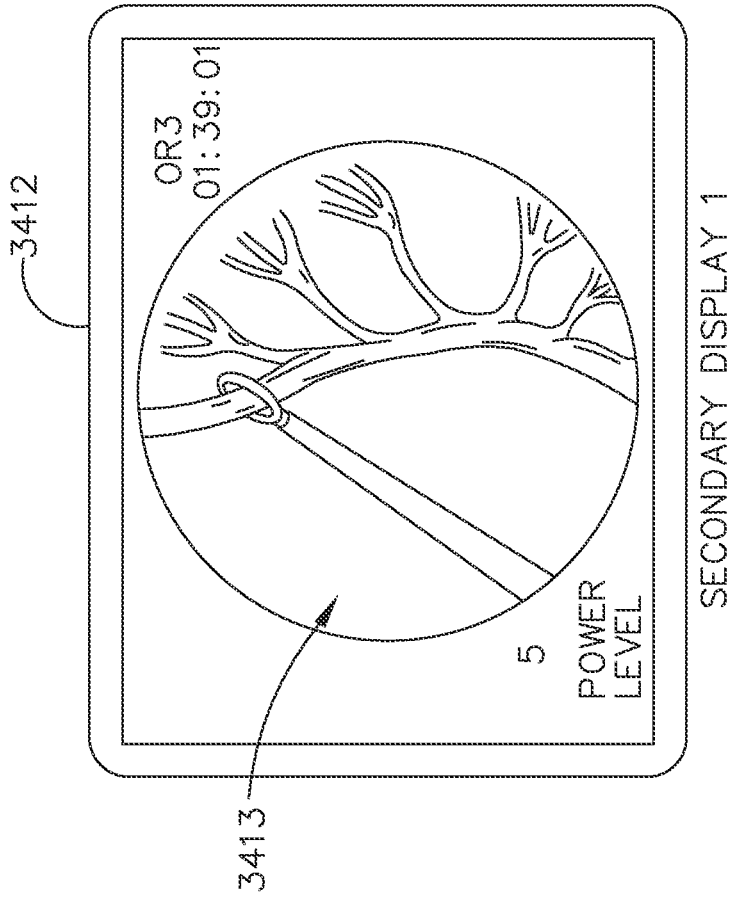


FIG. 57

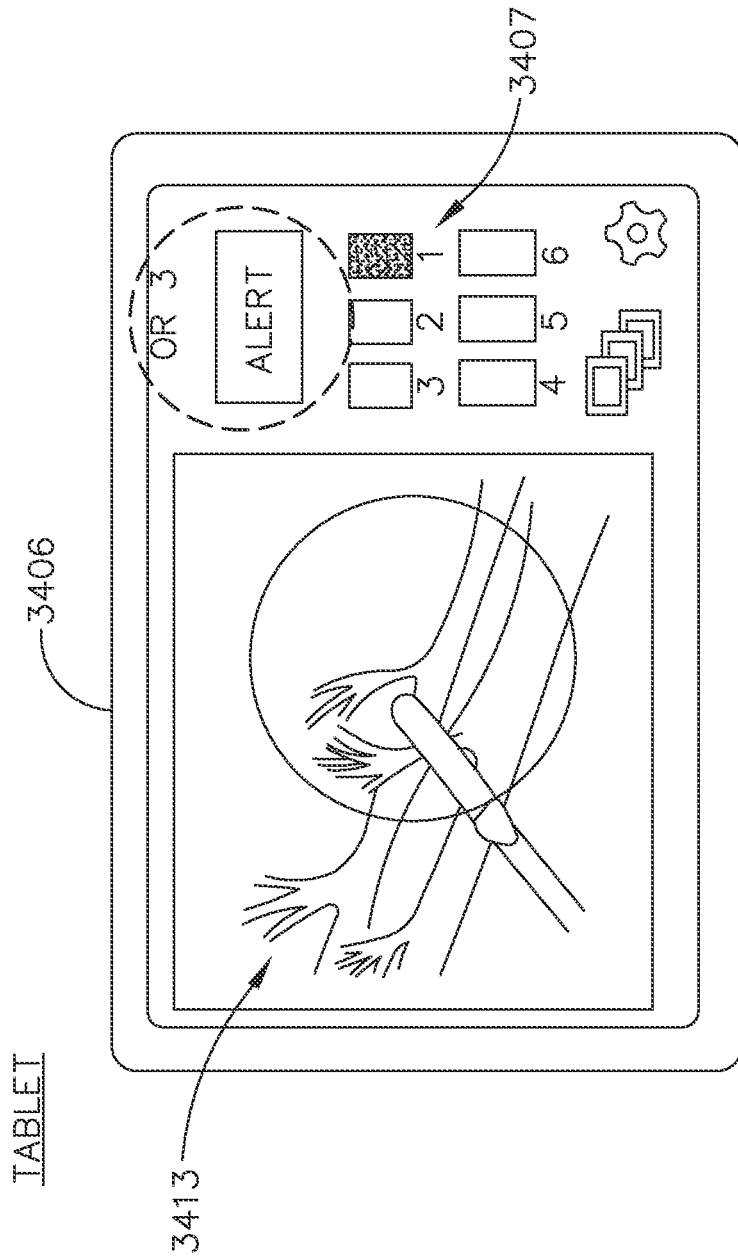


FIG. 58

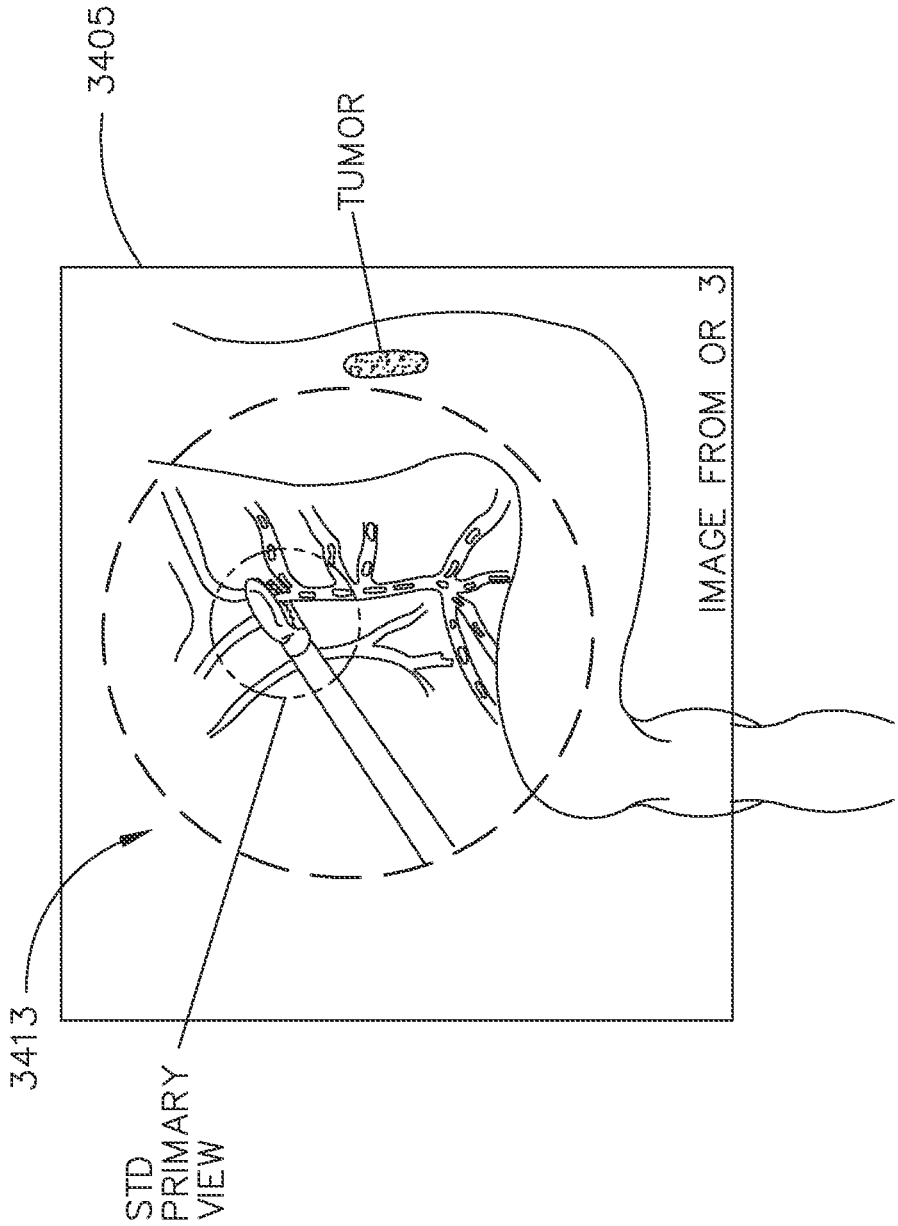


FIG. 59



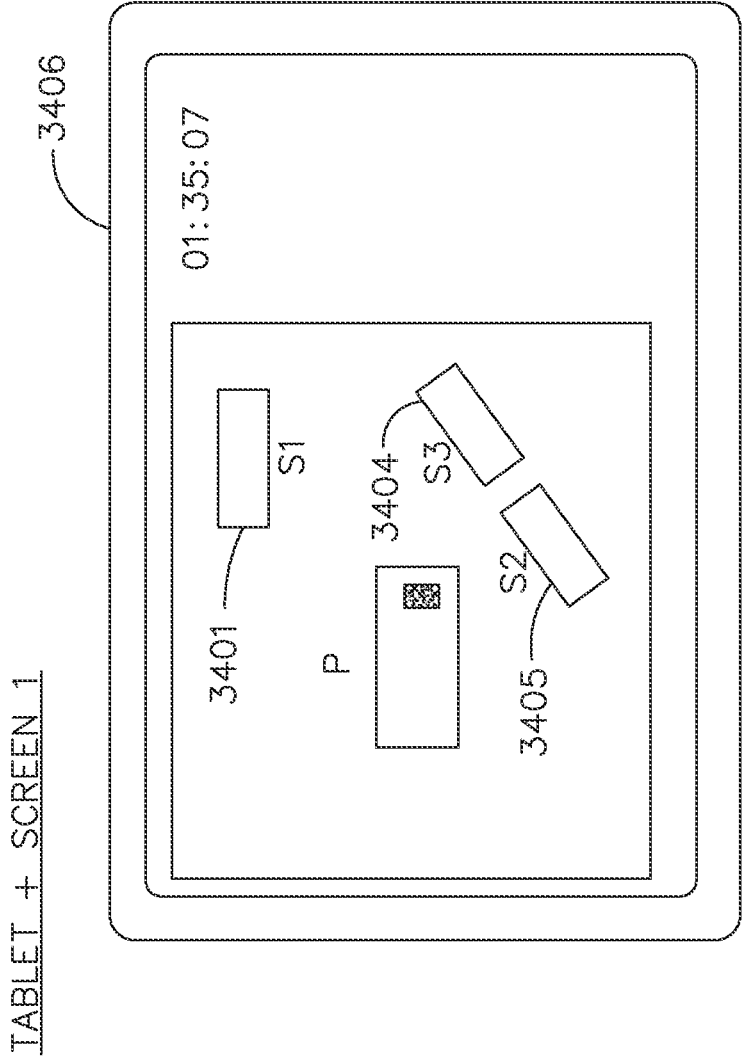


FIG. 60

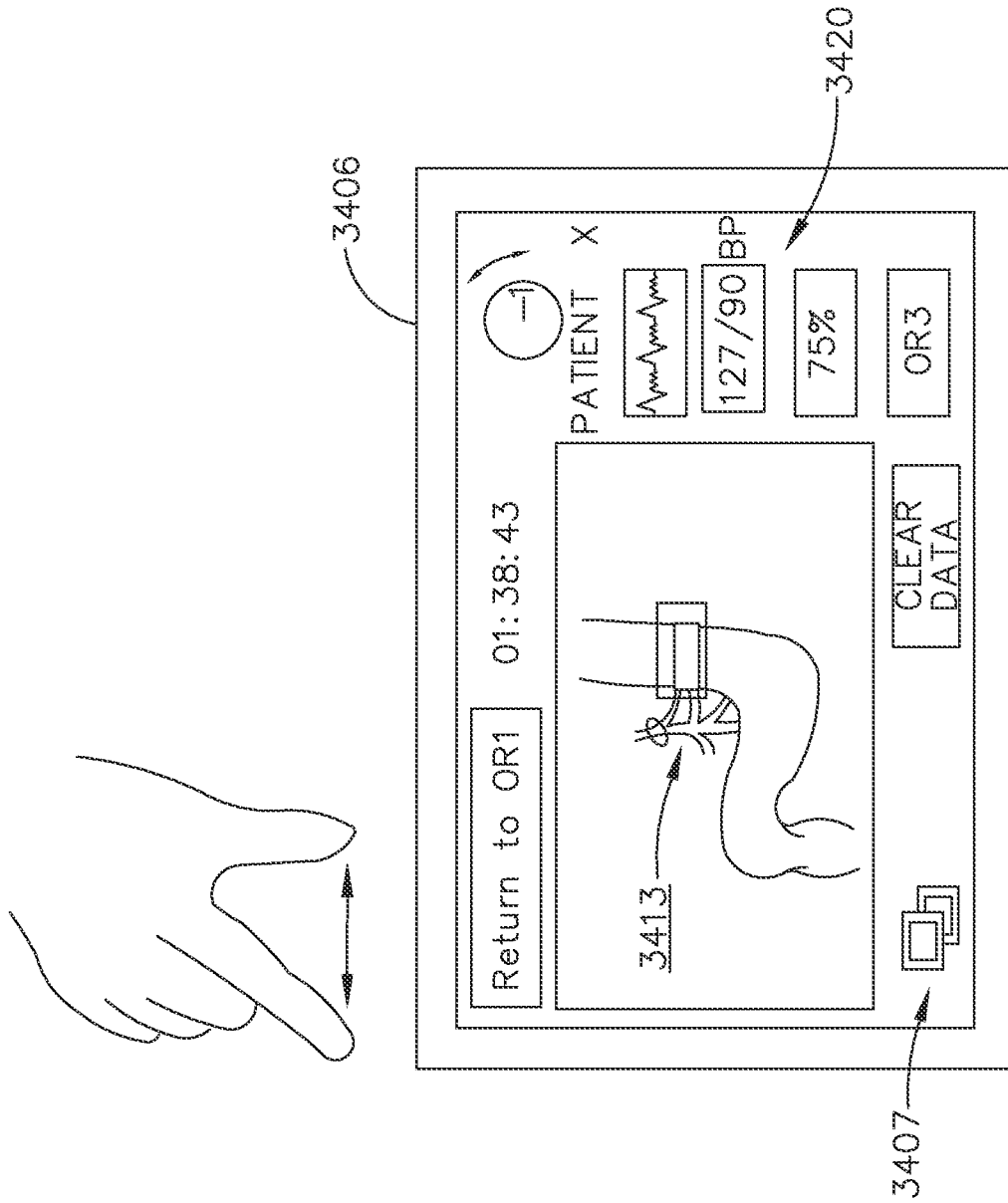


FIG. 61

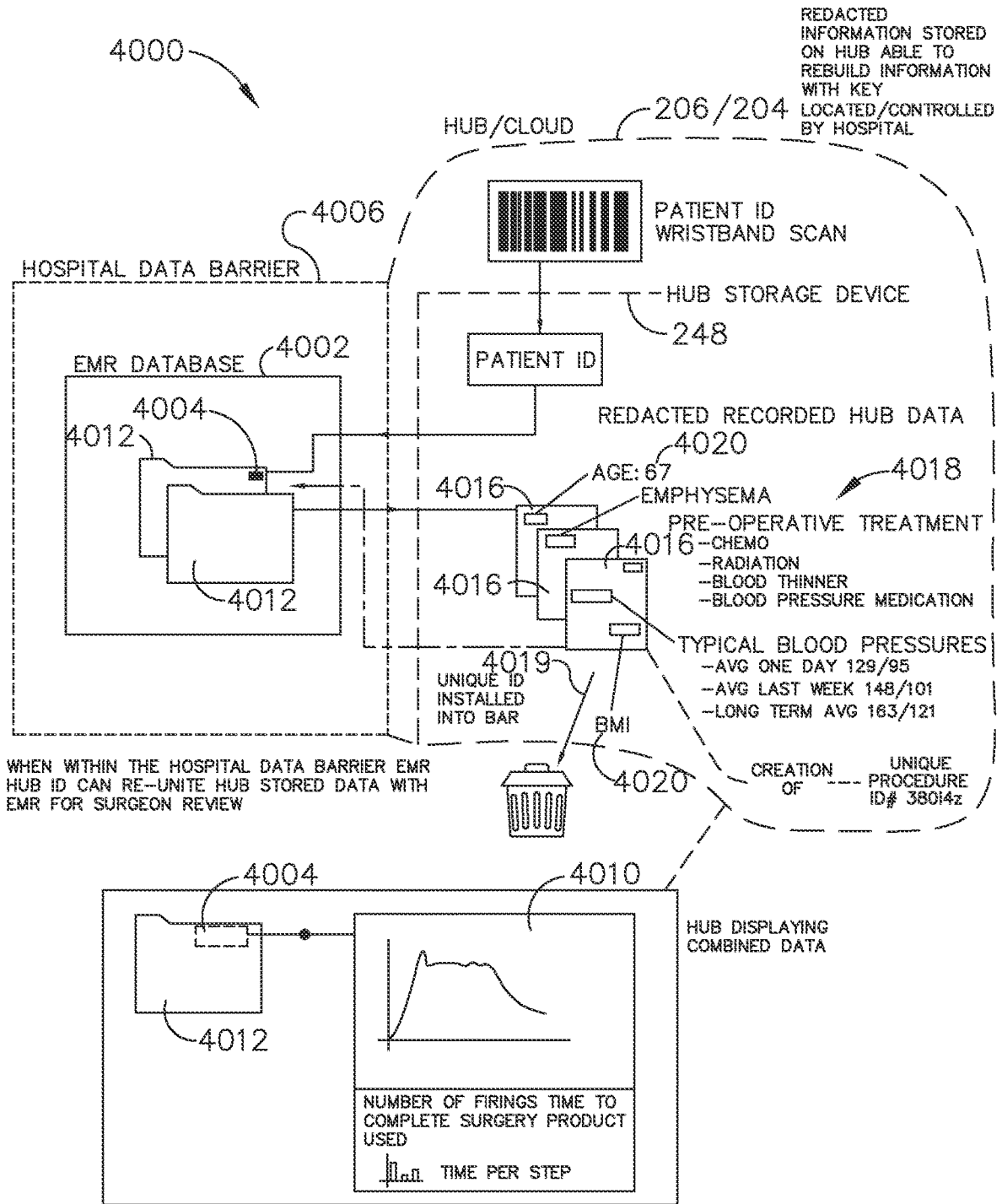


FIG. 62

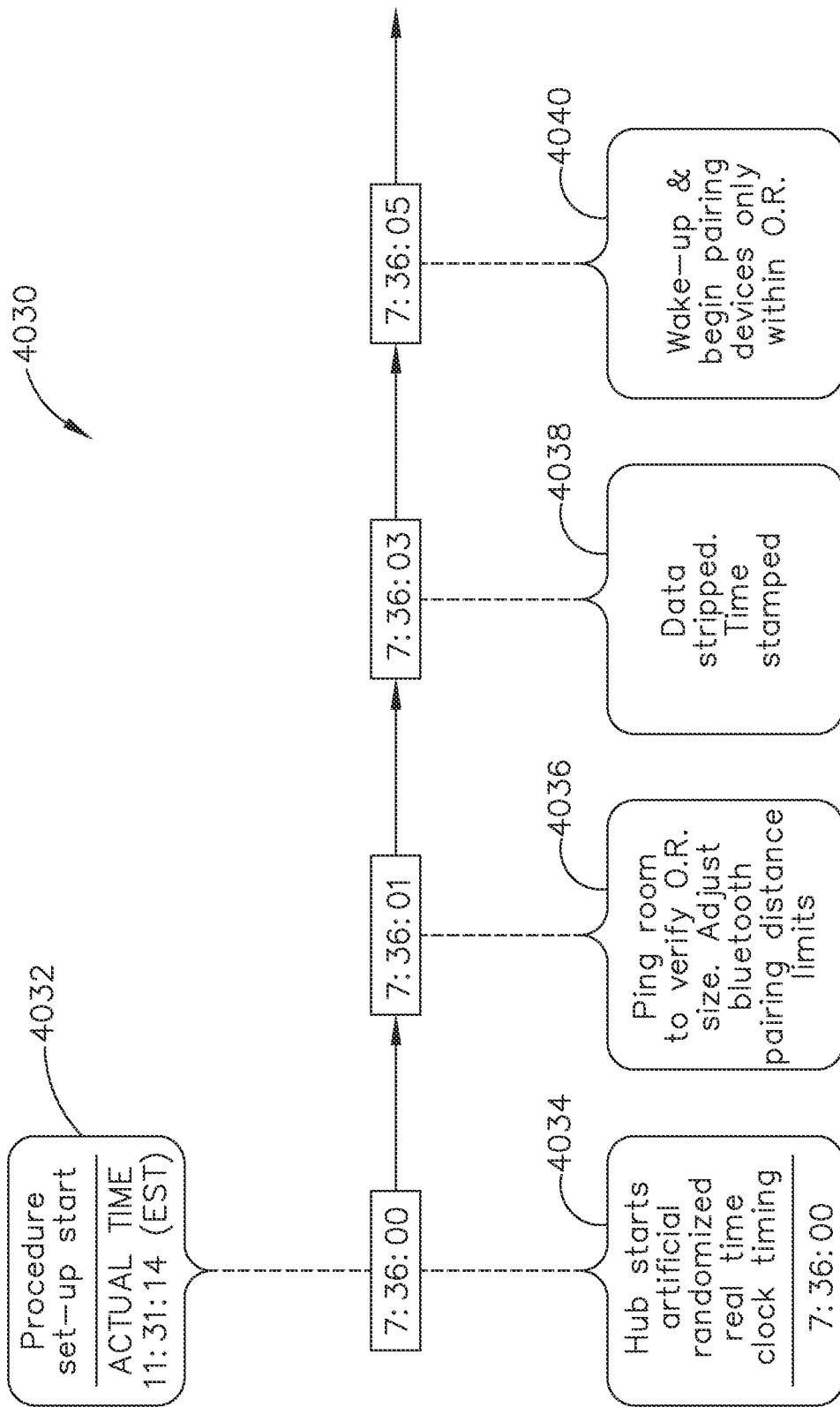


FIG. 63

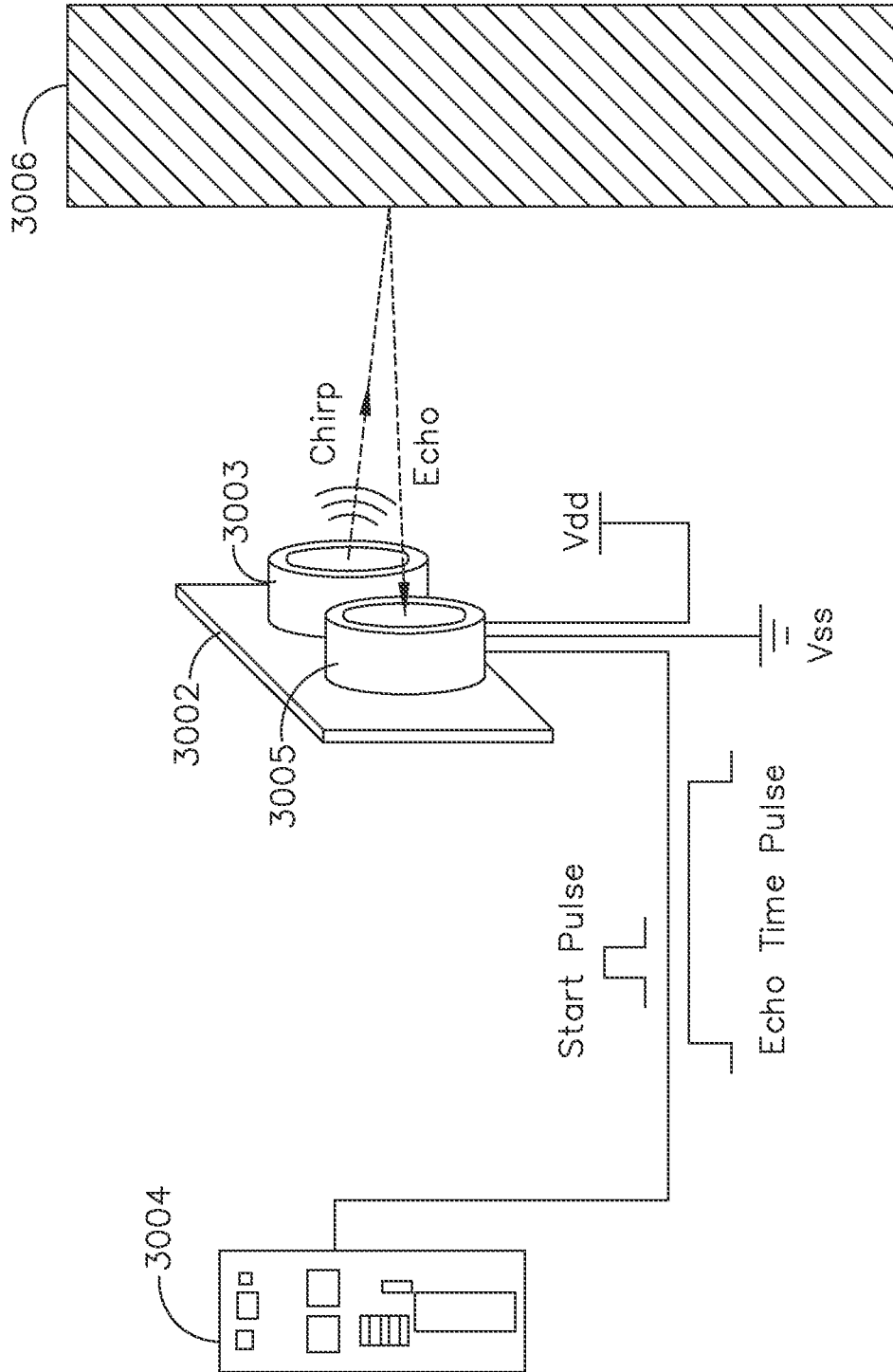


FIG. 64

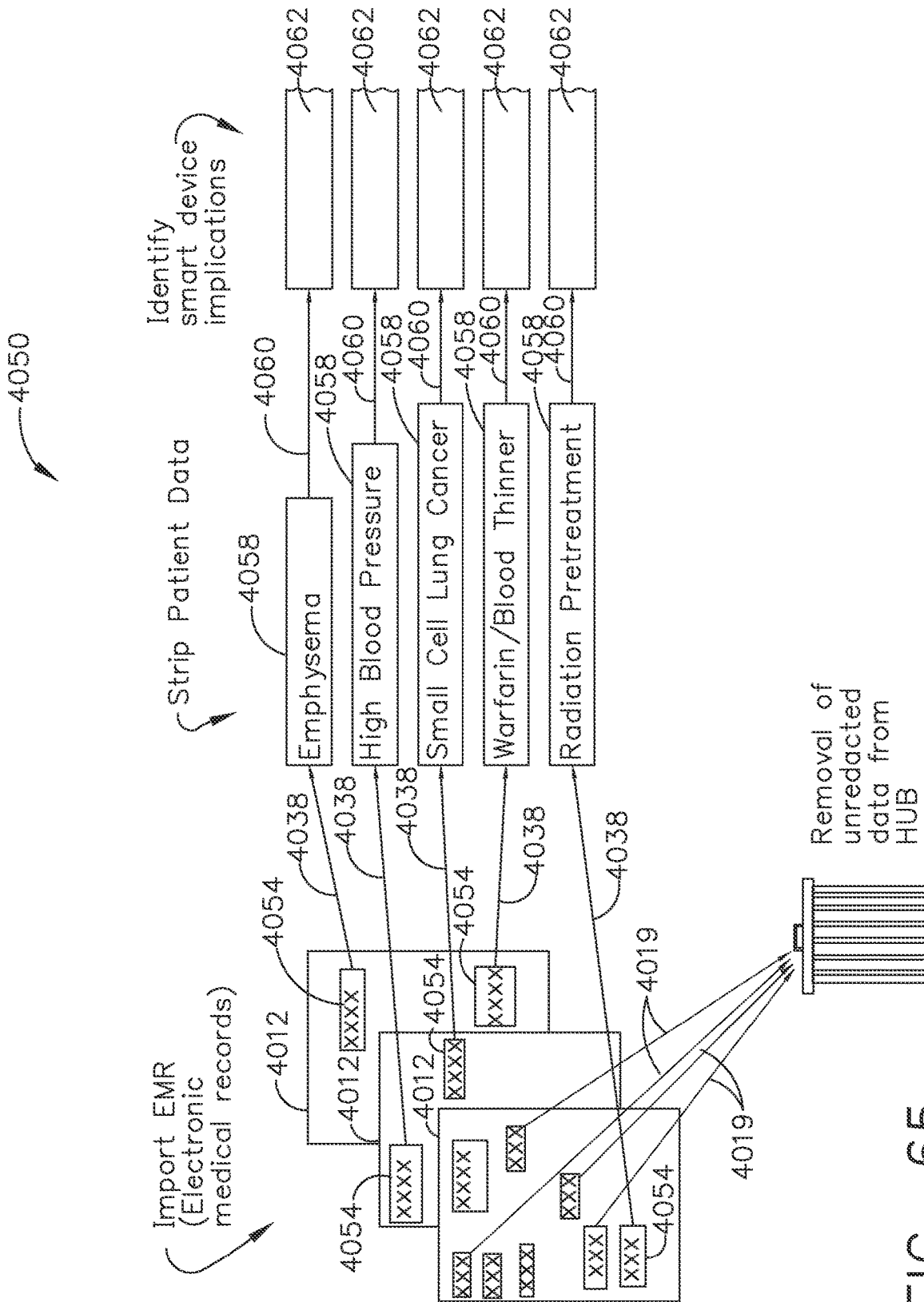


FIG. 65

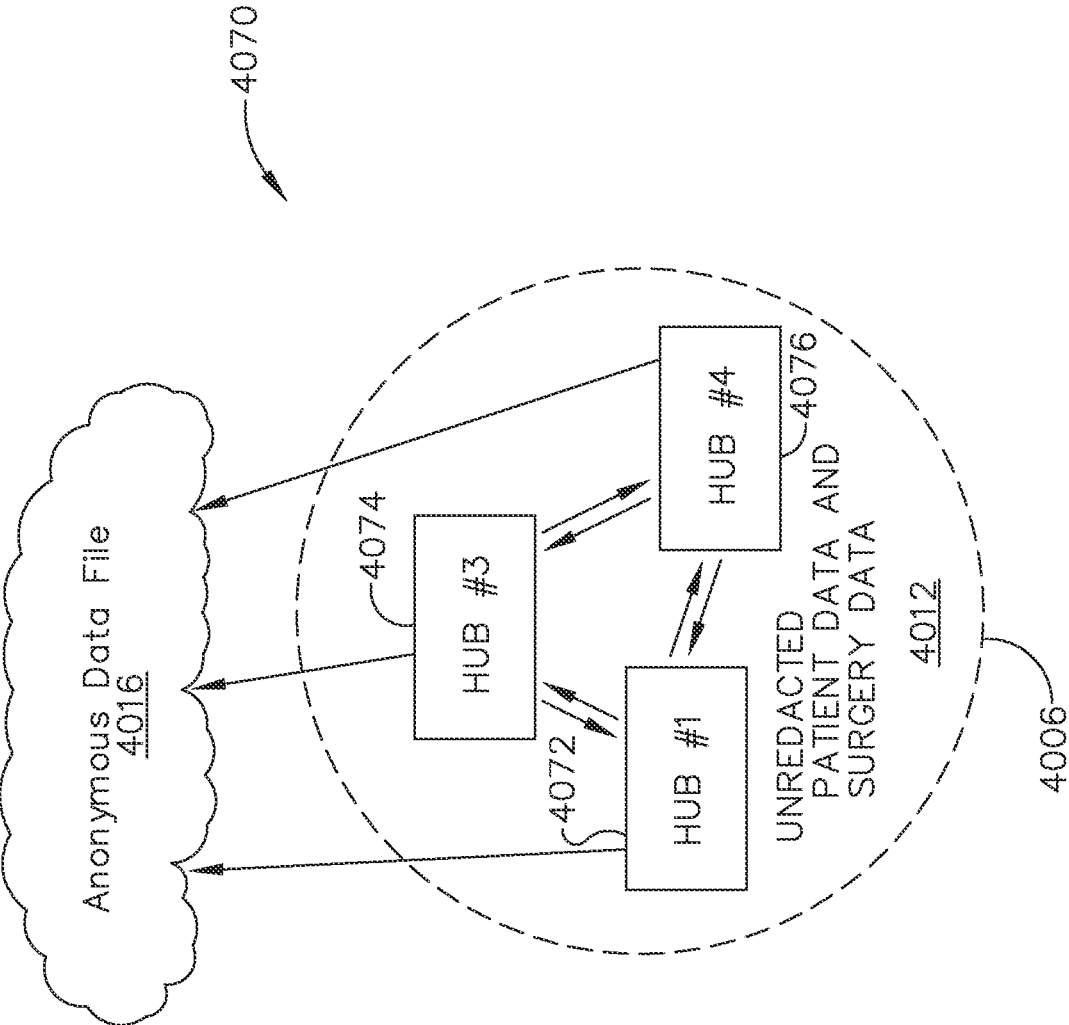


FIG. 66

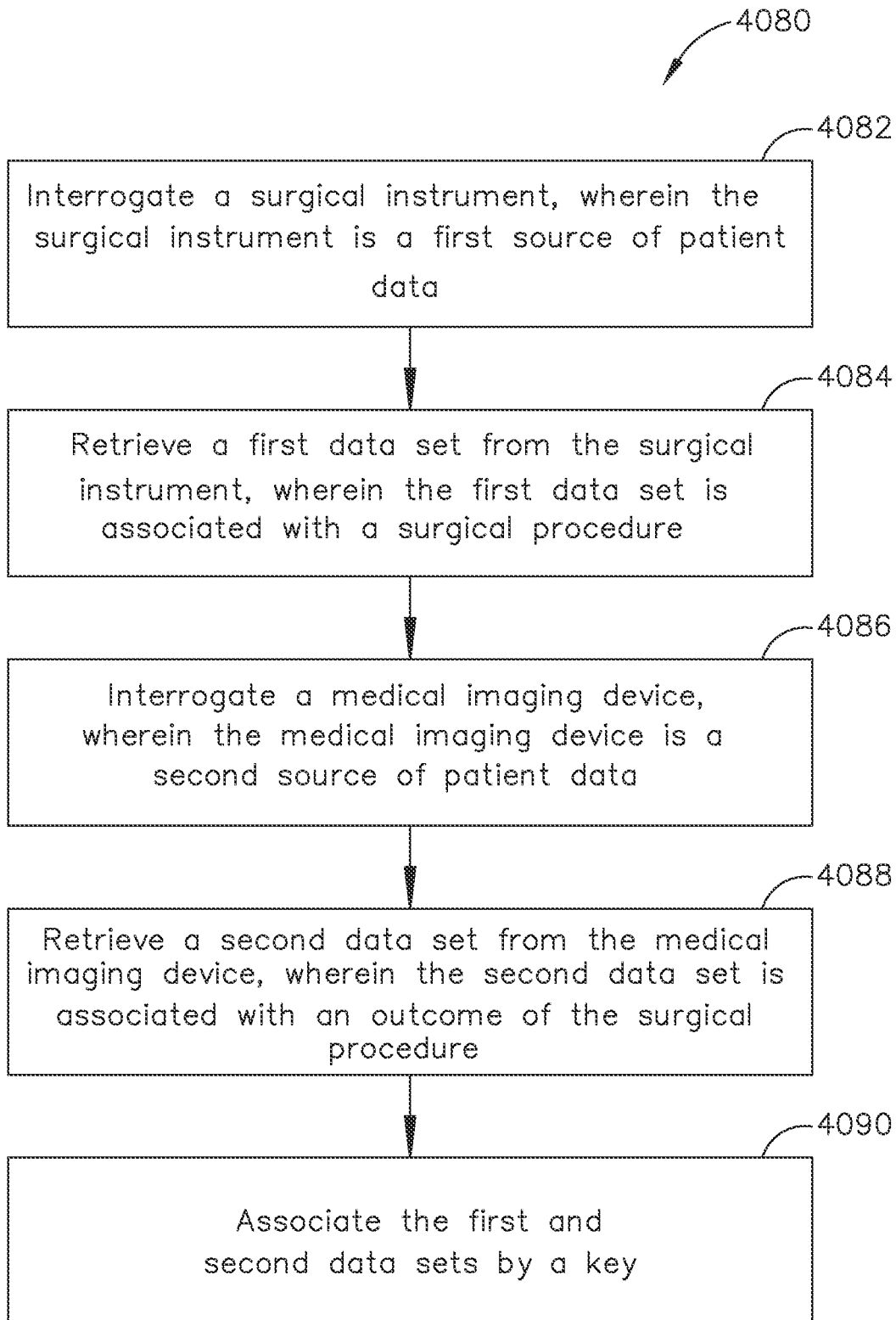


FIG. 67



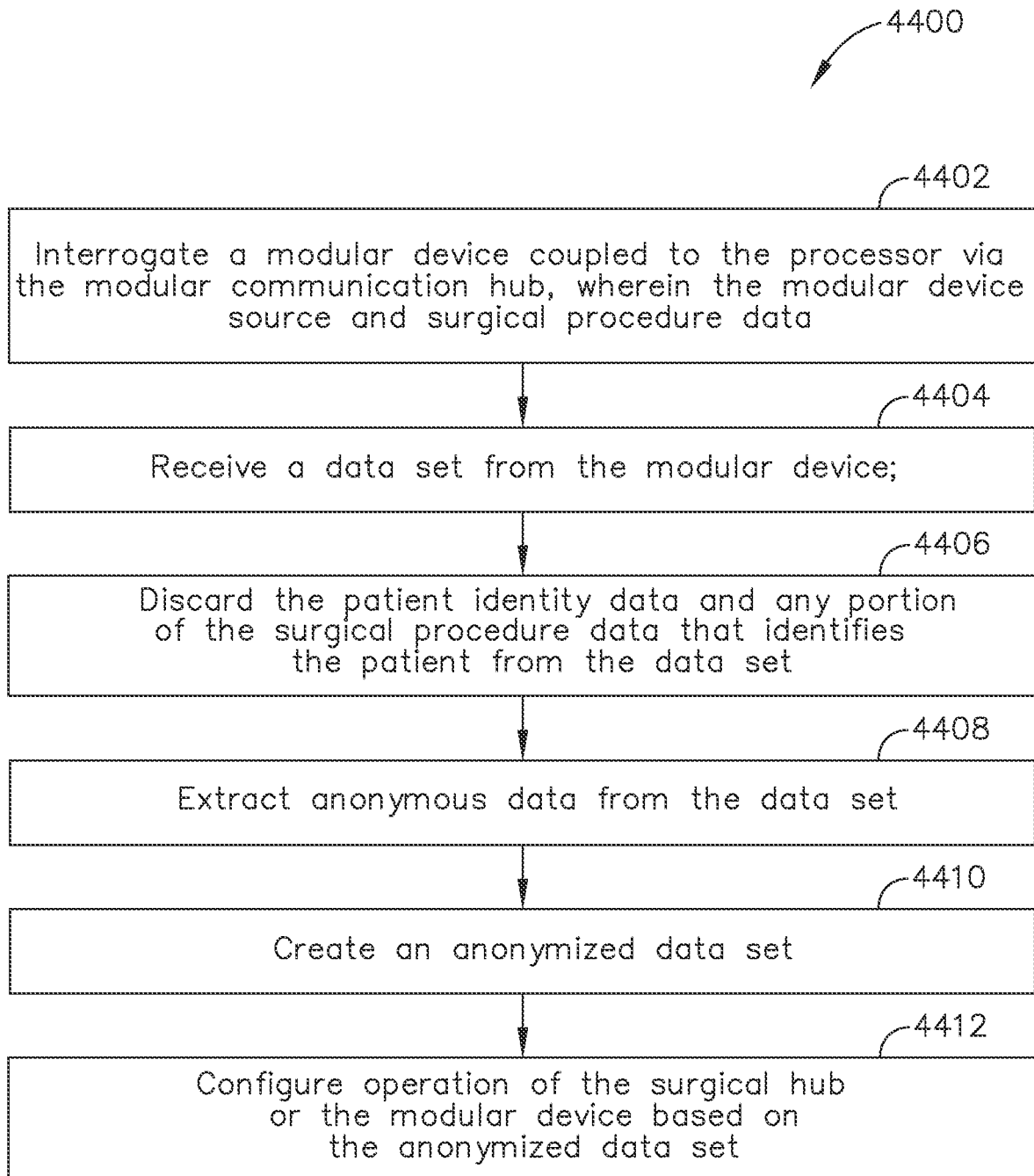


FIG. 68

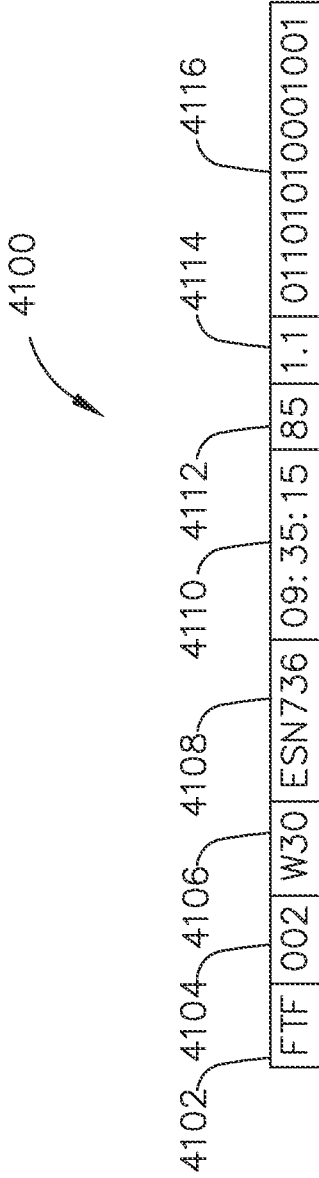


FIG. 69

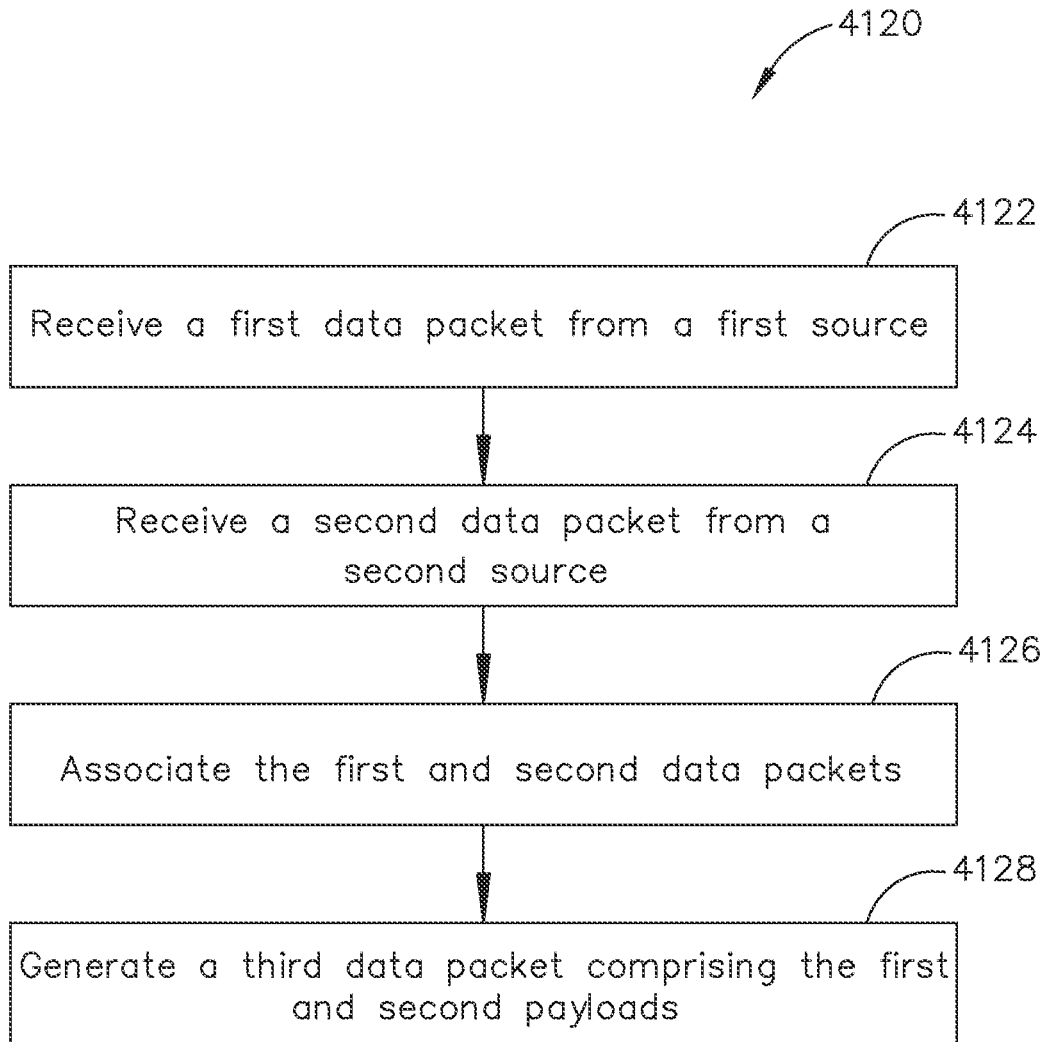


FIG. 70

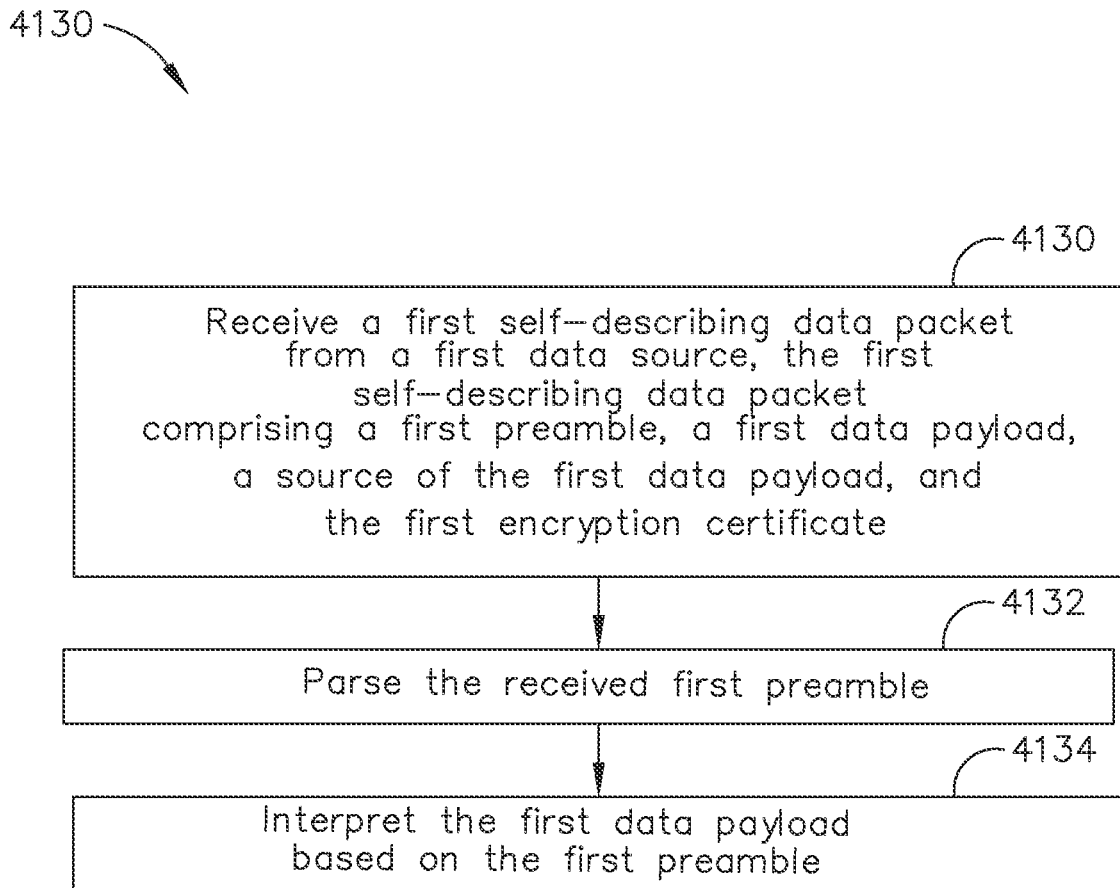


FIG. 71

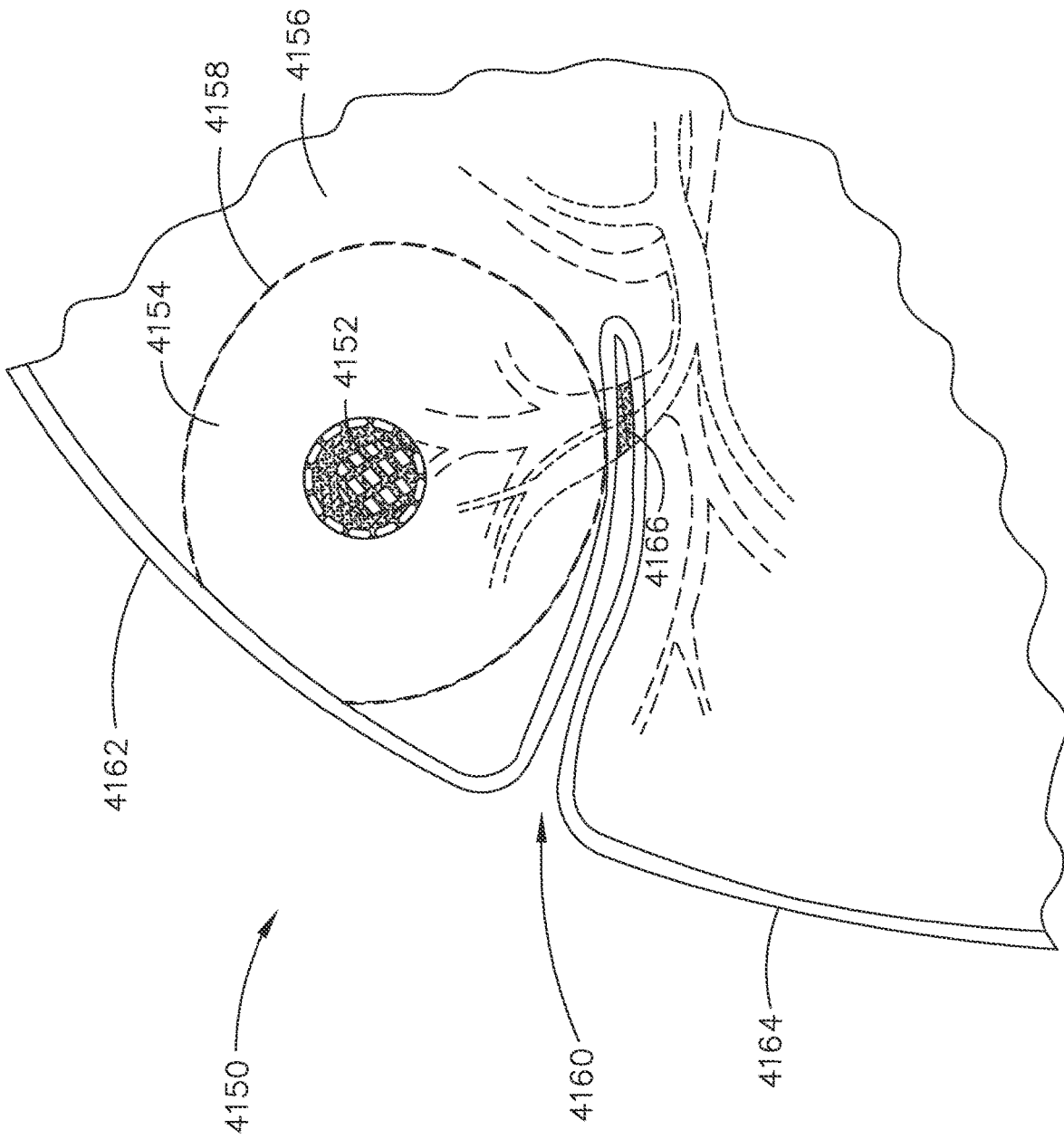


FIG. 72

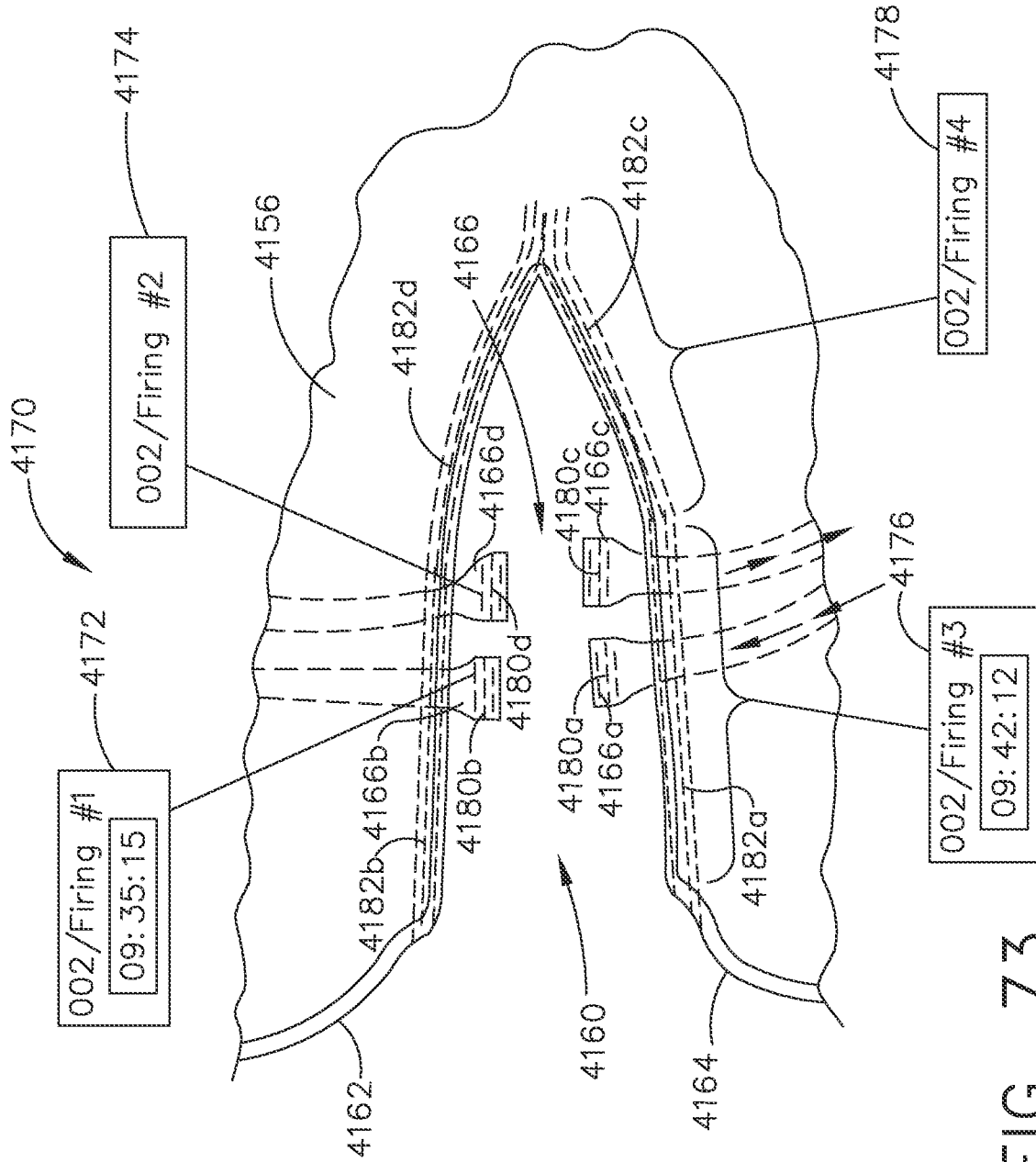


FIG. 73

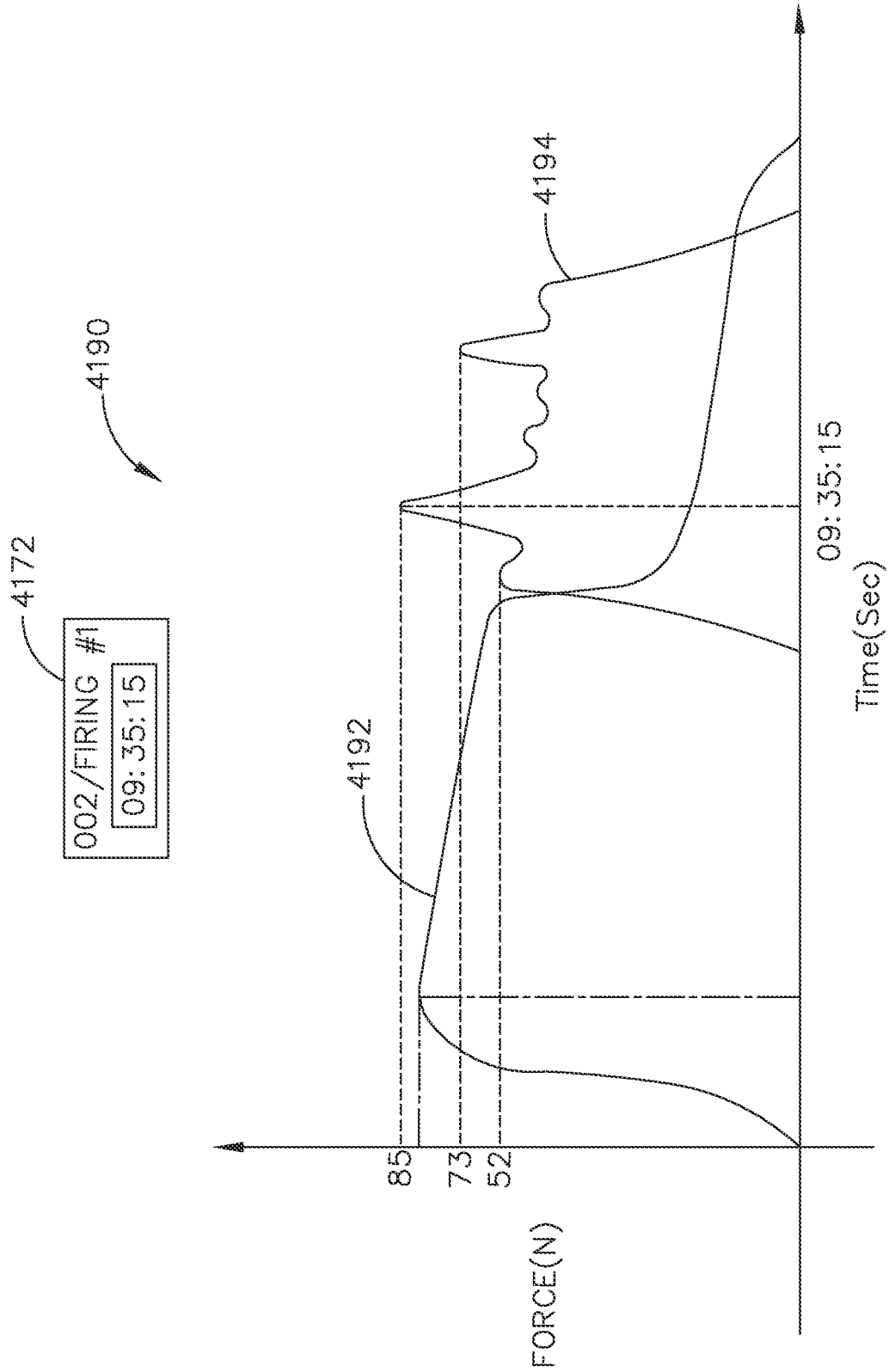


FIG. 74

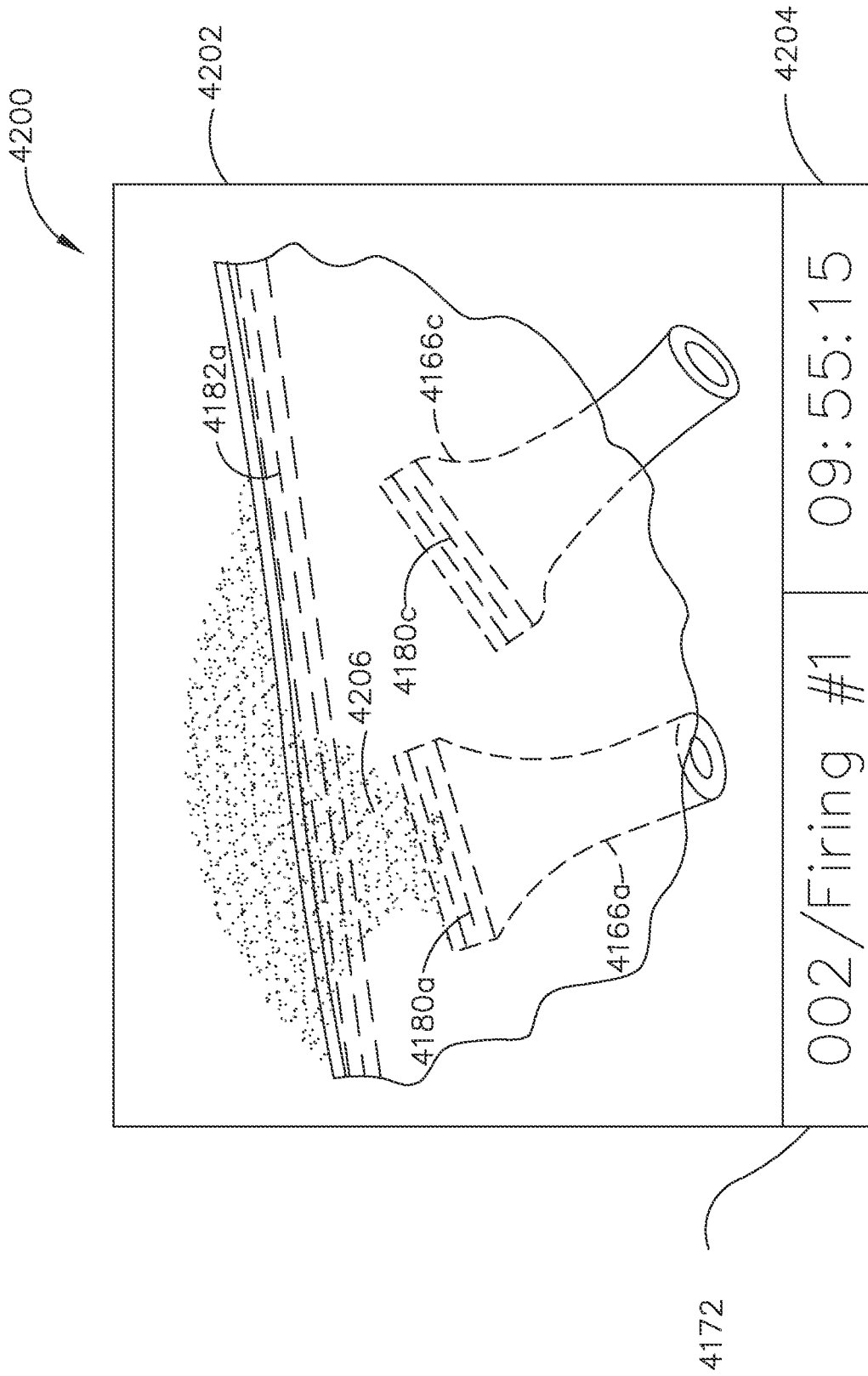


FIG. 75



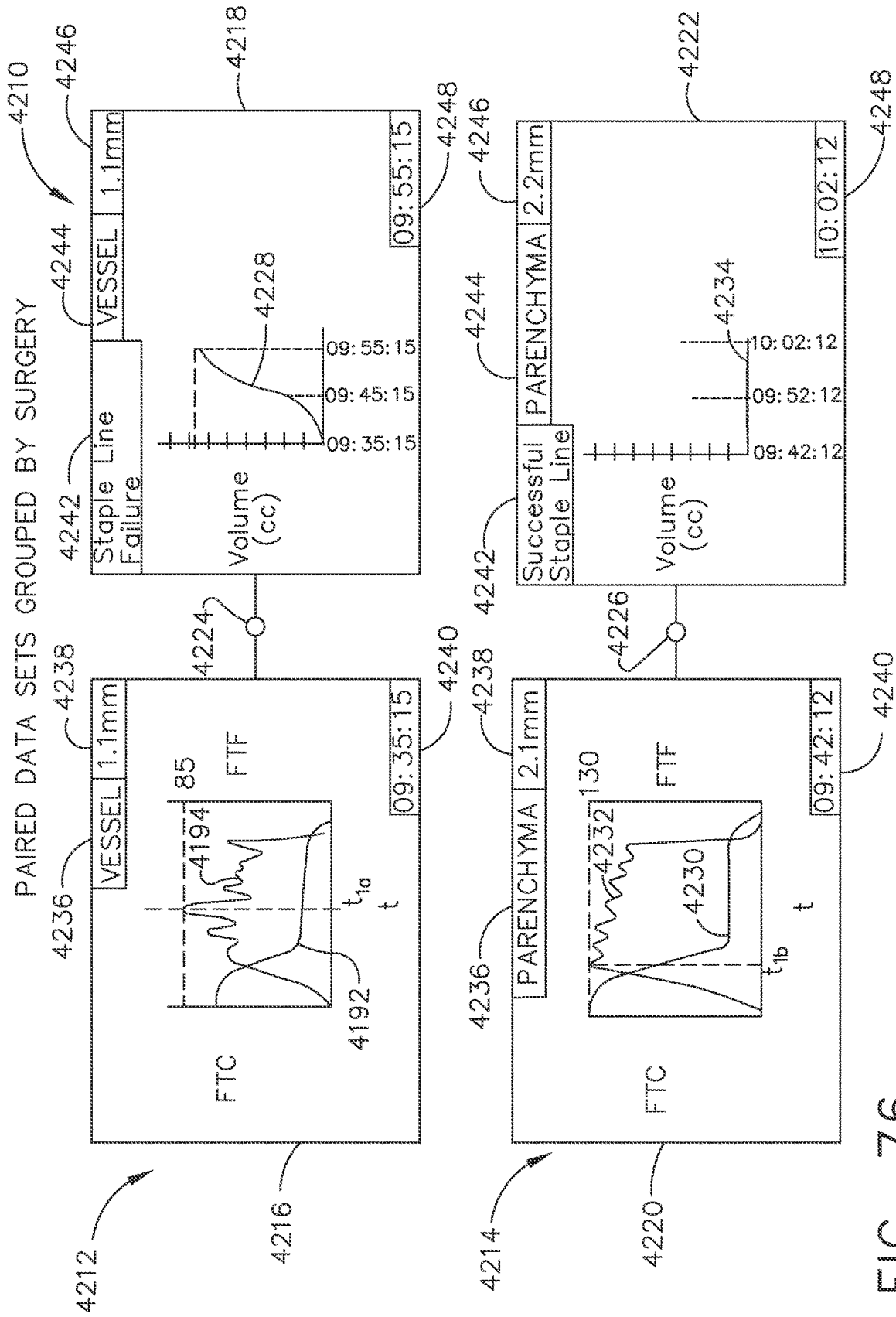


FIG. 76

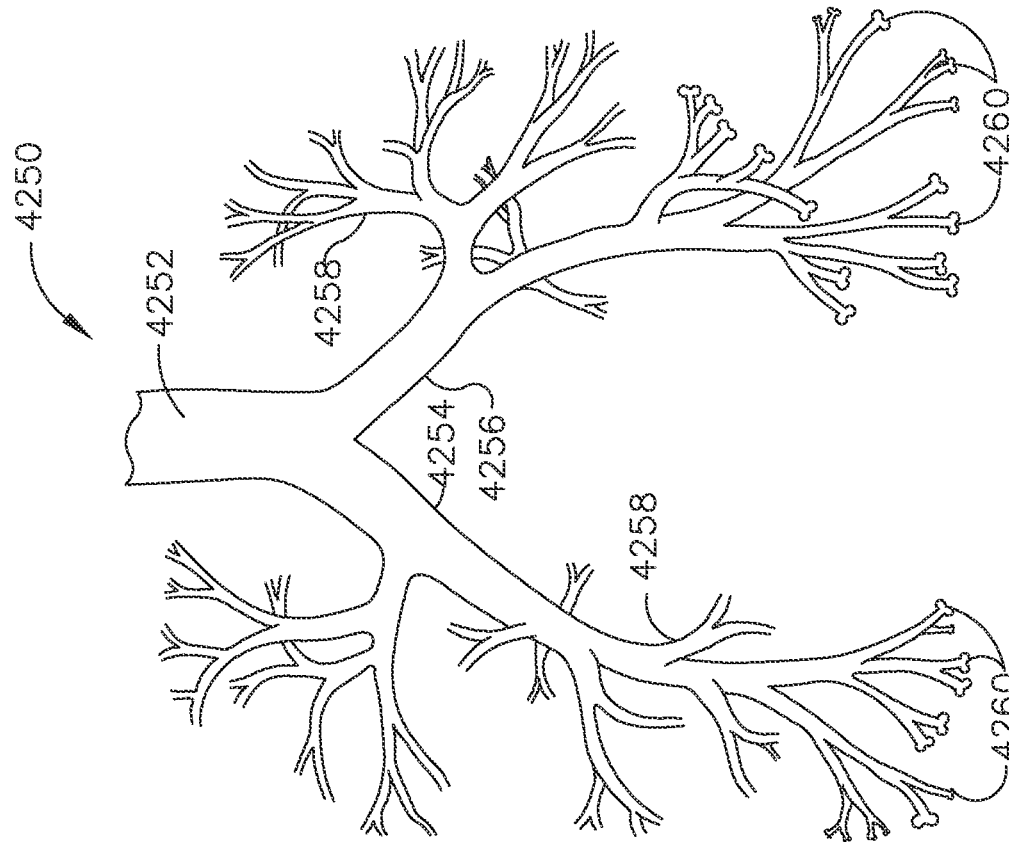


FIG. 78

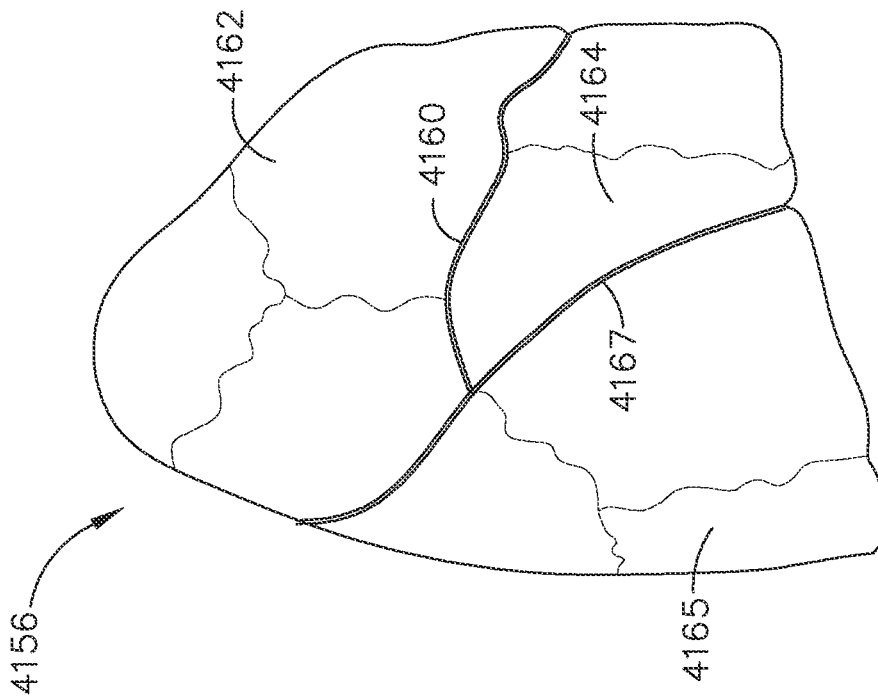


FIG. 77

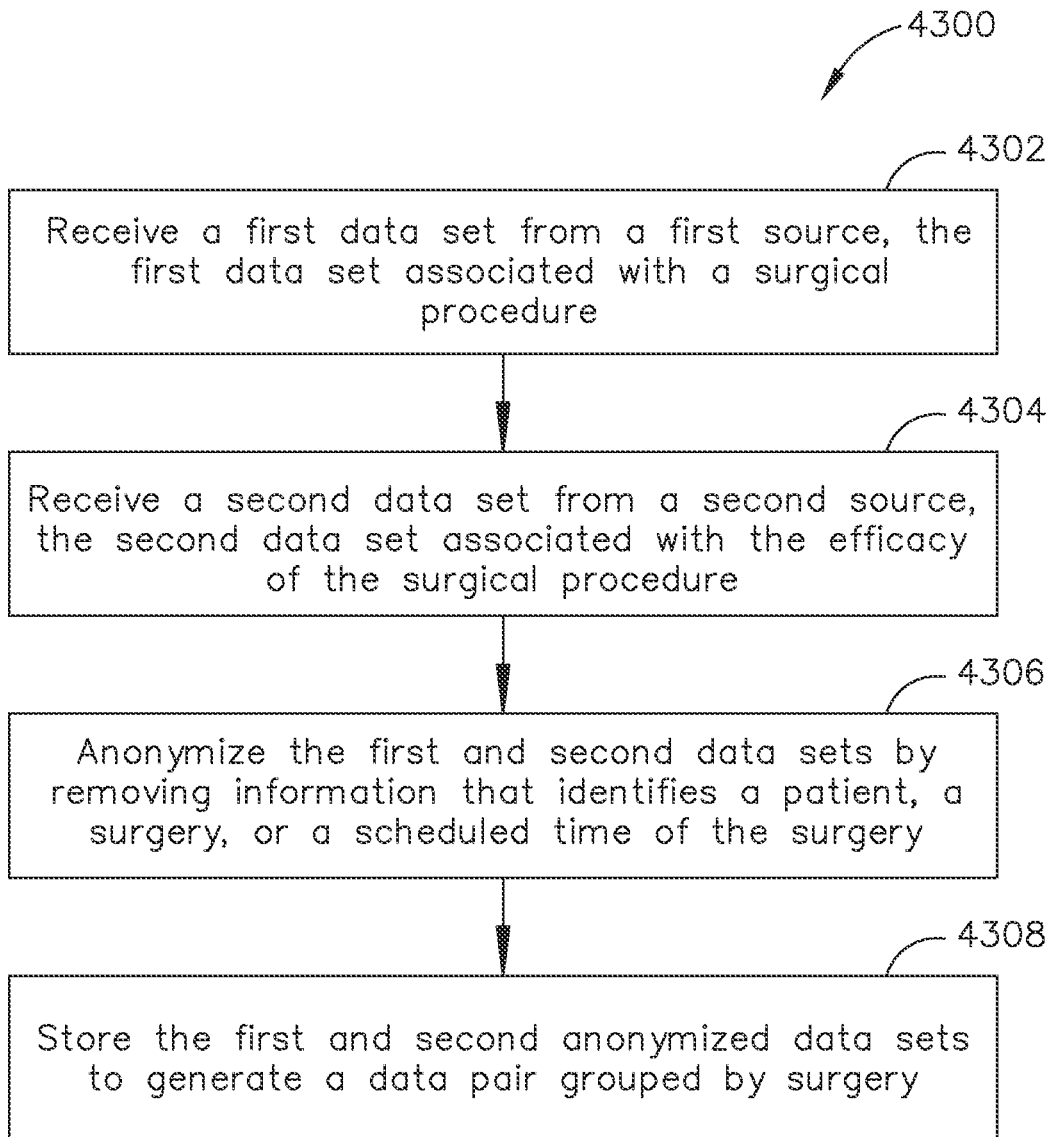


FIG. 79

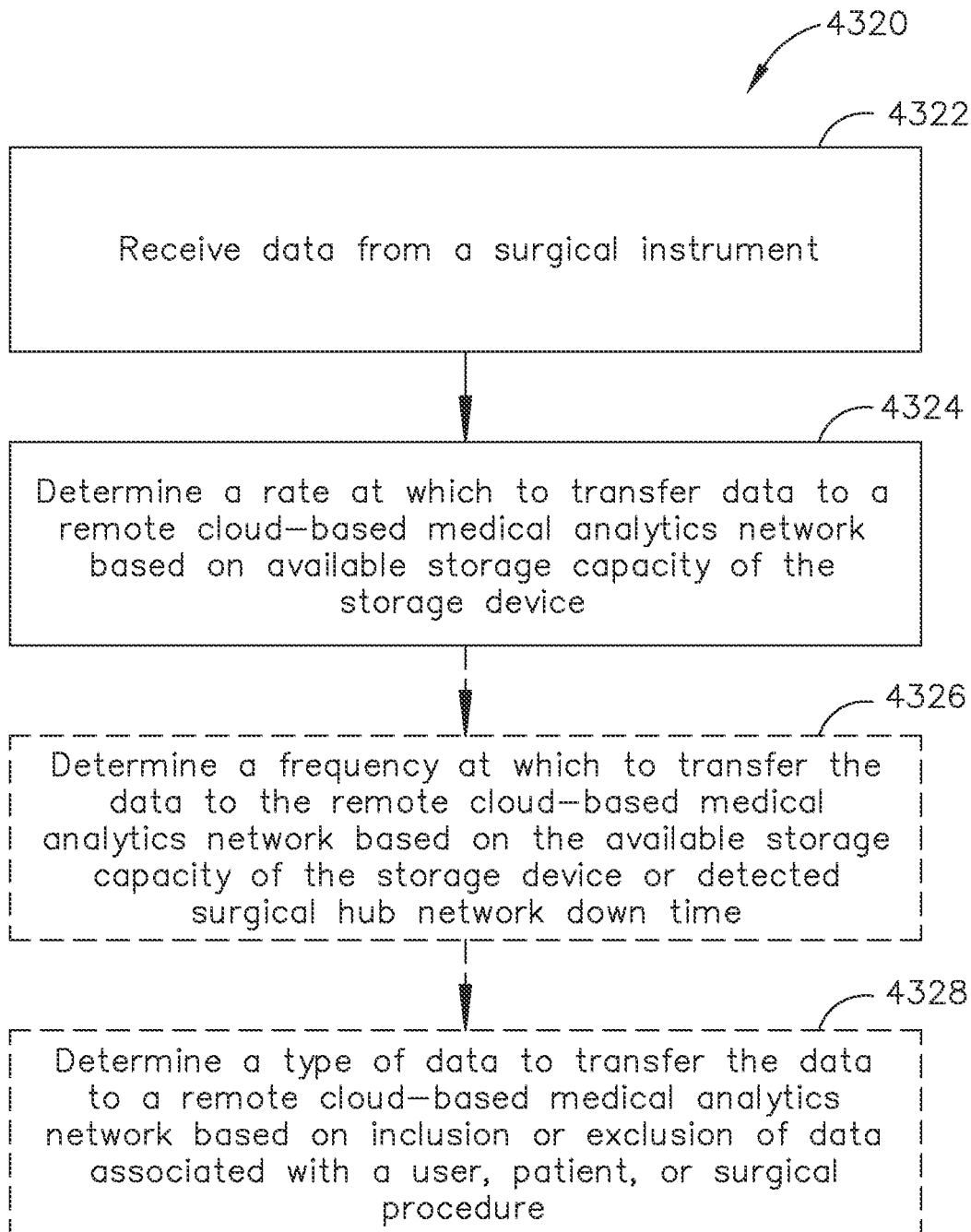


FIG. 80

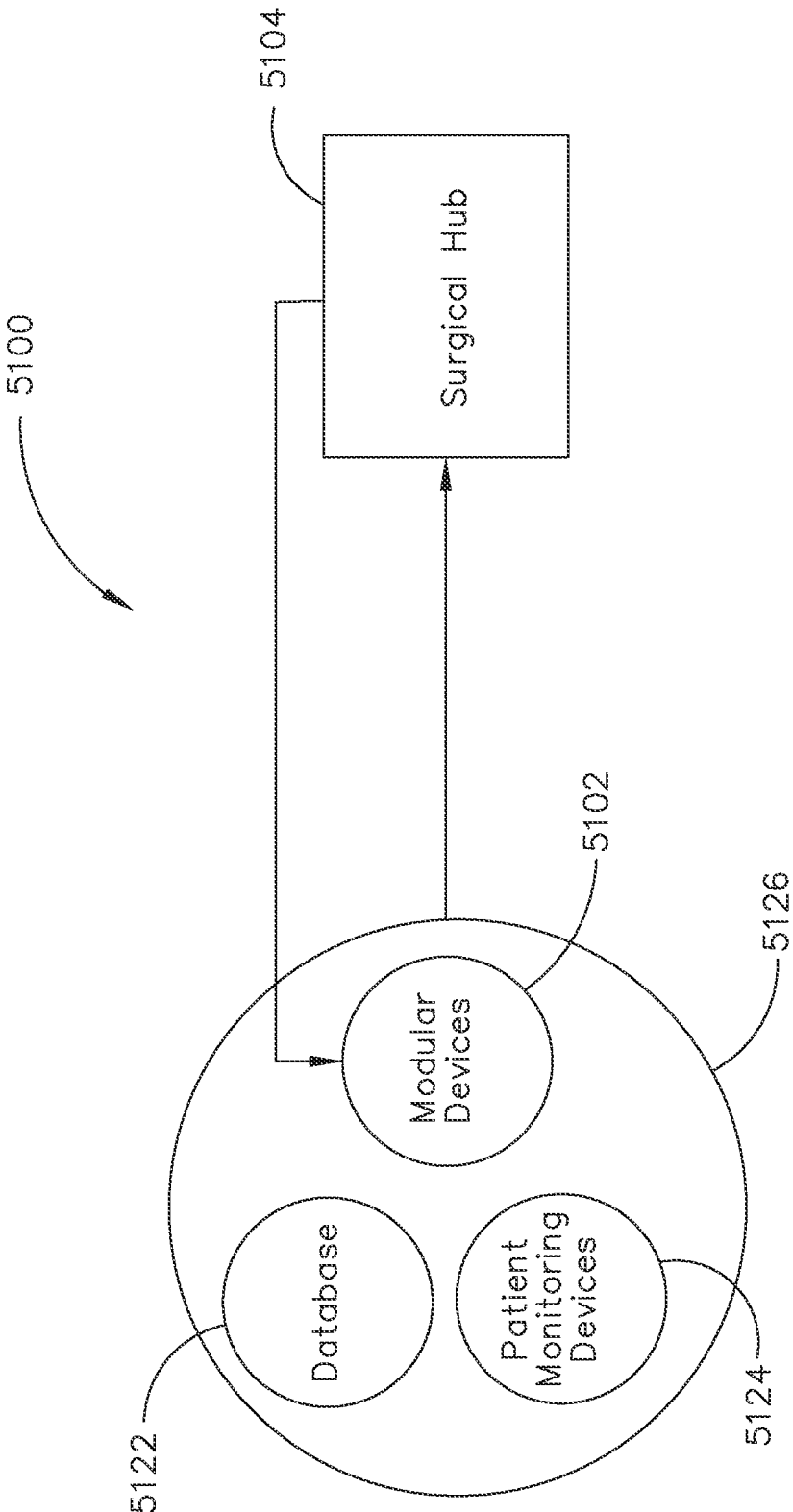


FIG. 81

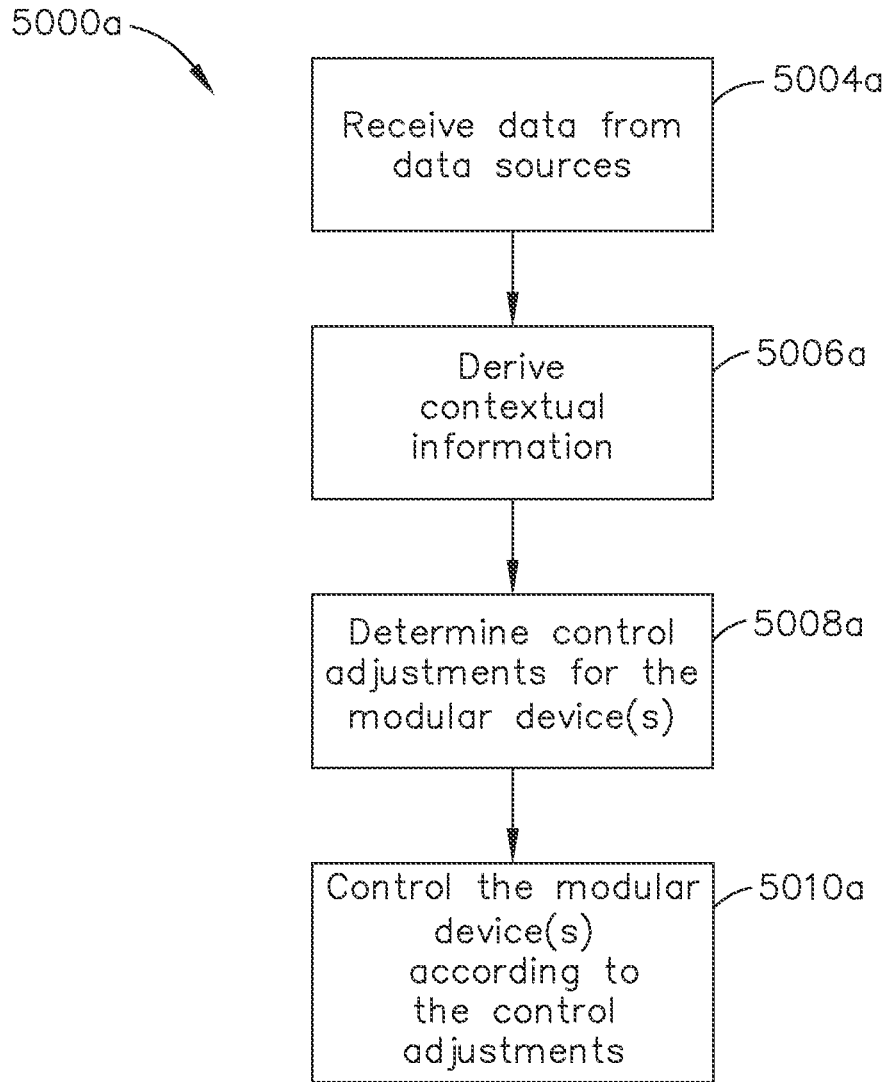


FIG. 82A

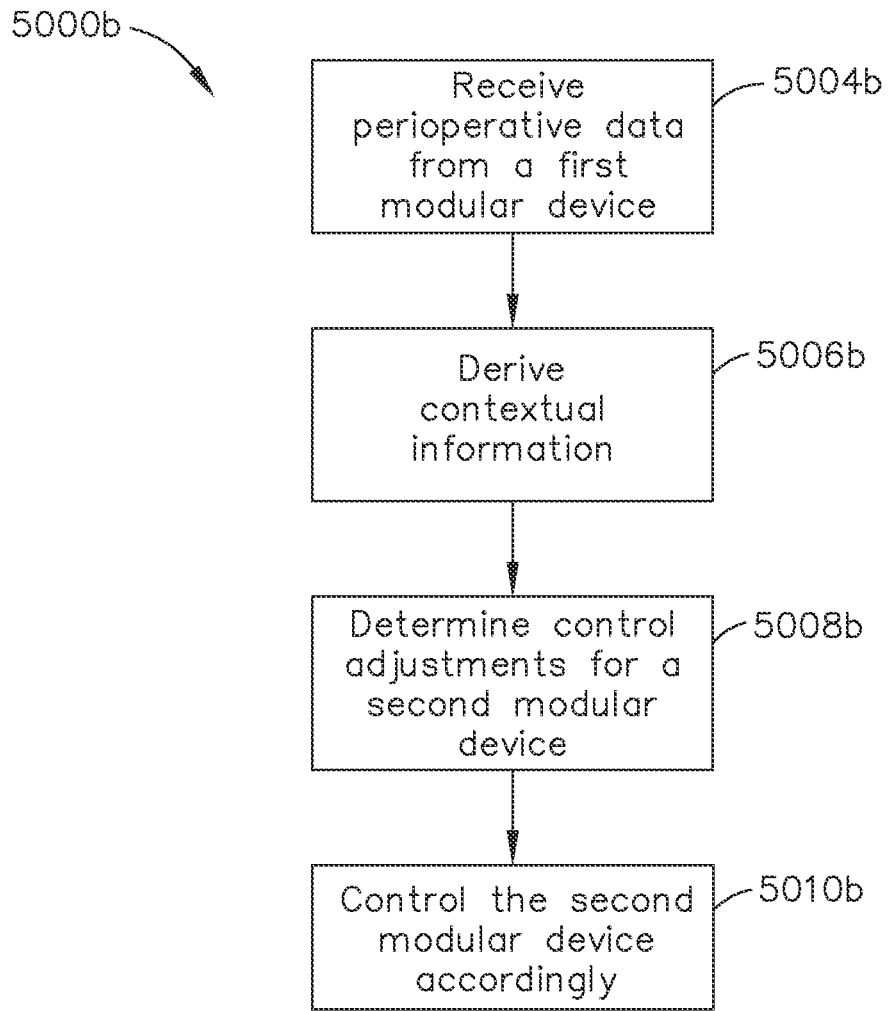


FIG. 82B

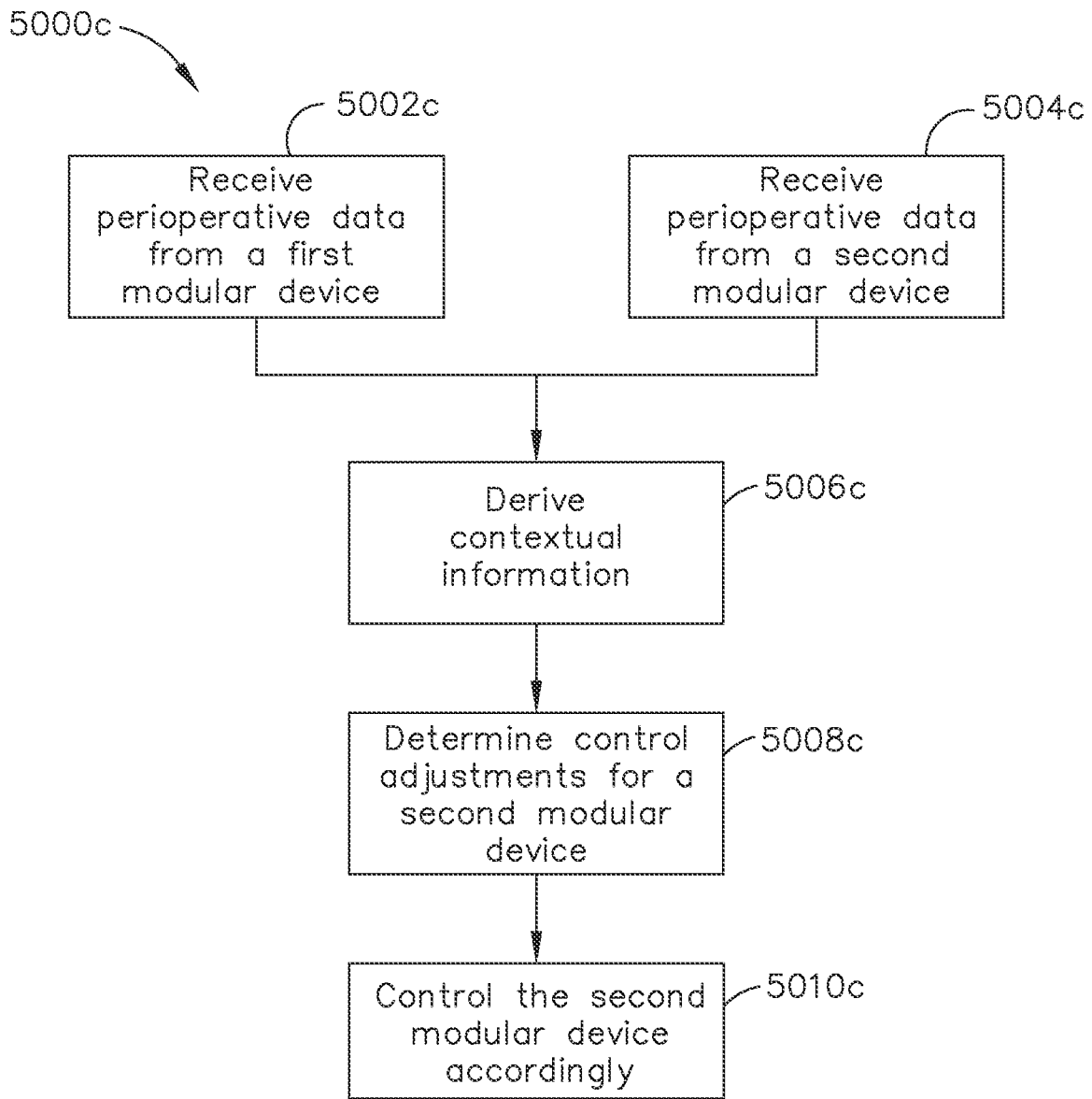


FIG. 82C



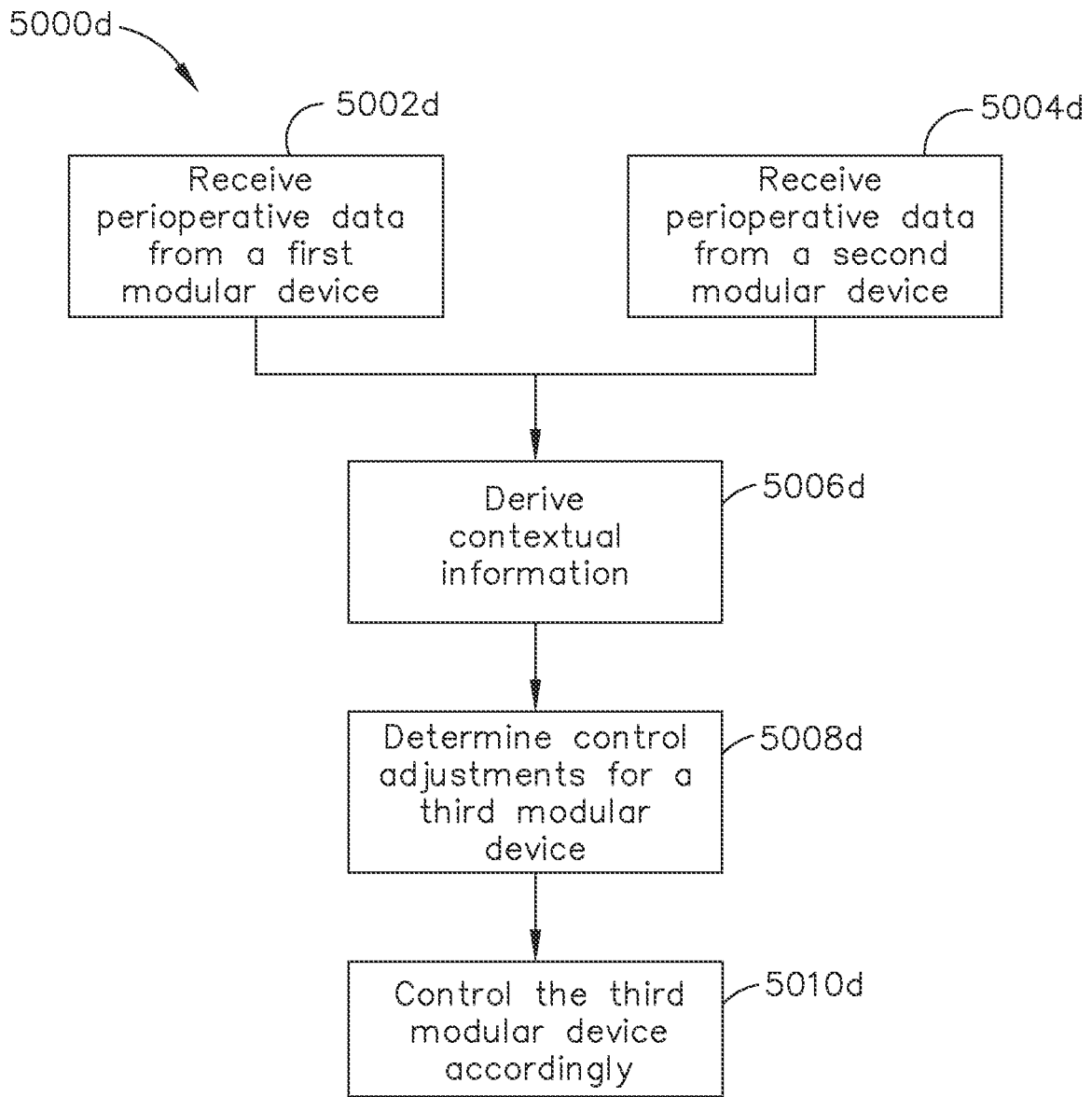
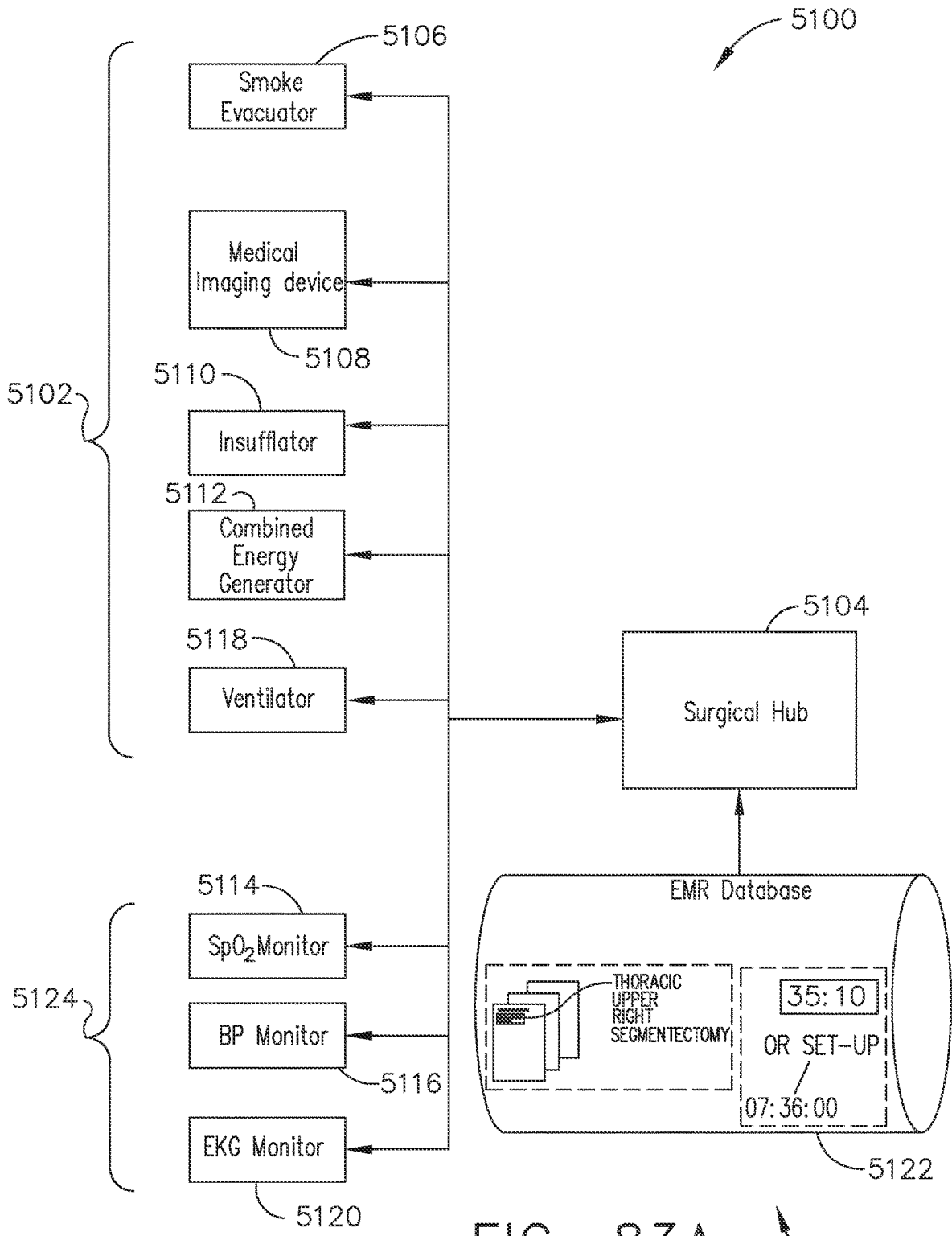


FIG. 82D



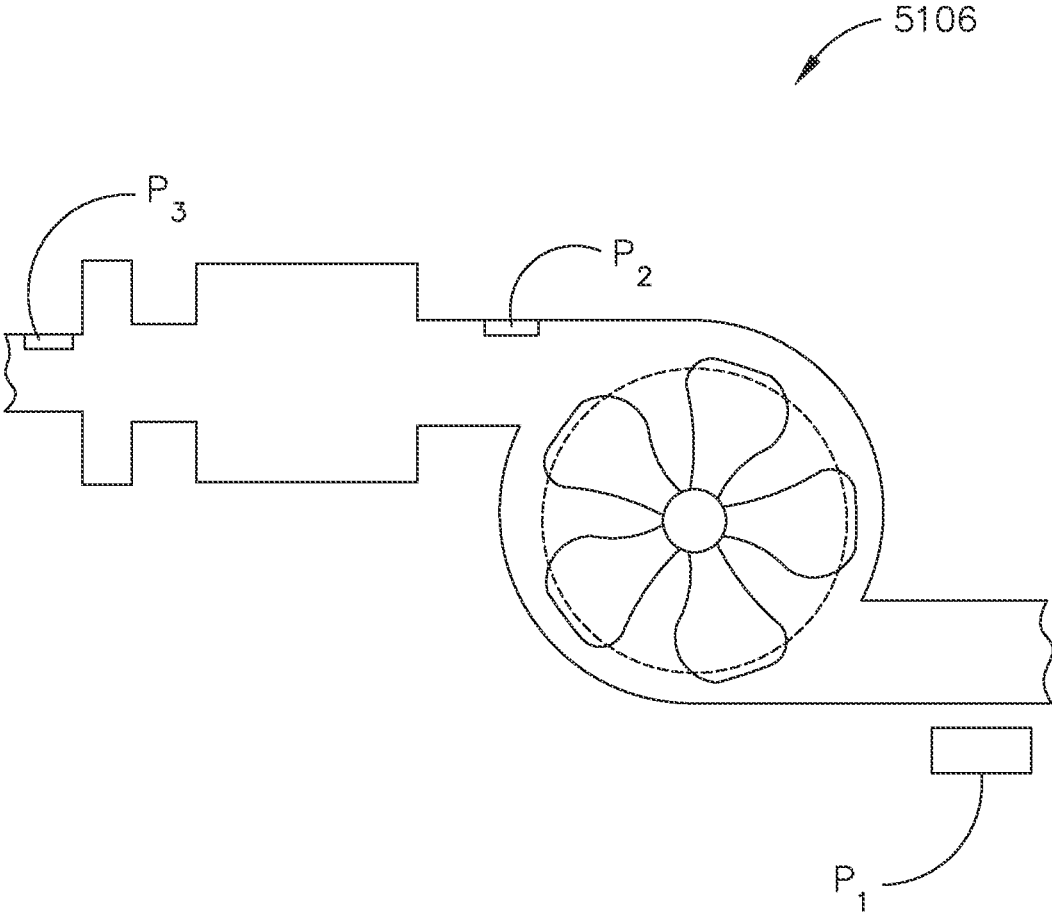


FIG. 83B

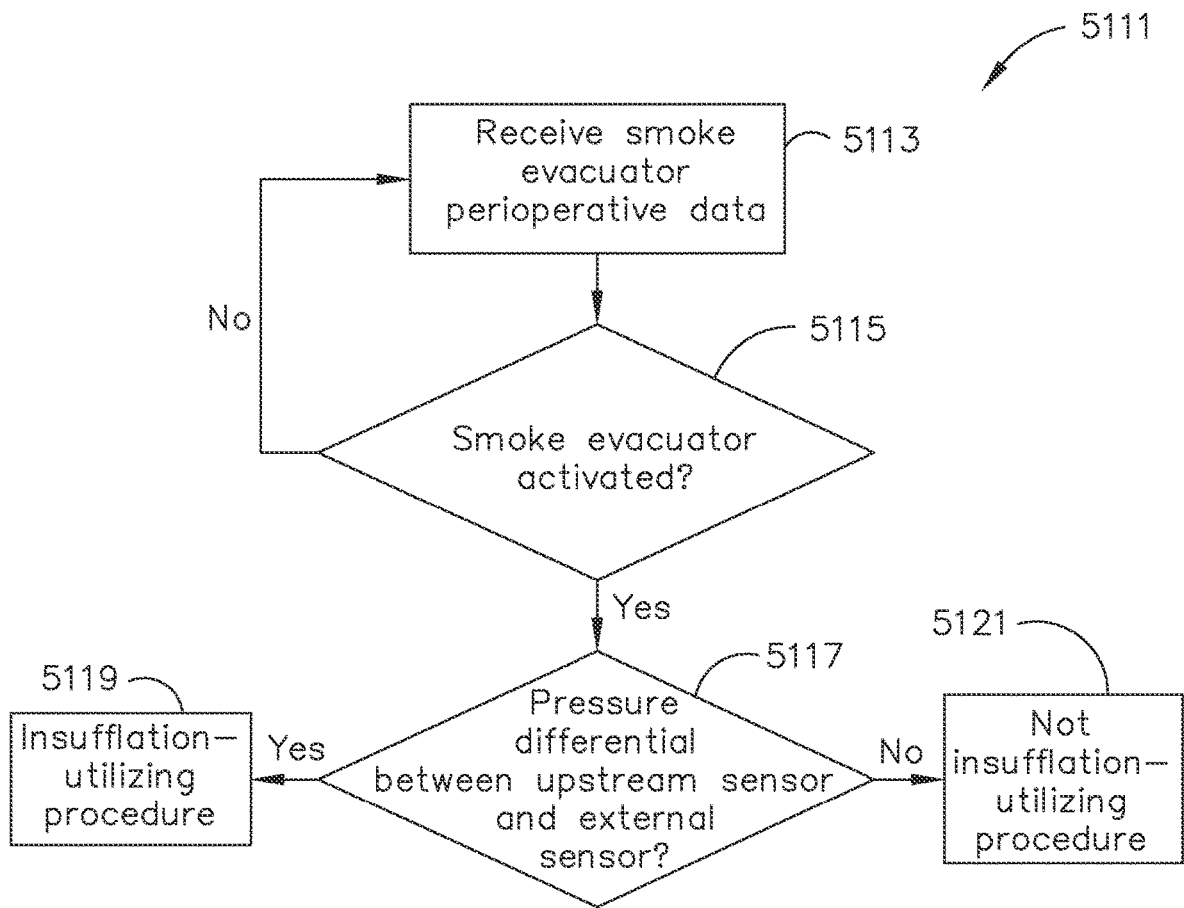


FIG. 84A

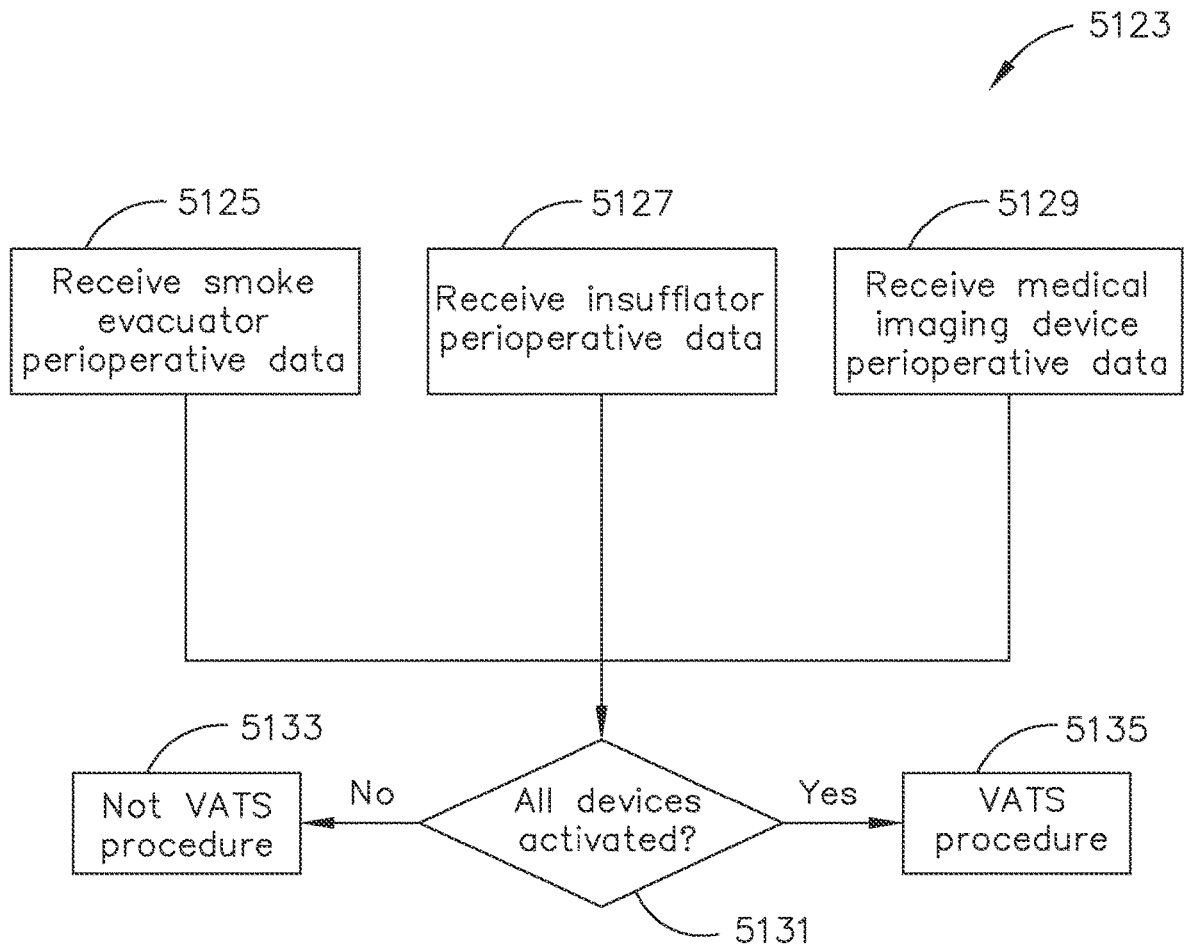


FIG. 84B

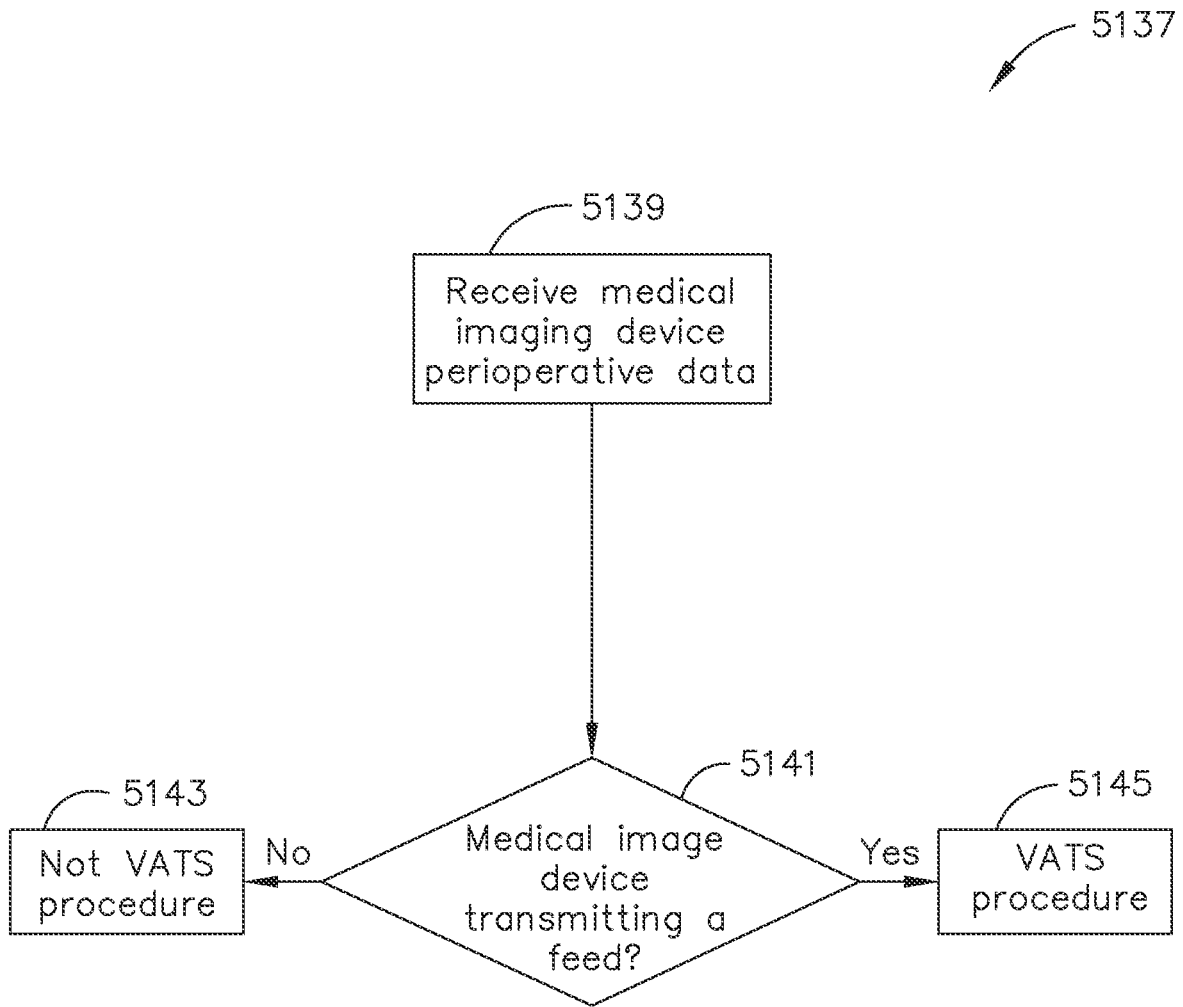


FIG. 84C

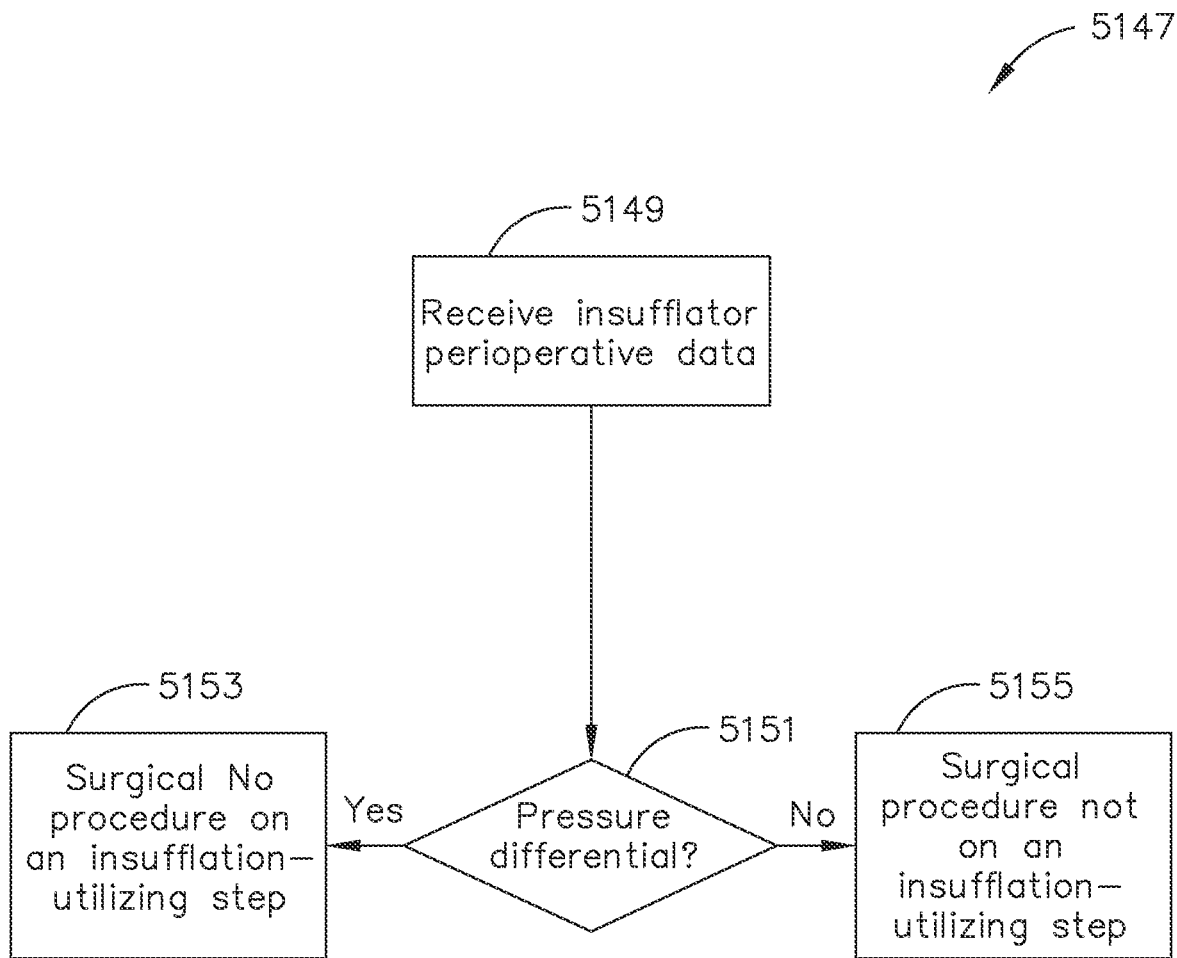


FIG. 84D

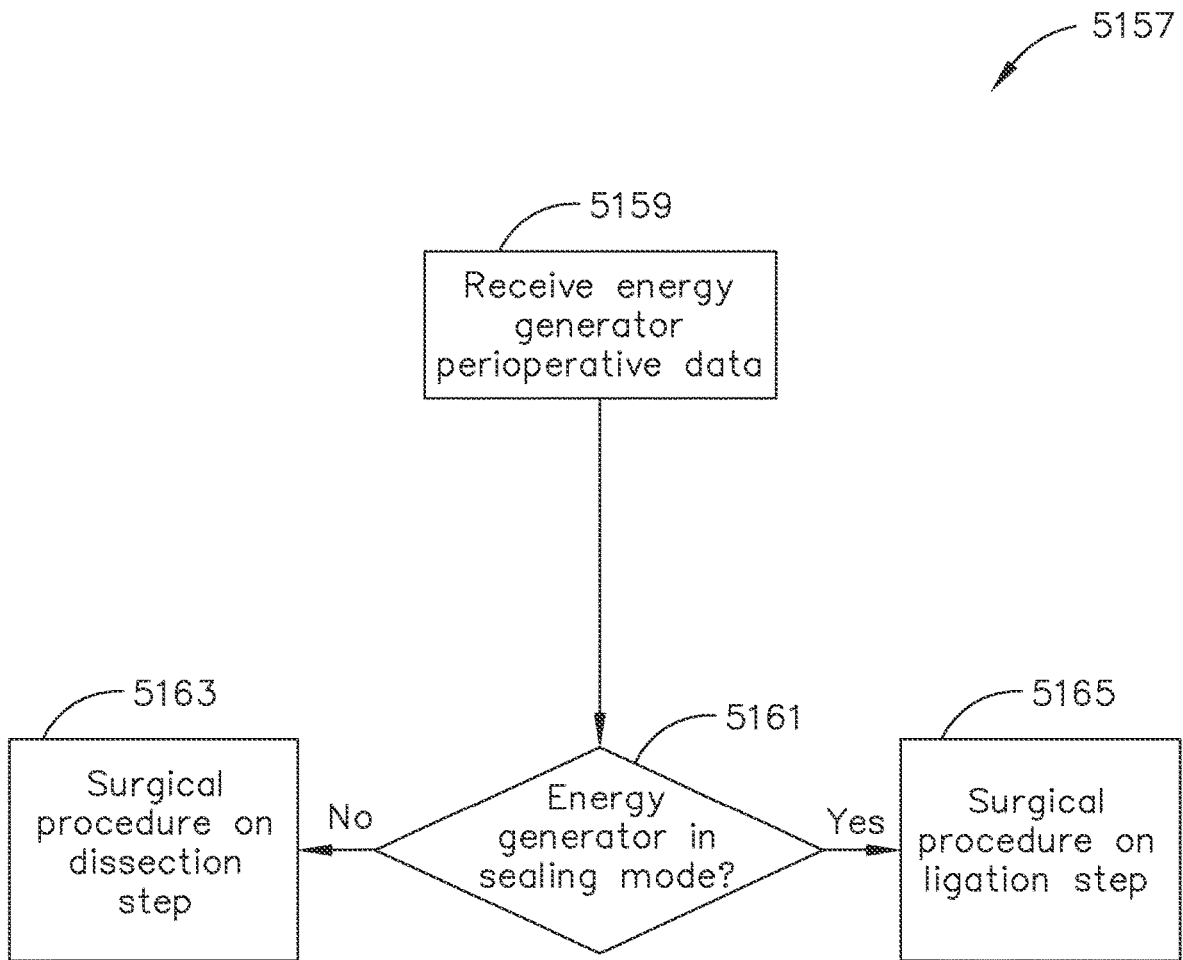


FIG. 84E



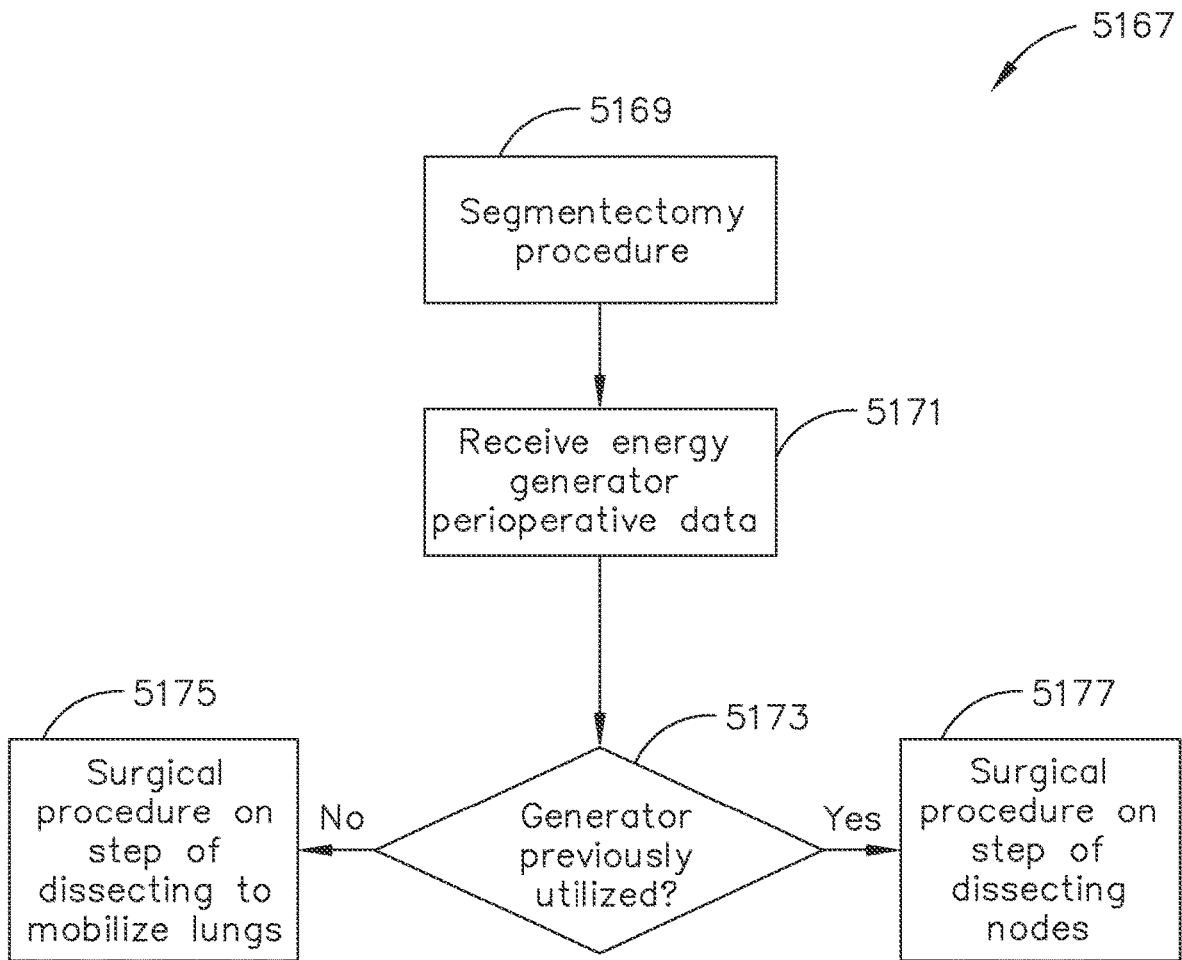


FIG. 84F

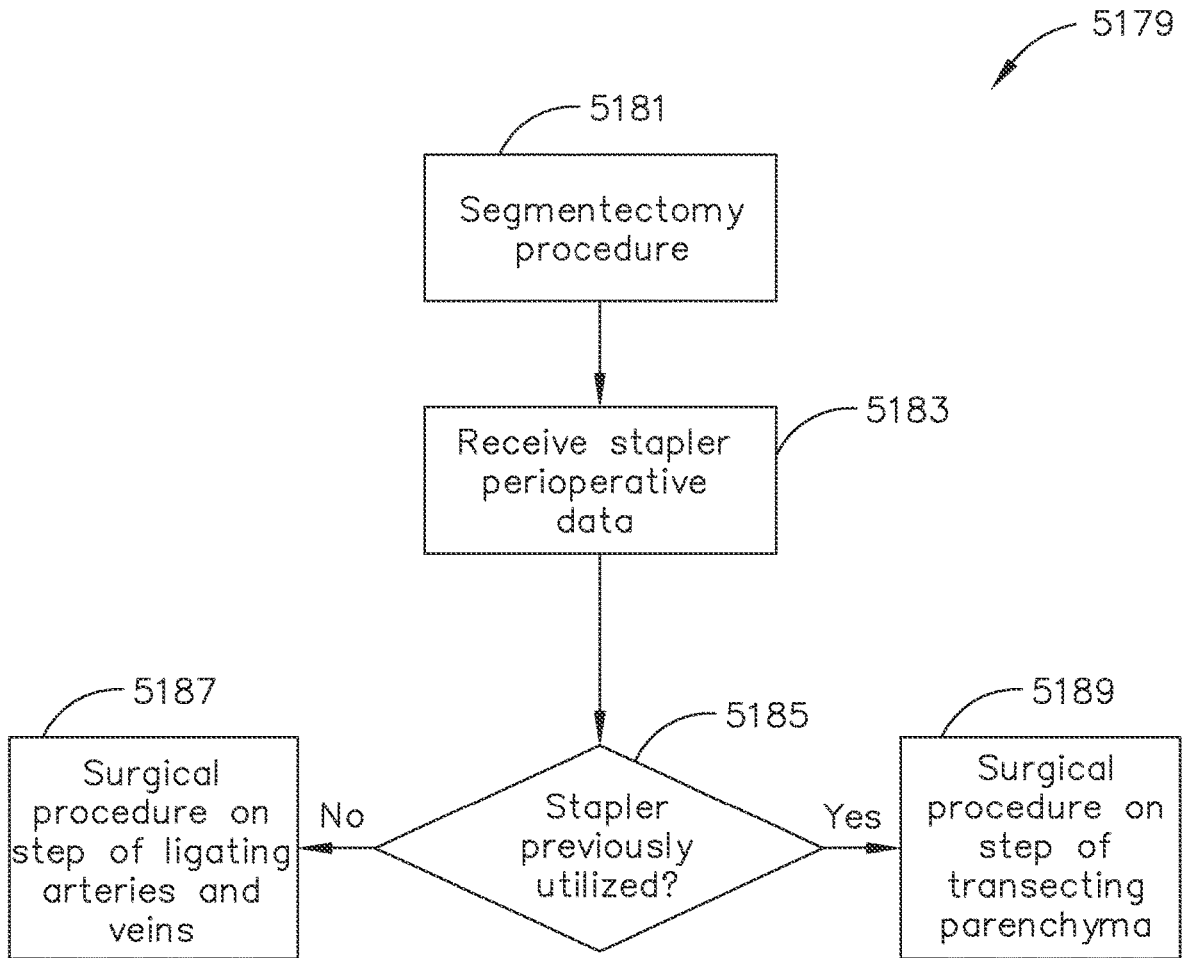


FIG. 84G

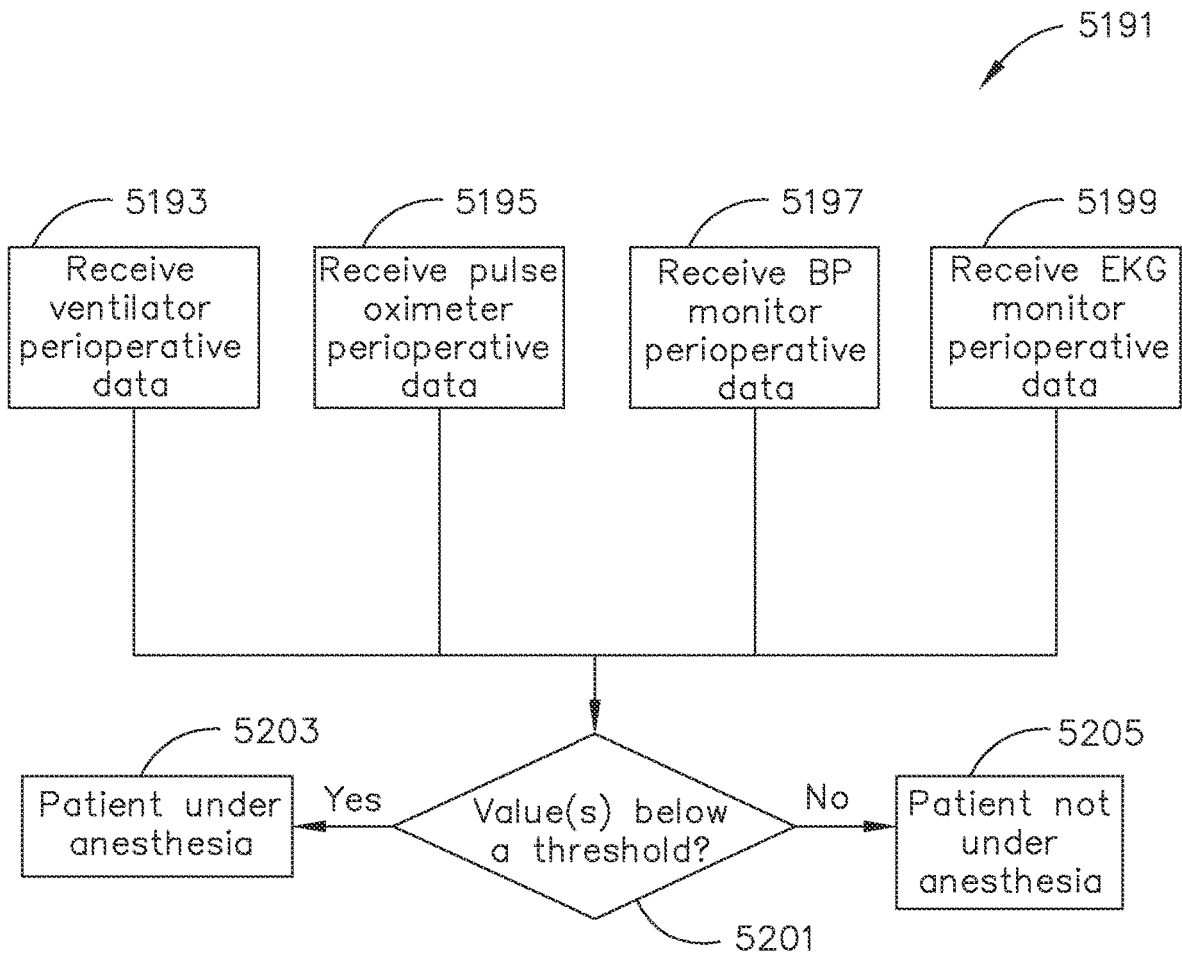


FIG. 84H

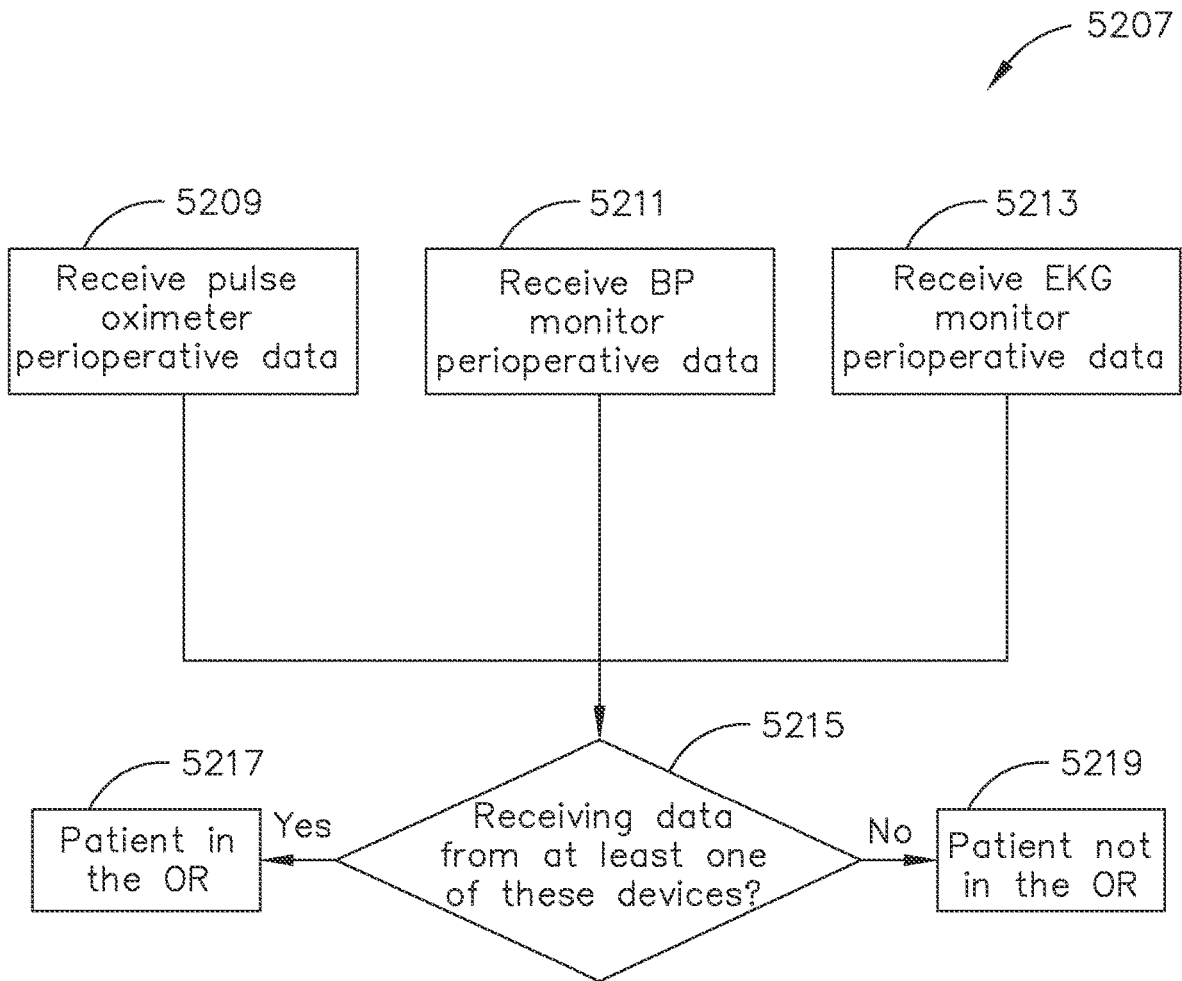


FIG. 84I

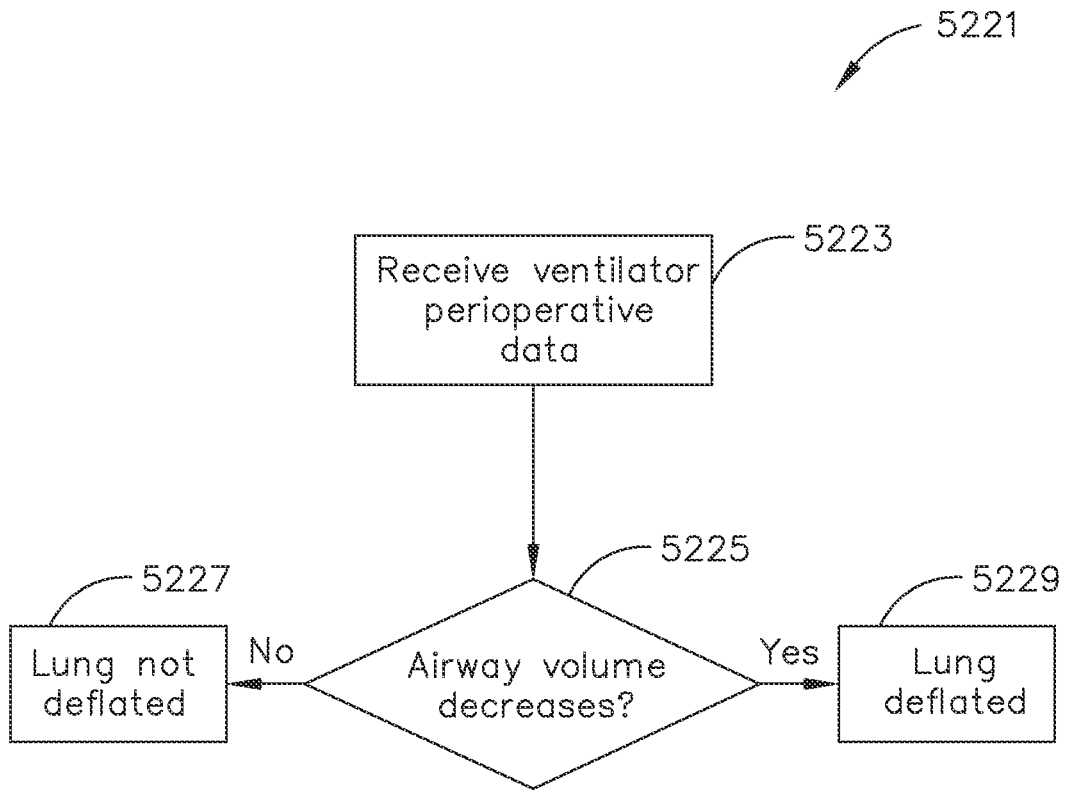


FIG. 84J

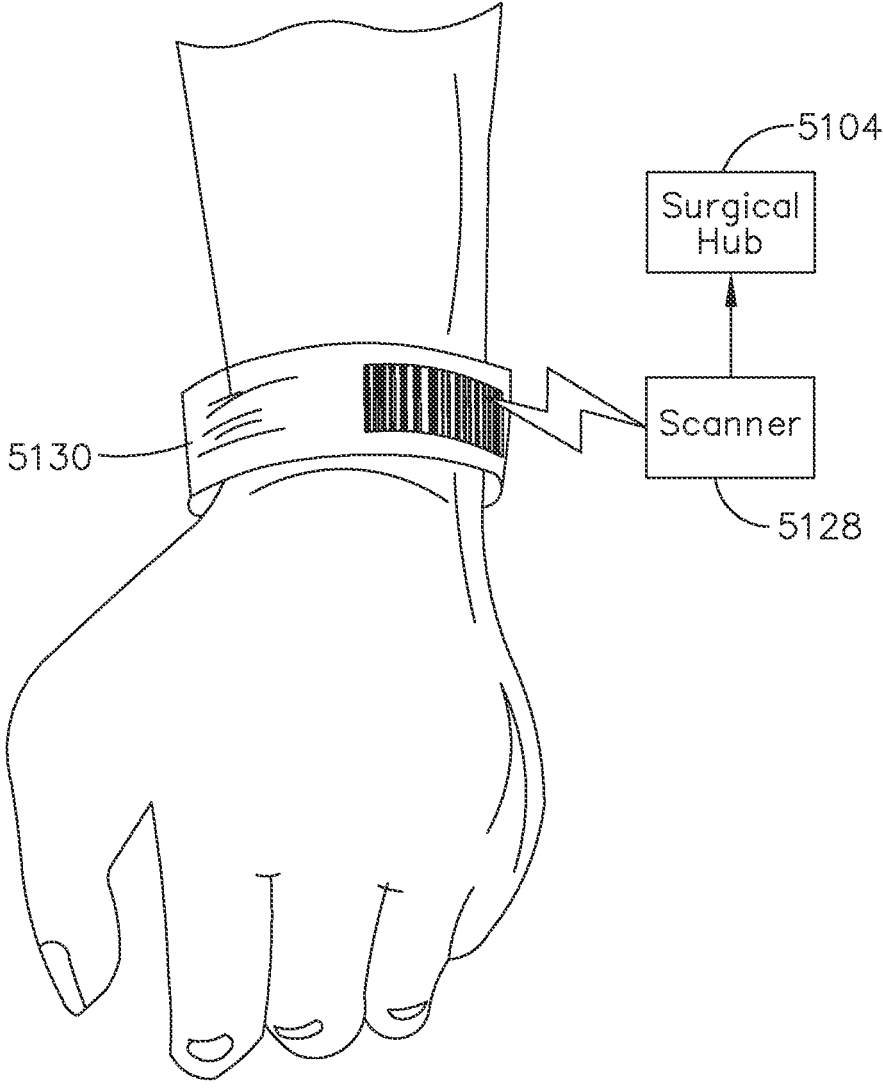


FIG. 85A

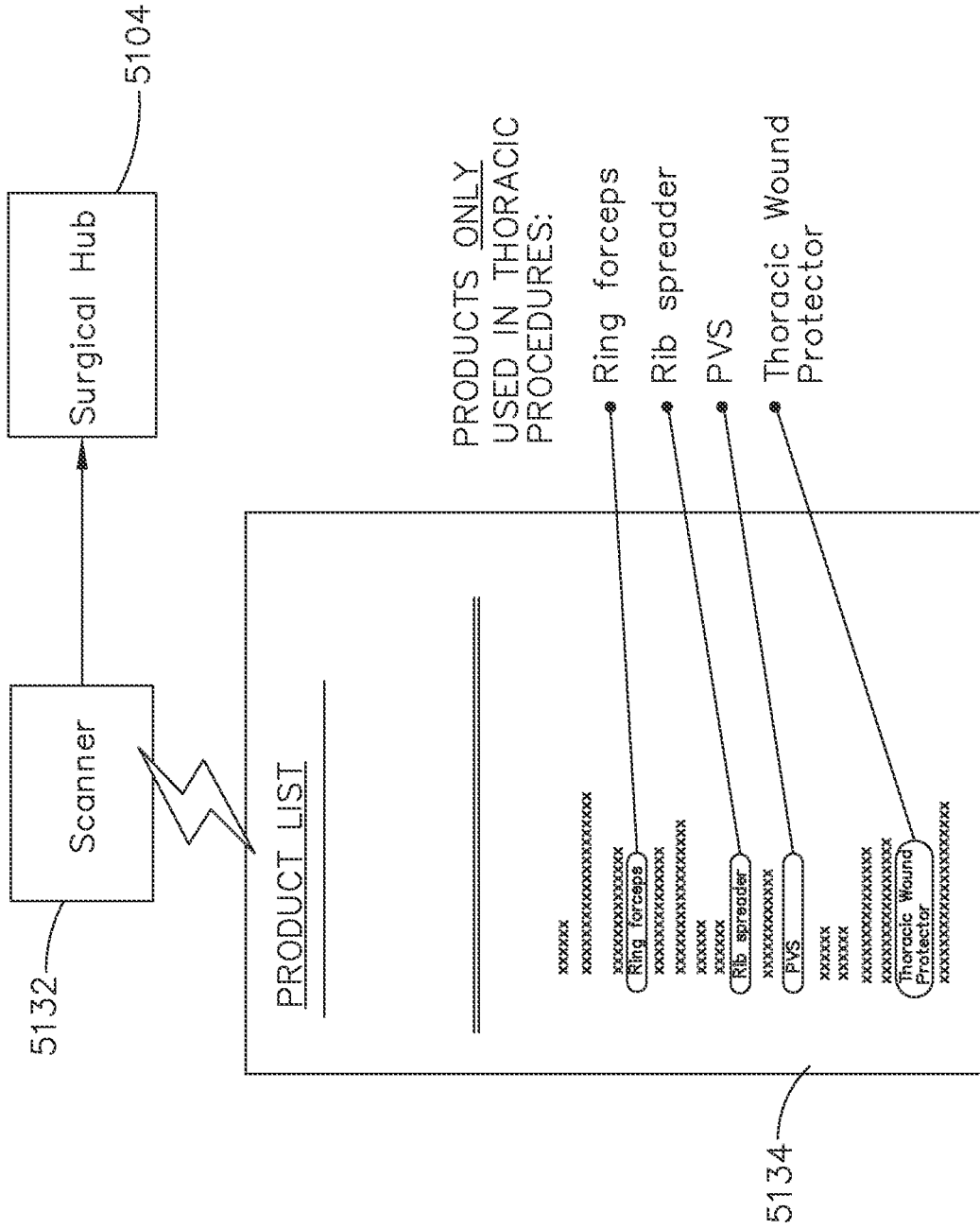


FIG. 85B

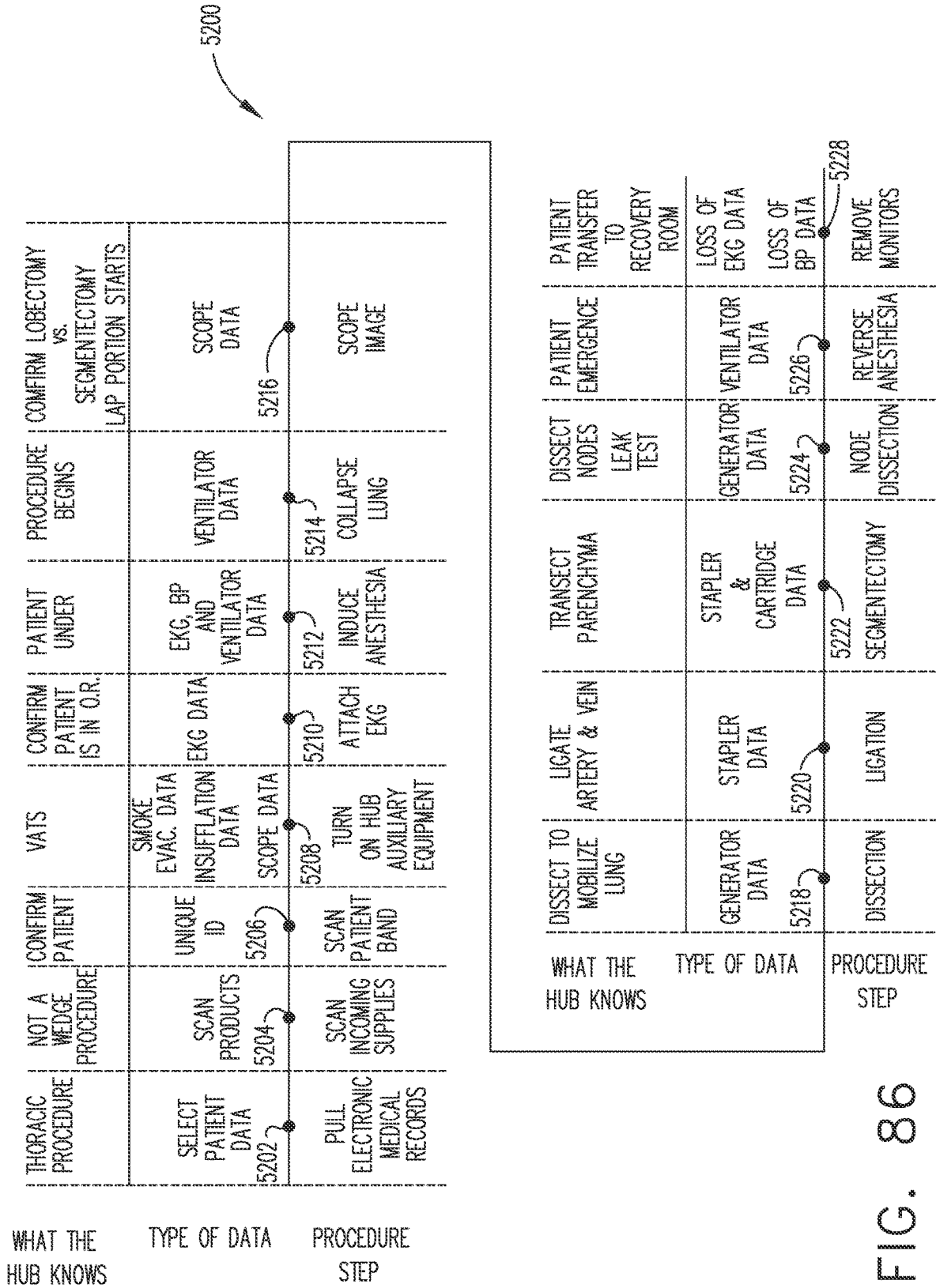


FIG. 86



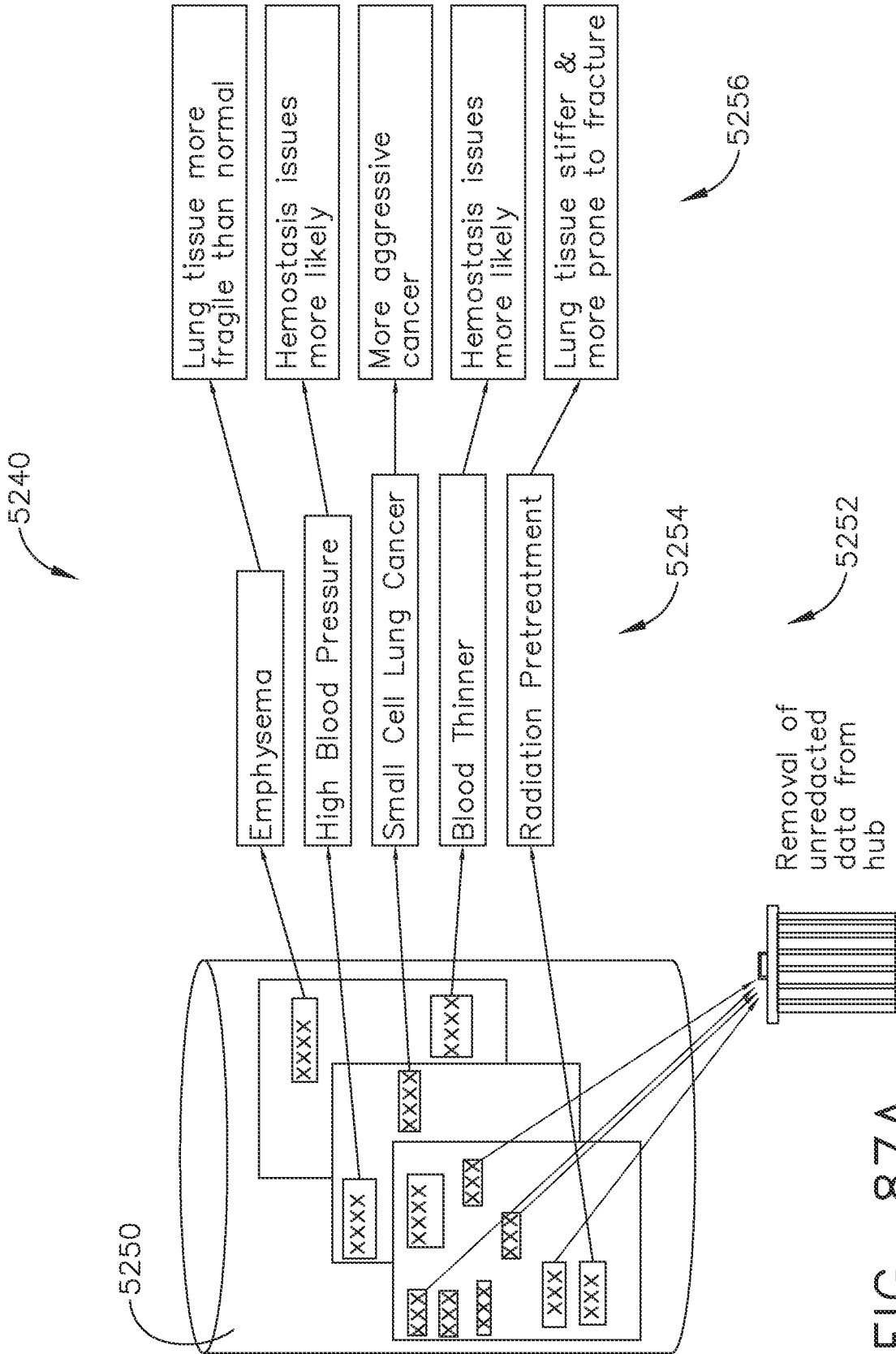


FIG. 87A

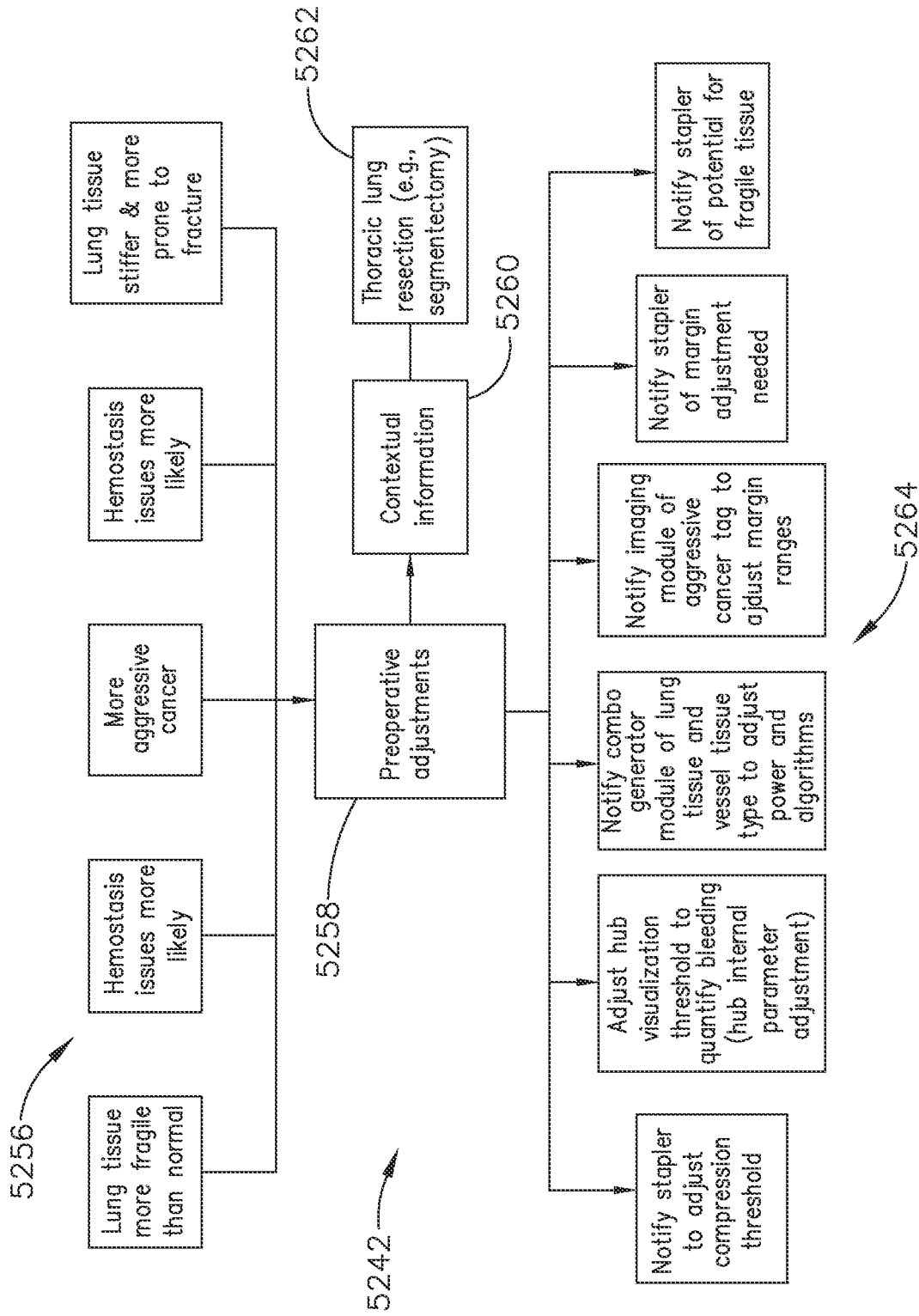


FIG. 87B

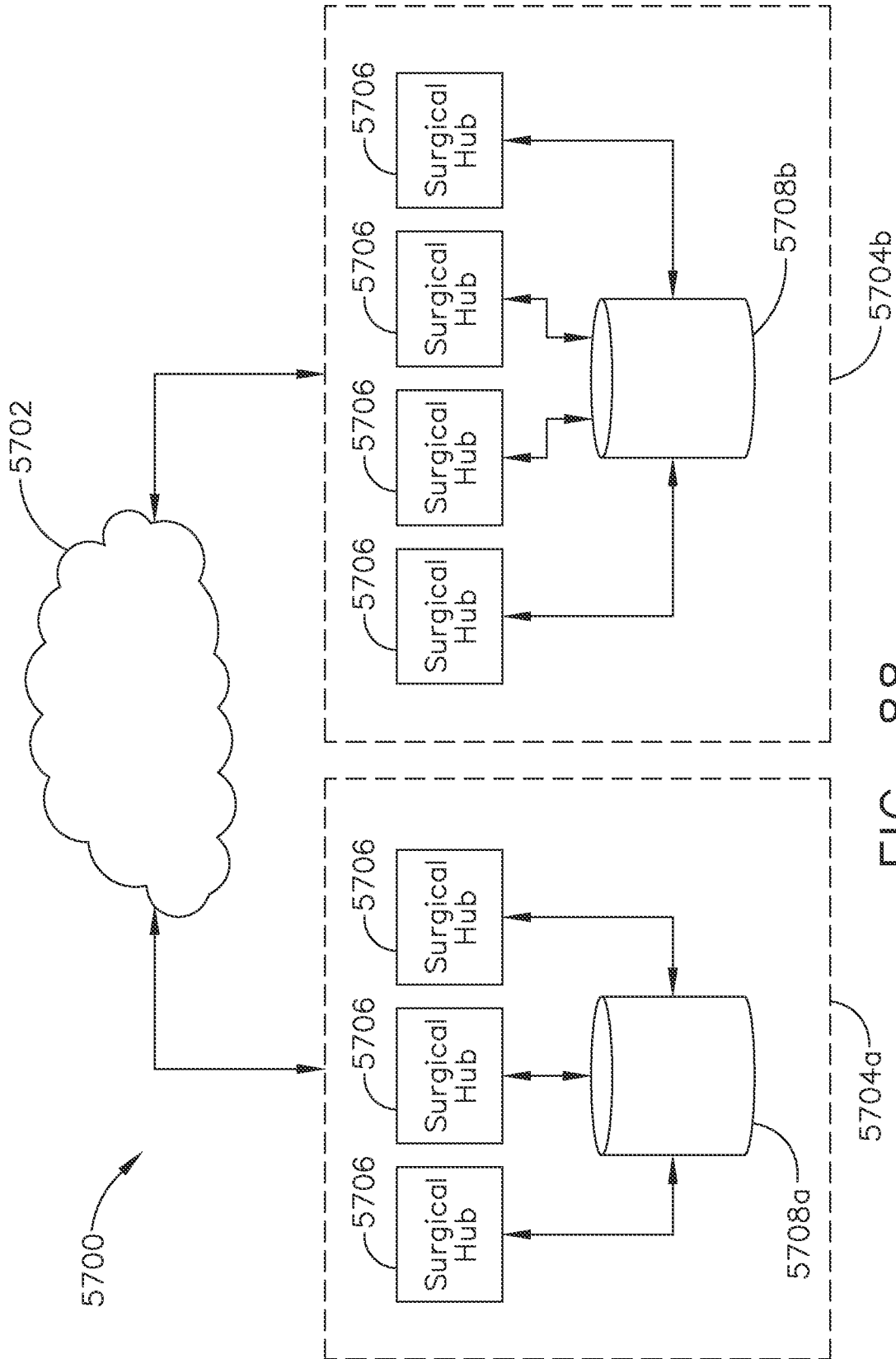


FIG. 88

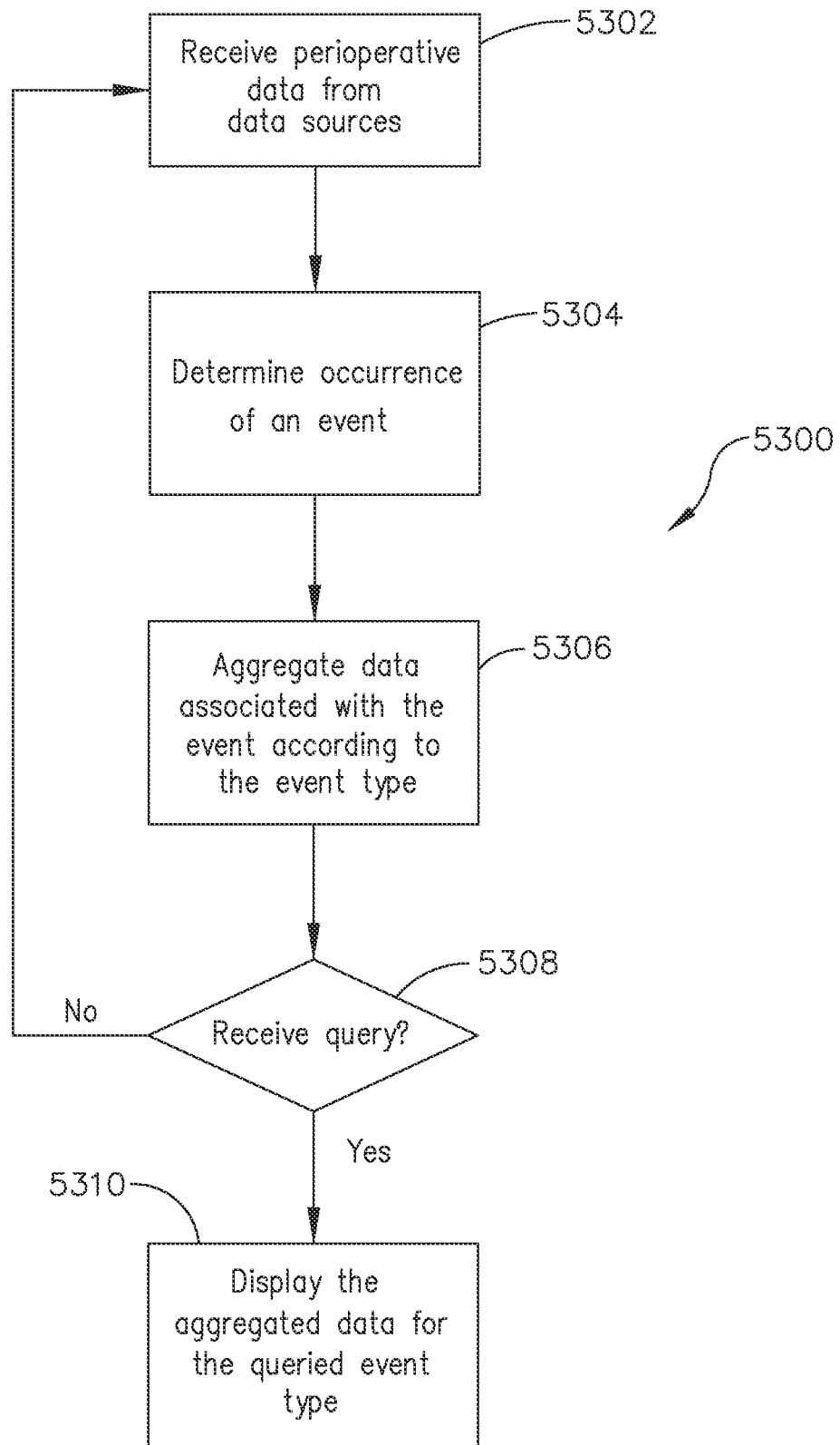


FIG. 89

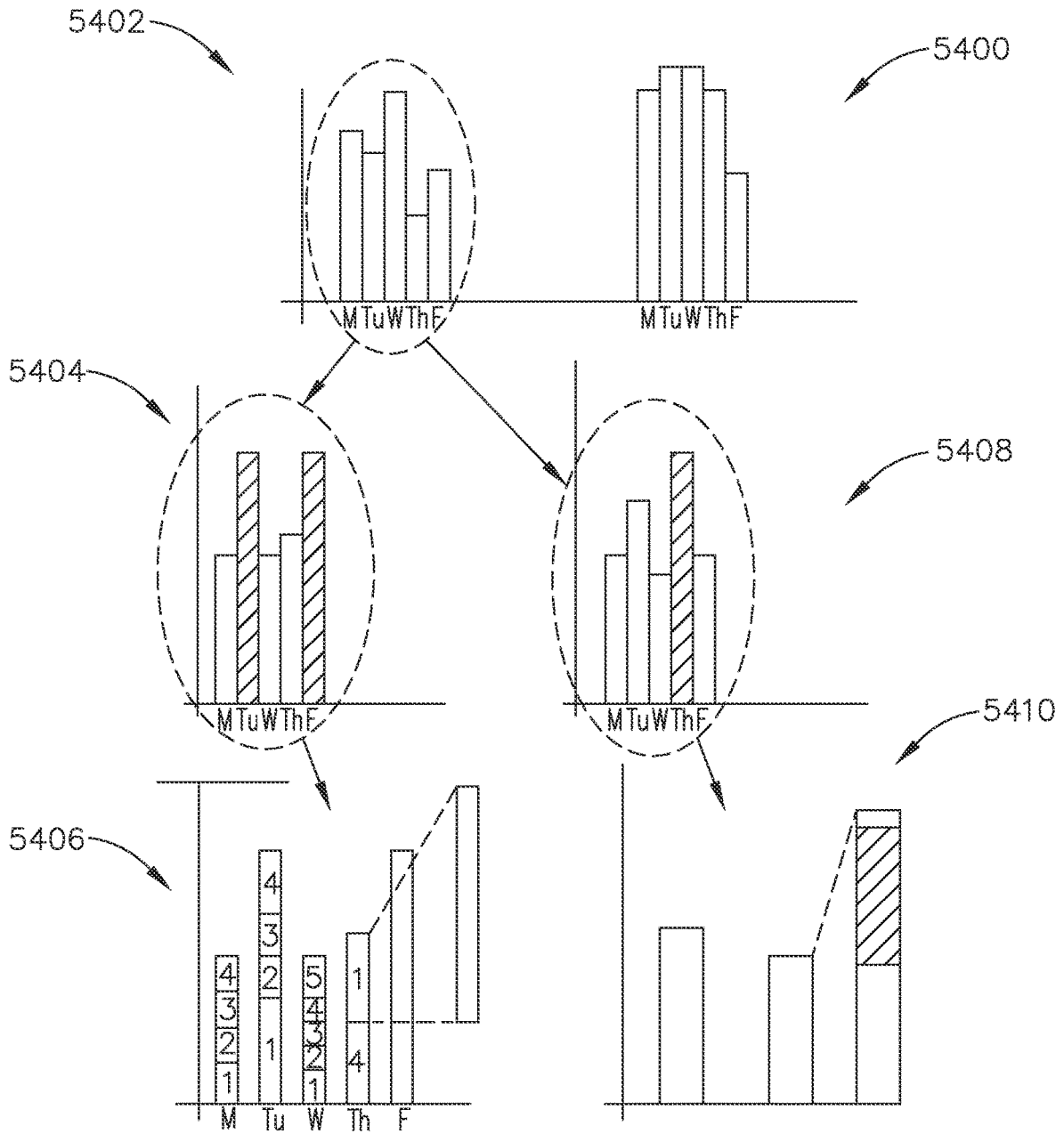


FIG. 90

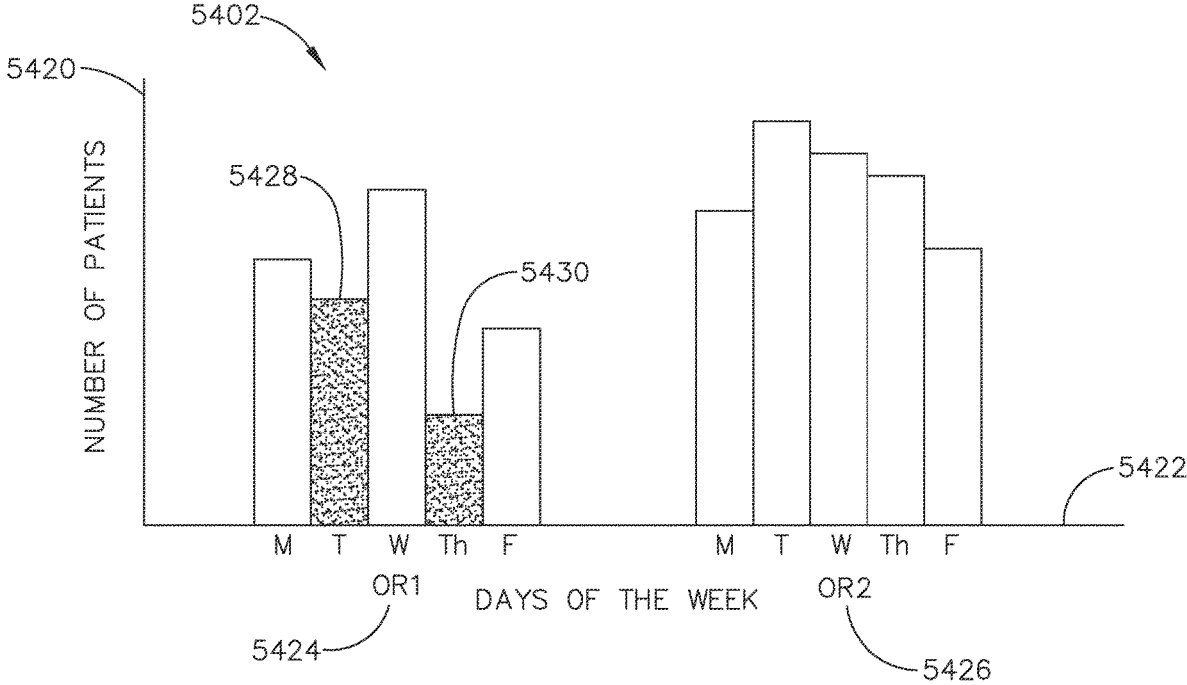


FIG. 91

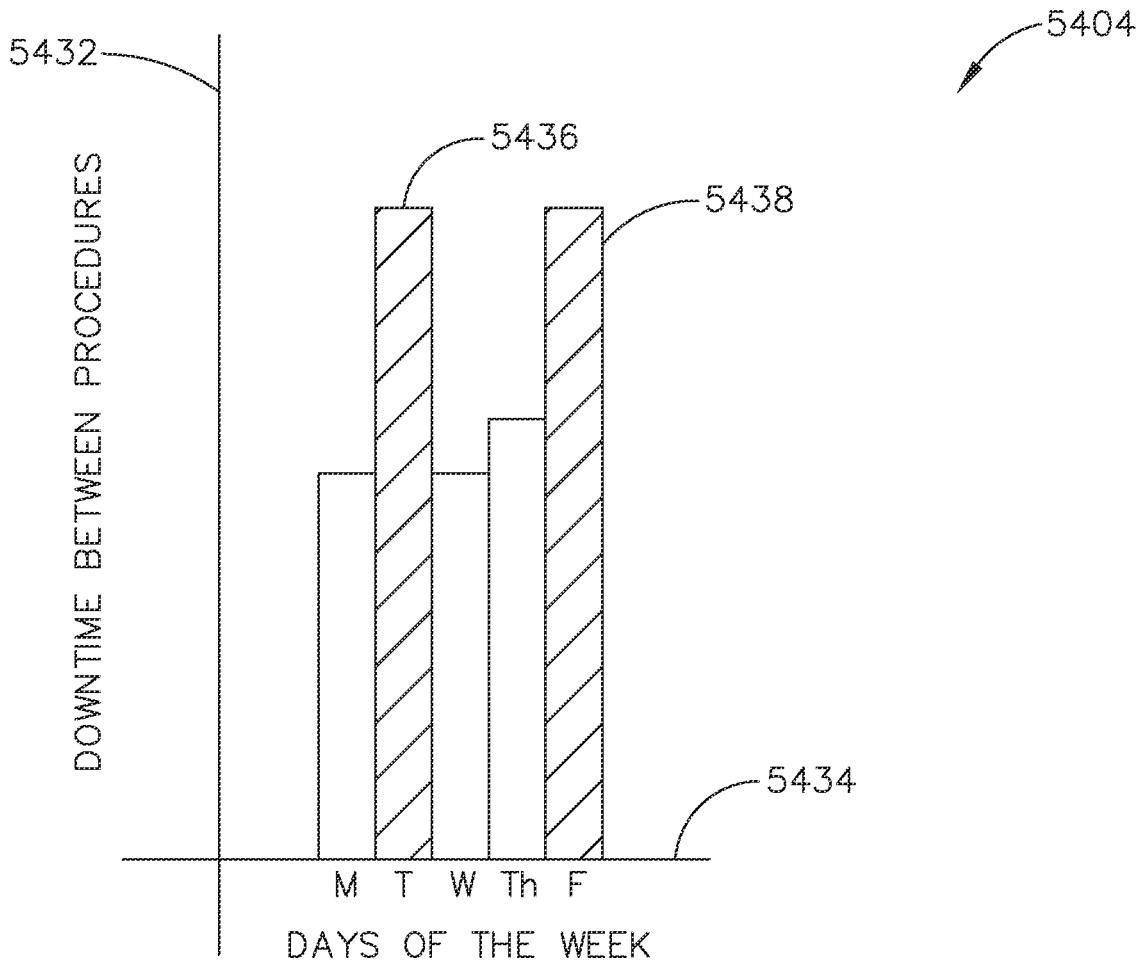


FIG. 92

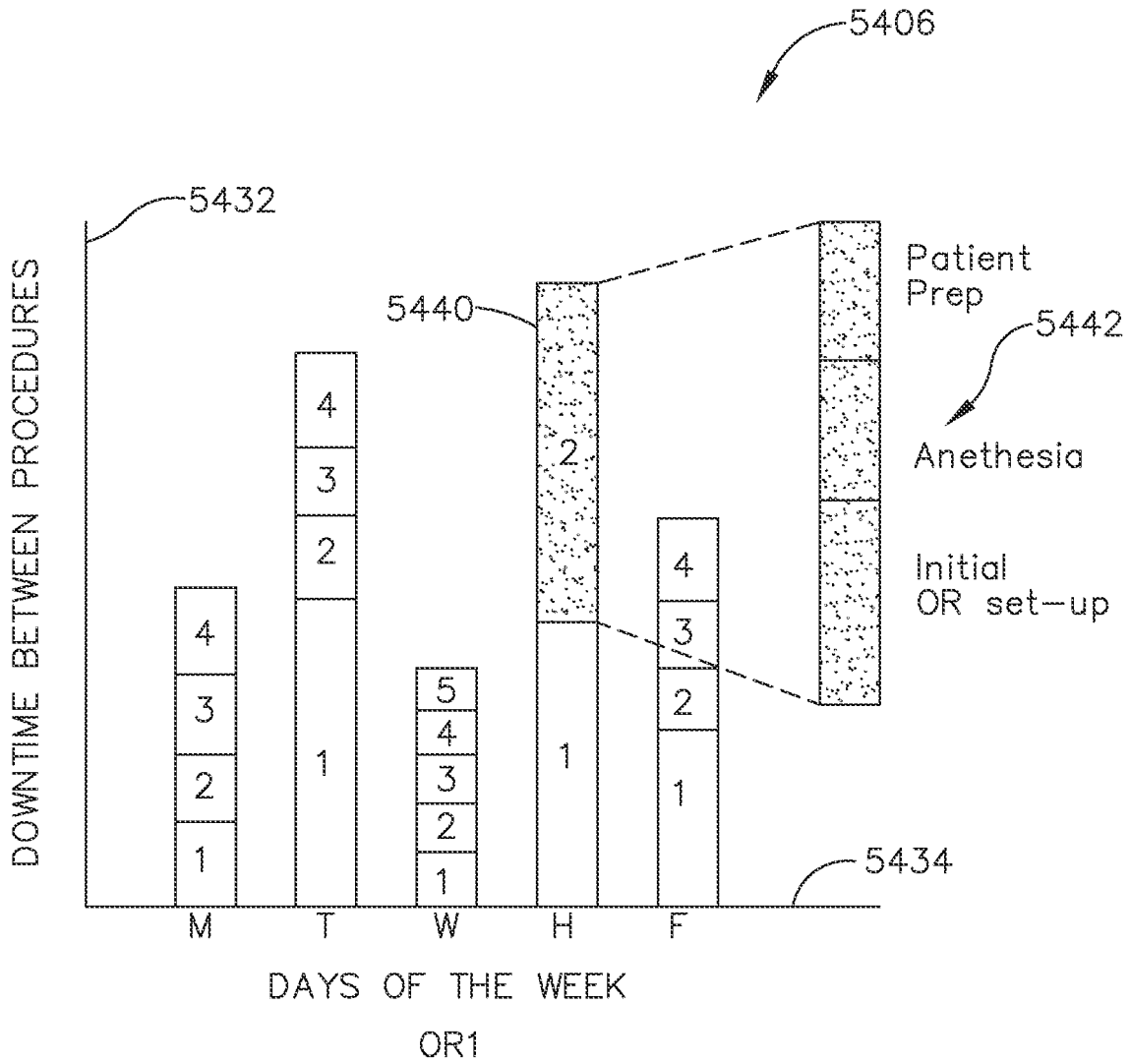


FIG. 93



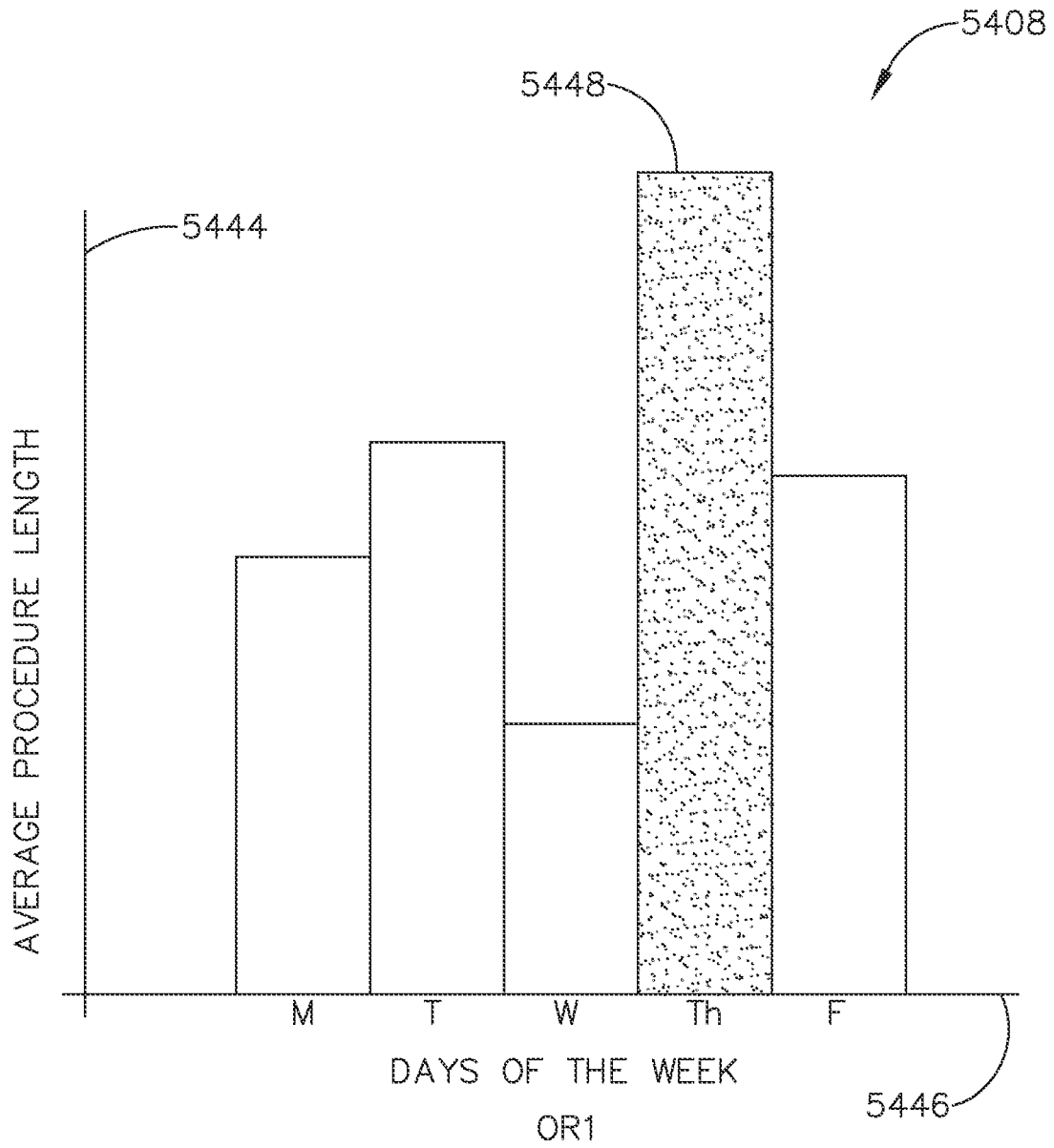


FIG. 94

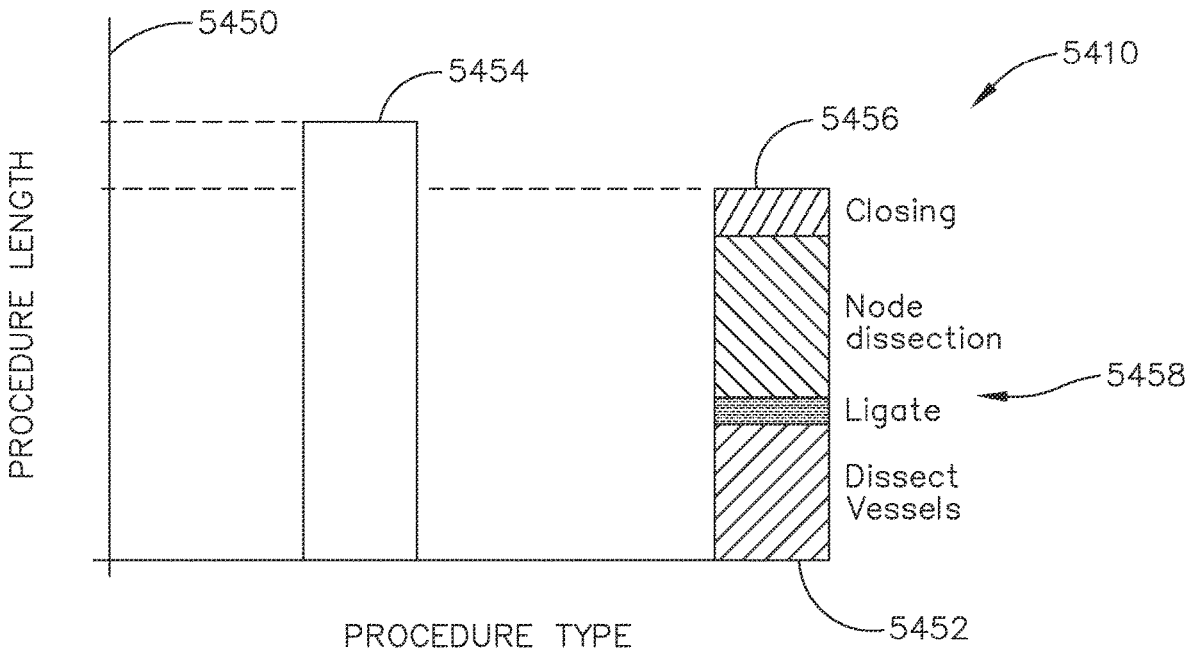


FIG. 95

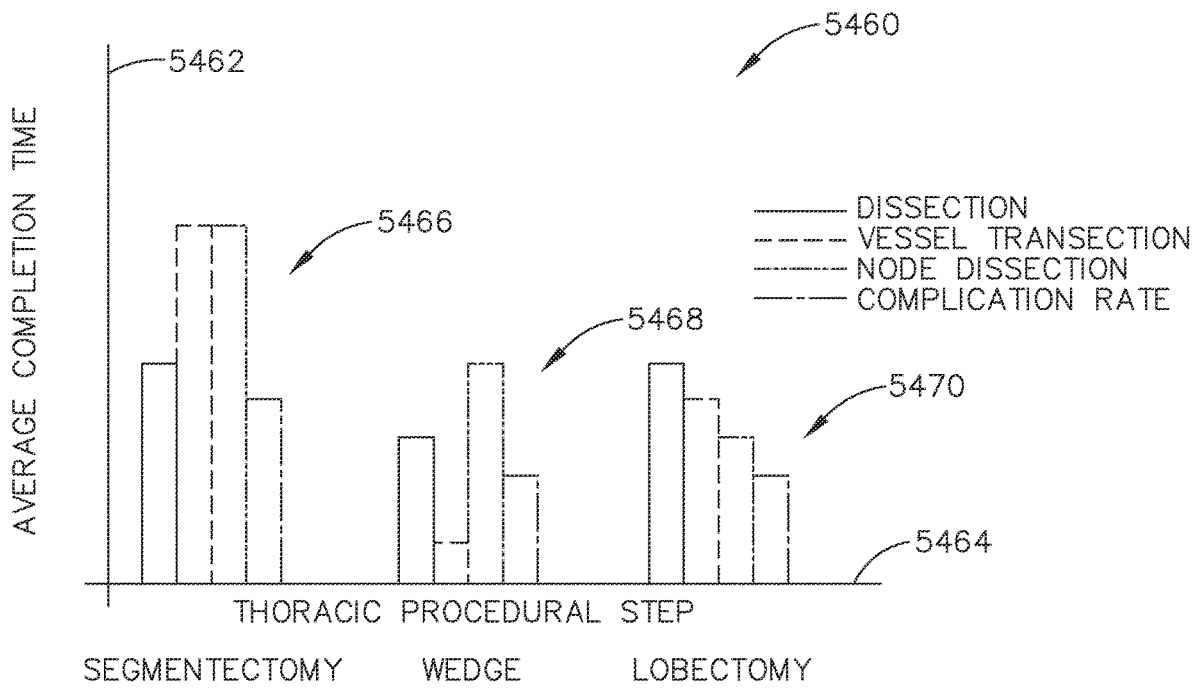


FIG. 96

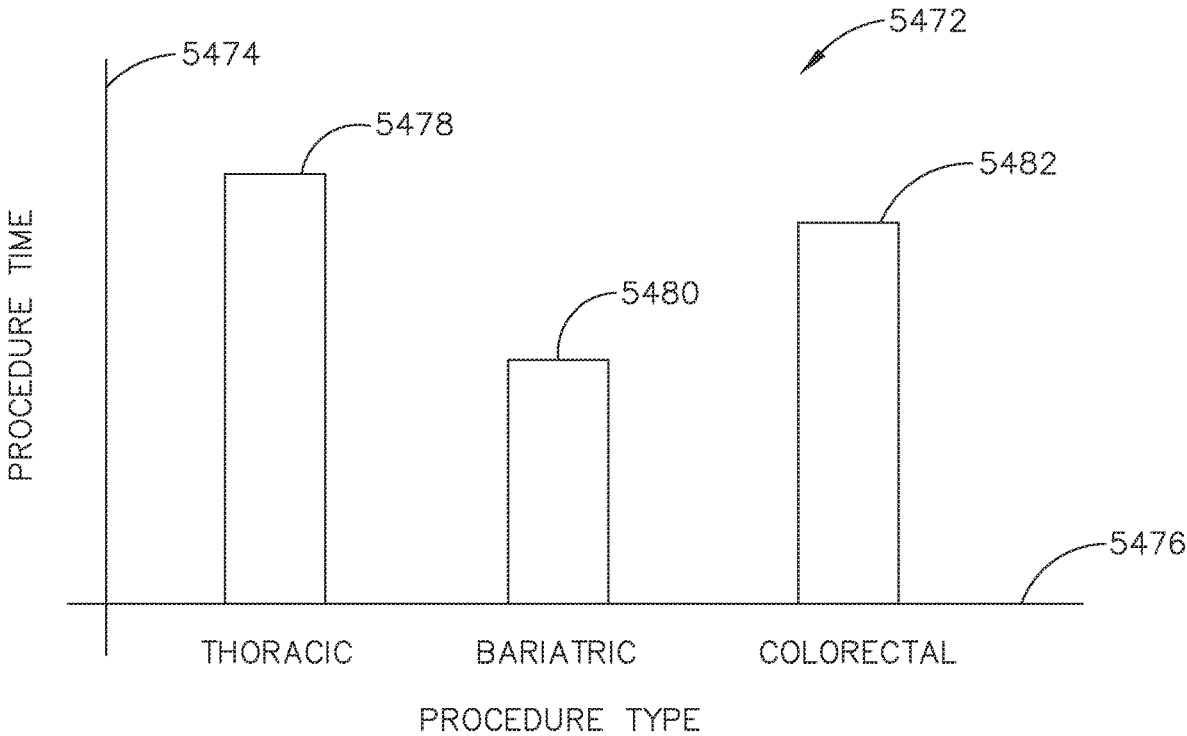


FIG. 97

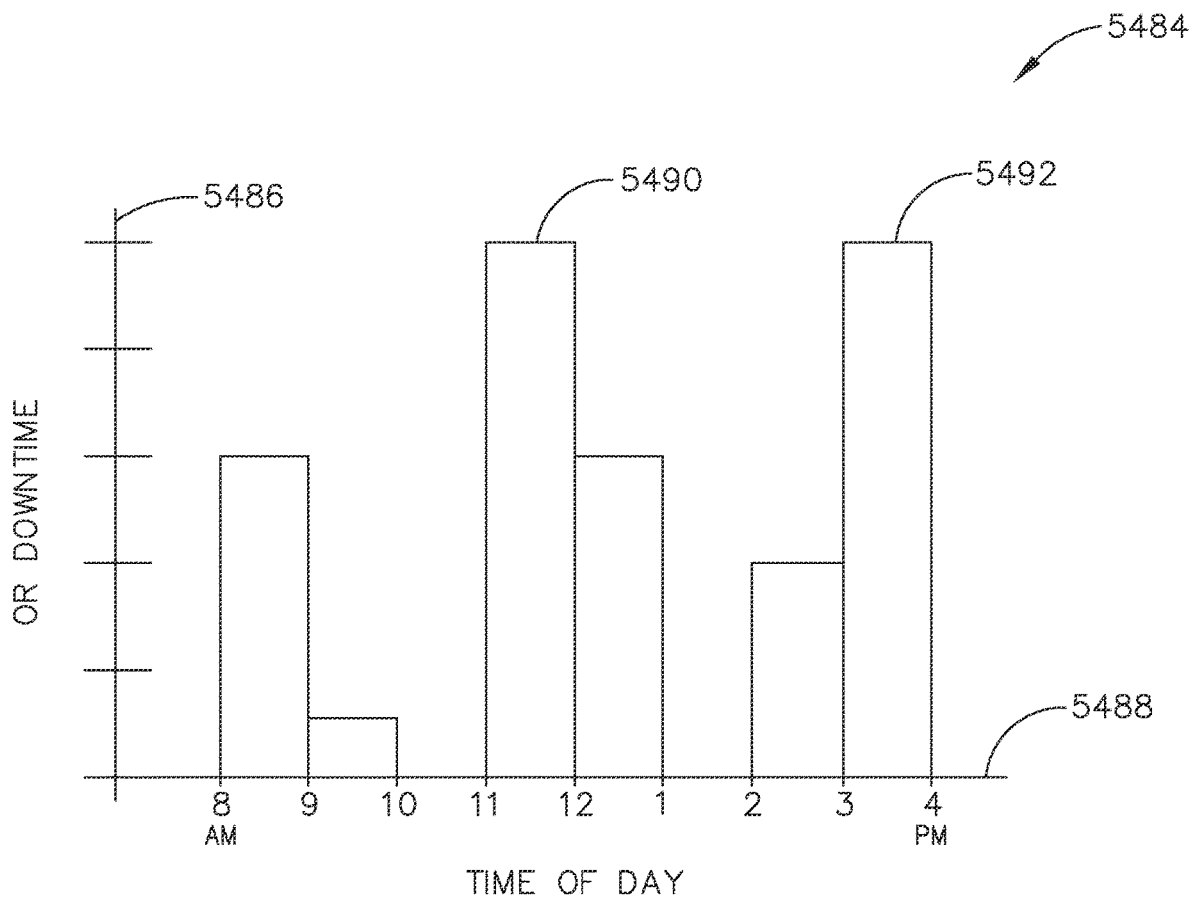


FIG. 98

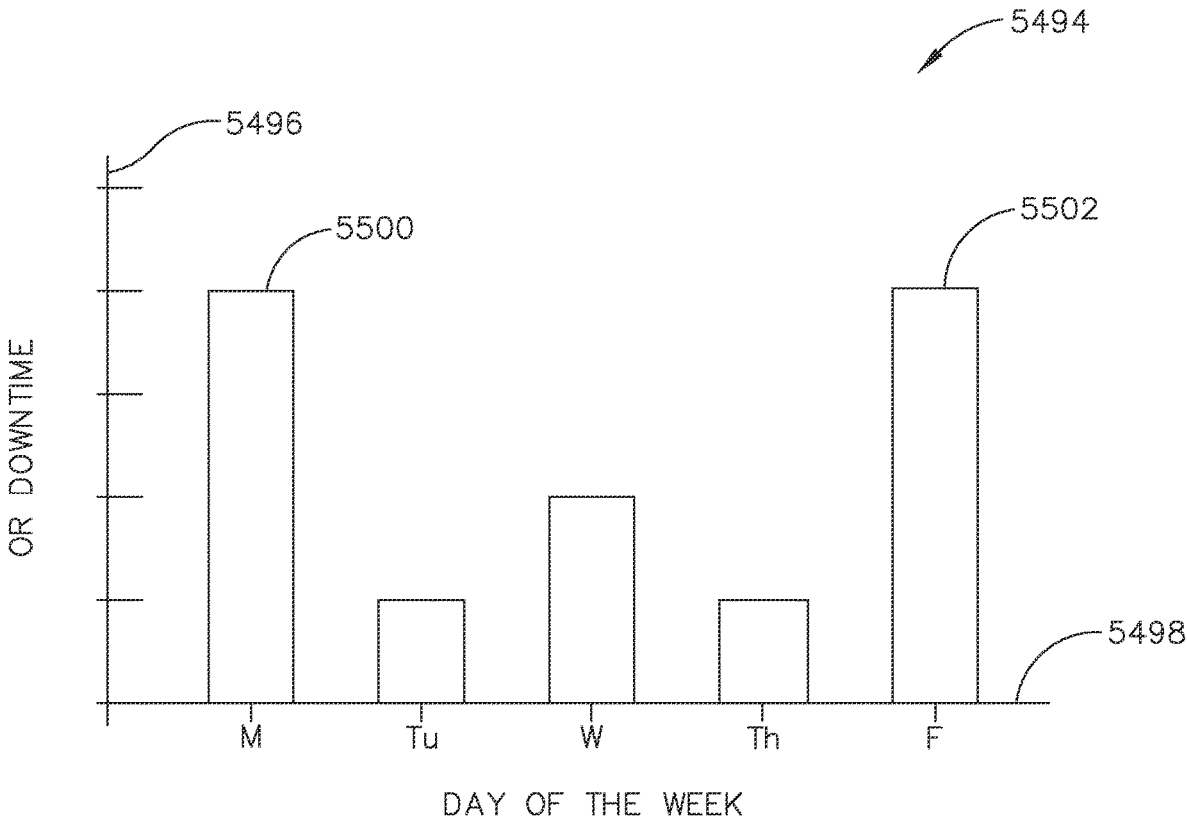


FIG. 99

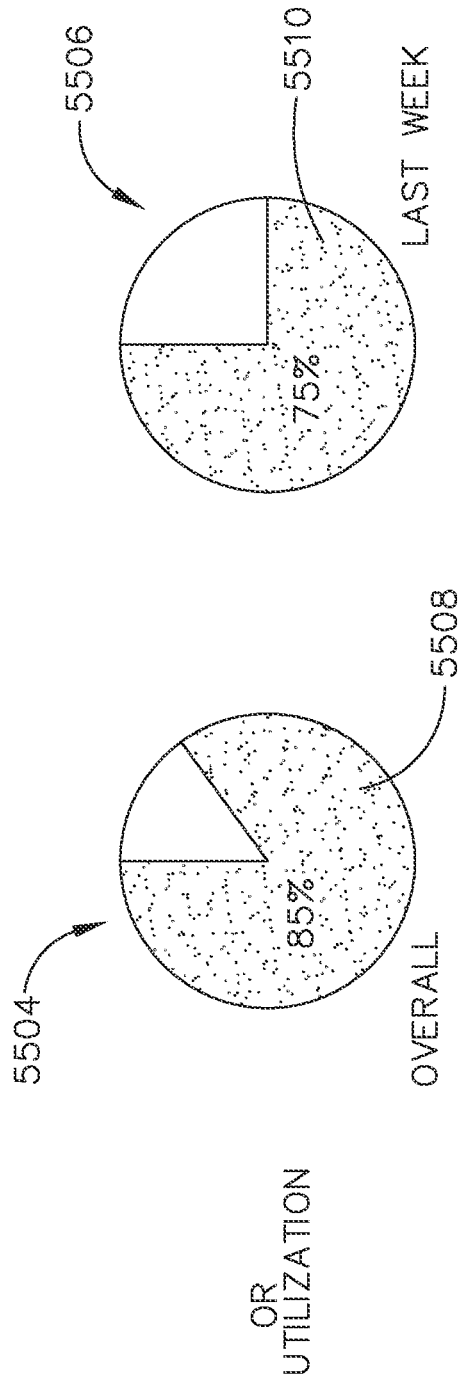


FIG. 100

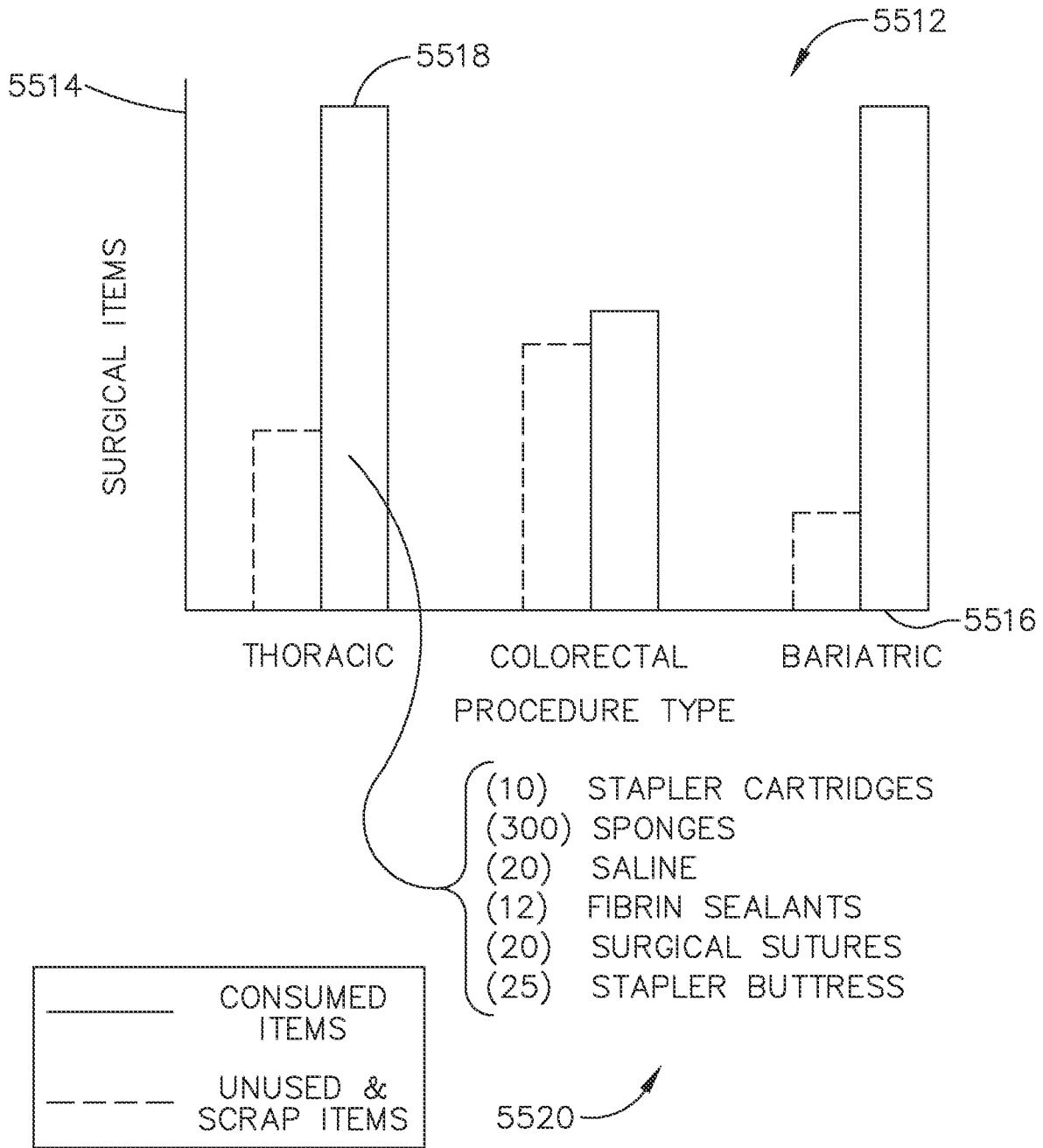


FIG. 101



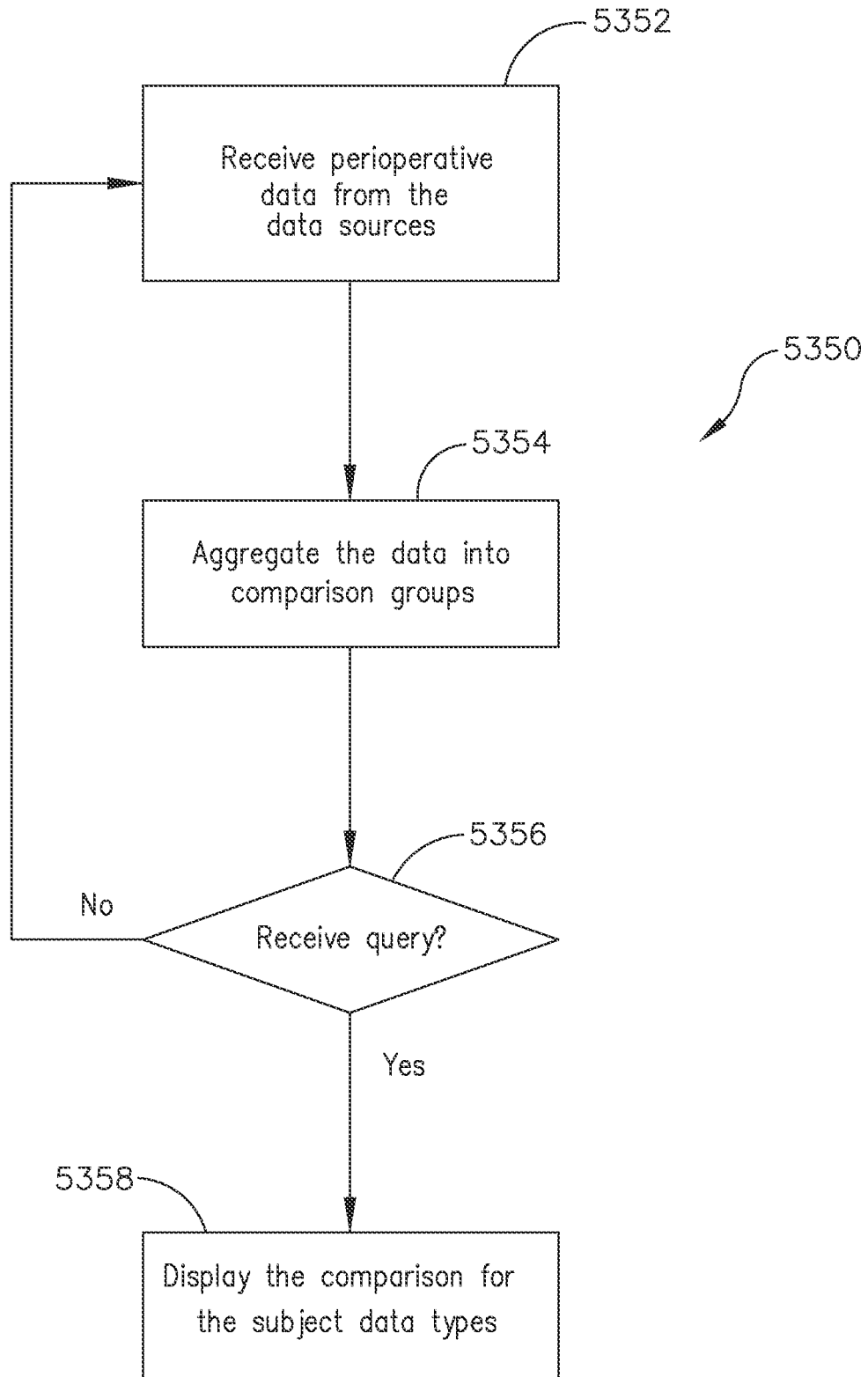


FIG. 102

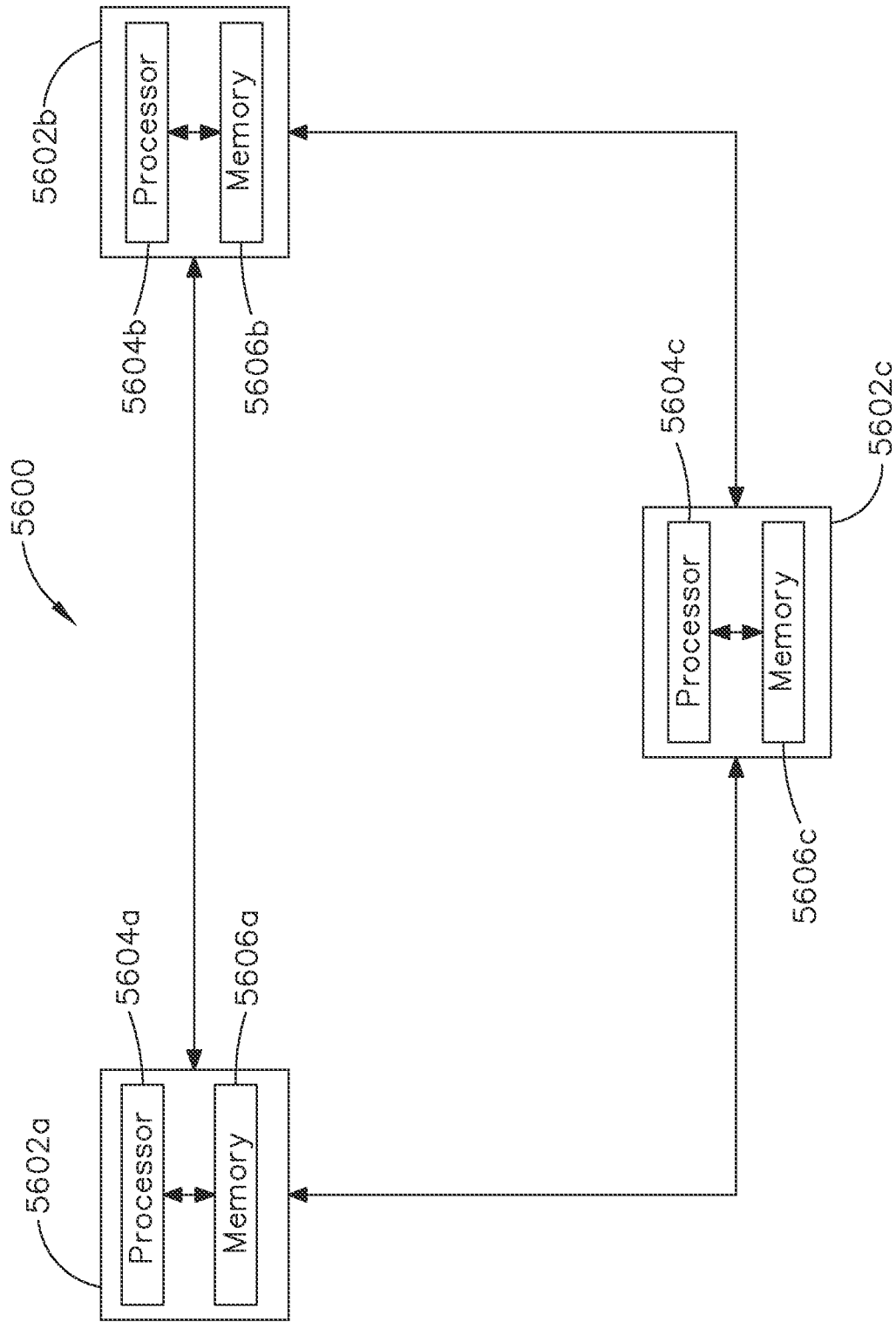


FIG. 103

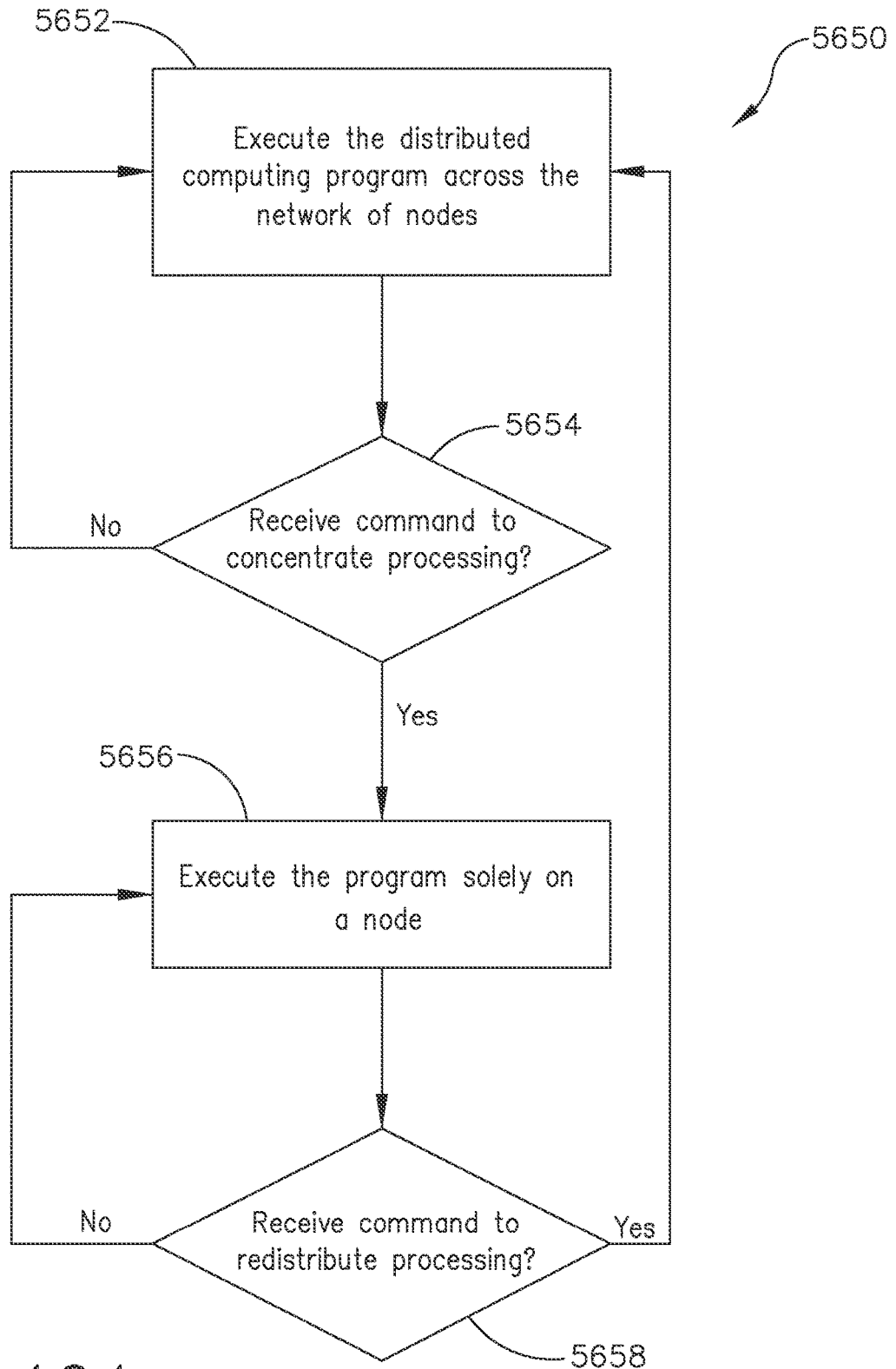


FIG. 104

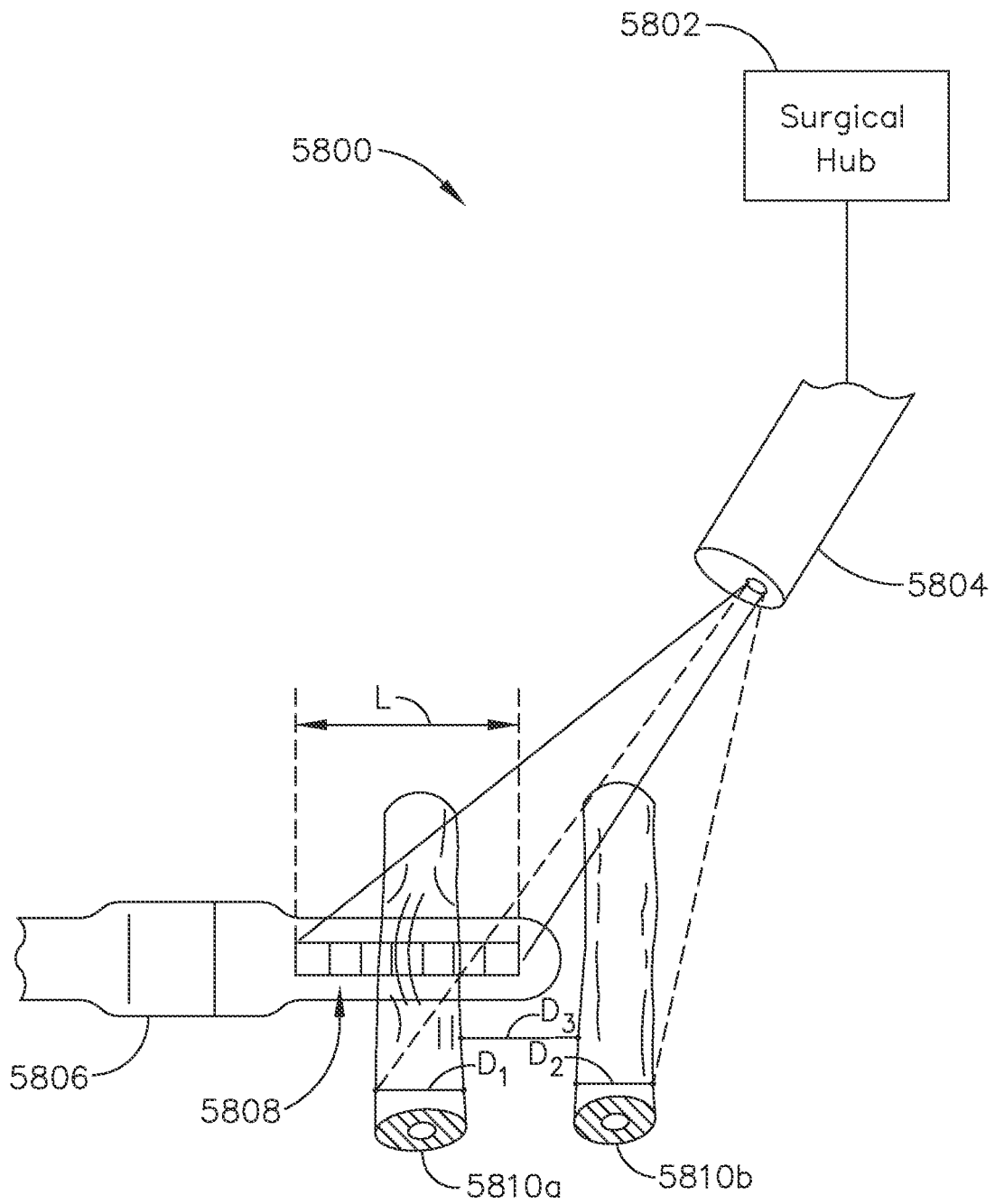


FIG. 105

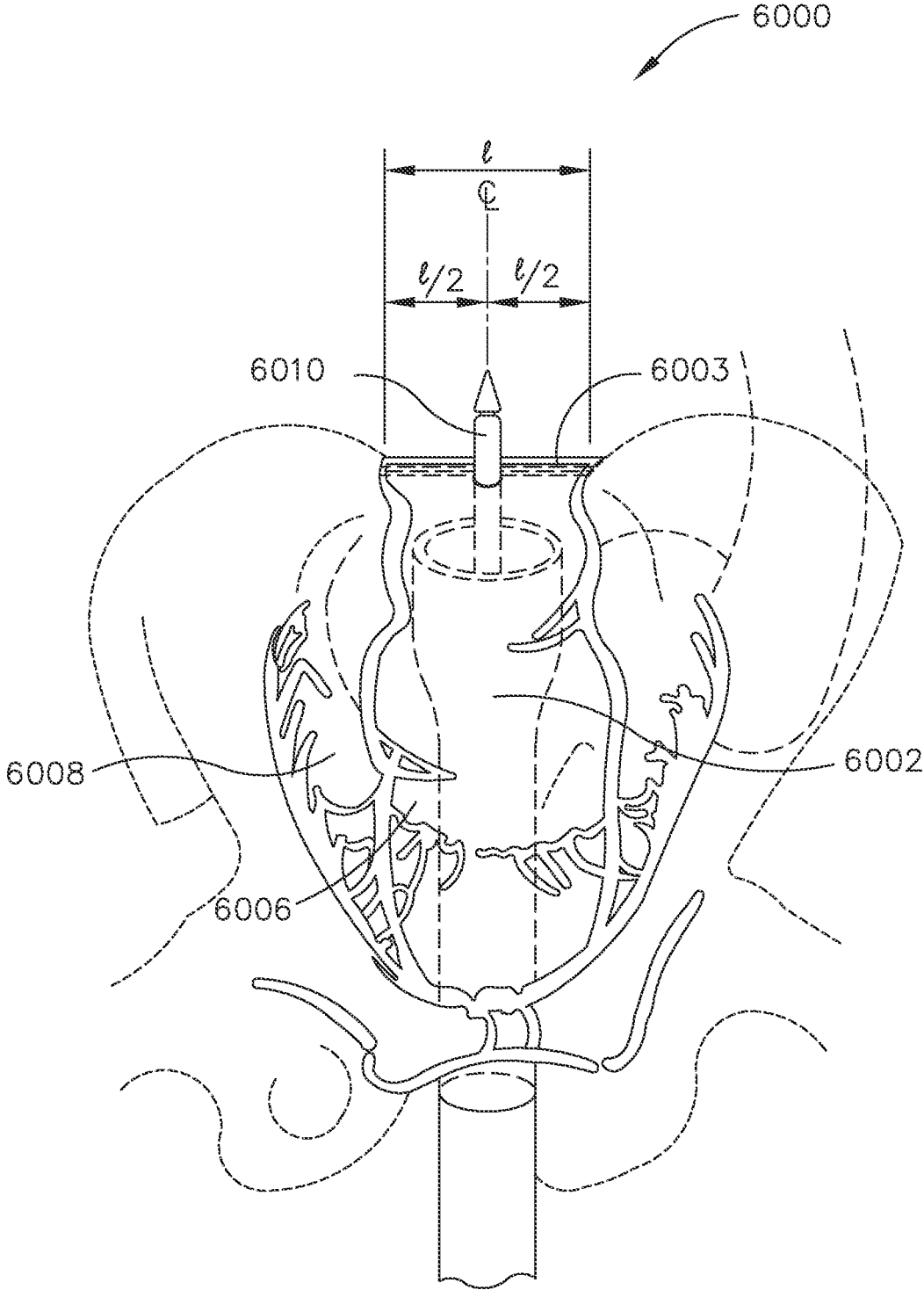


FIG. 106

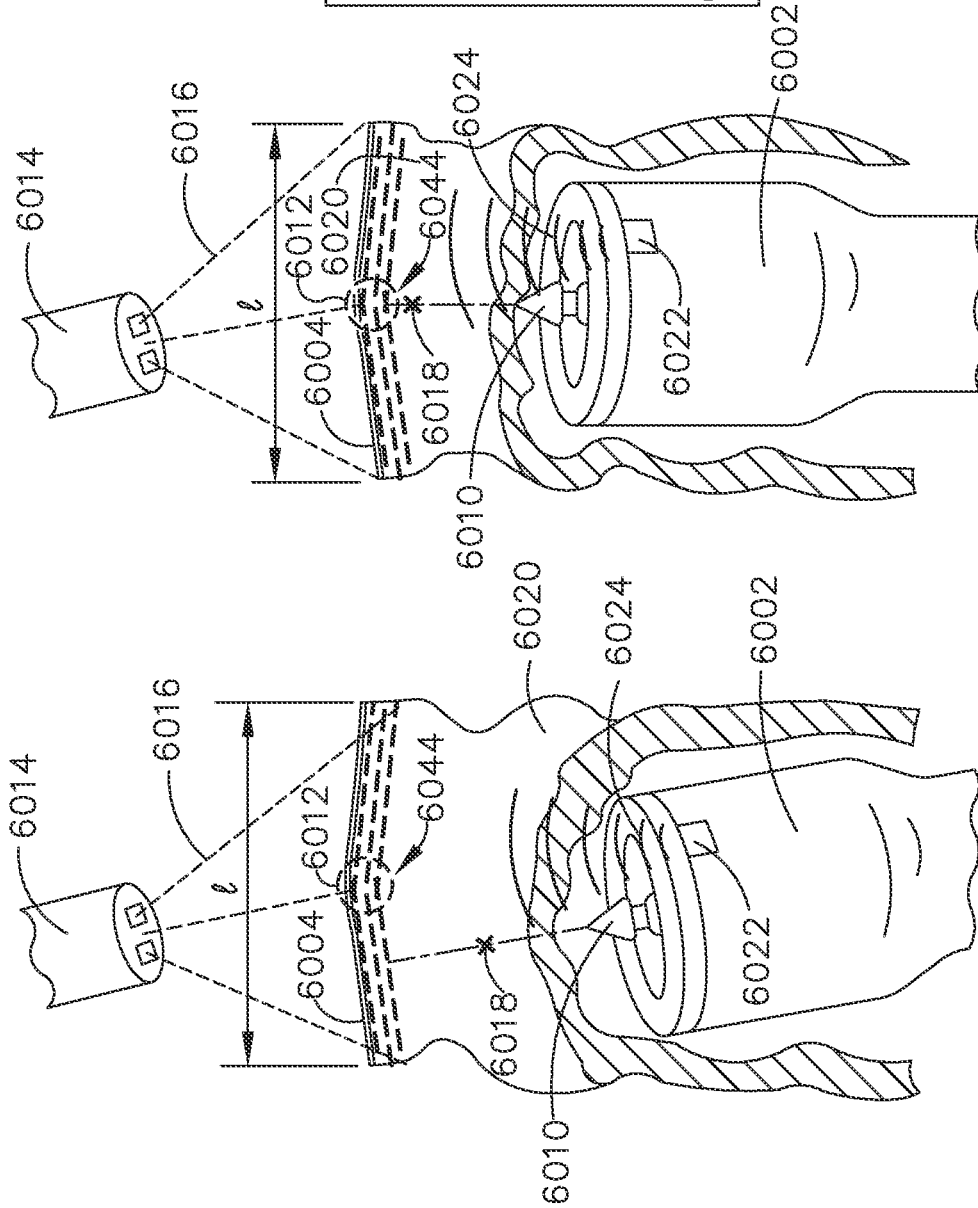


FIG. 107

FIG. 108

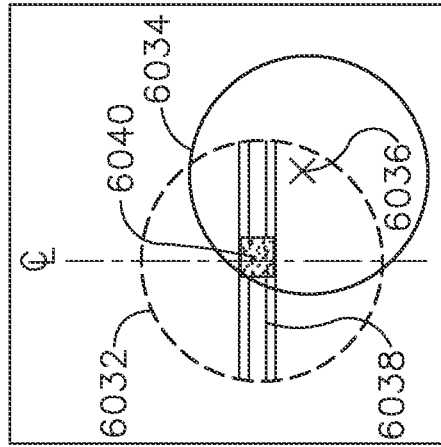


FIG. 109

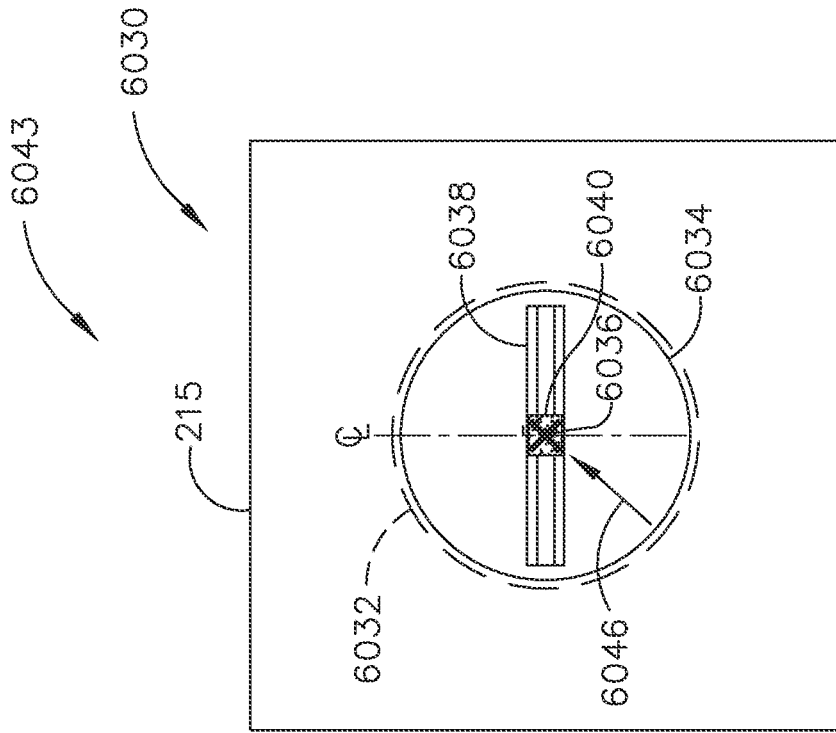


FIG. 110

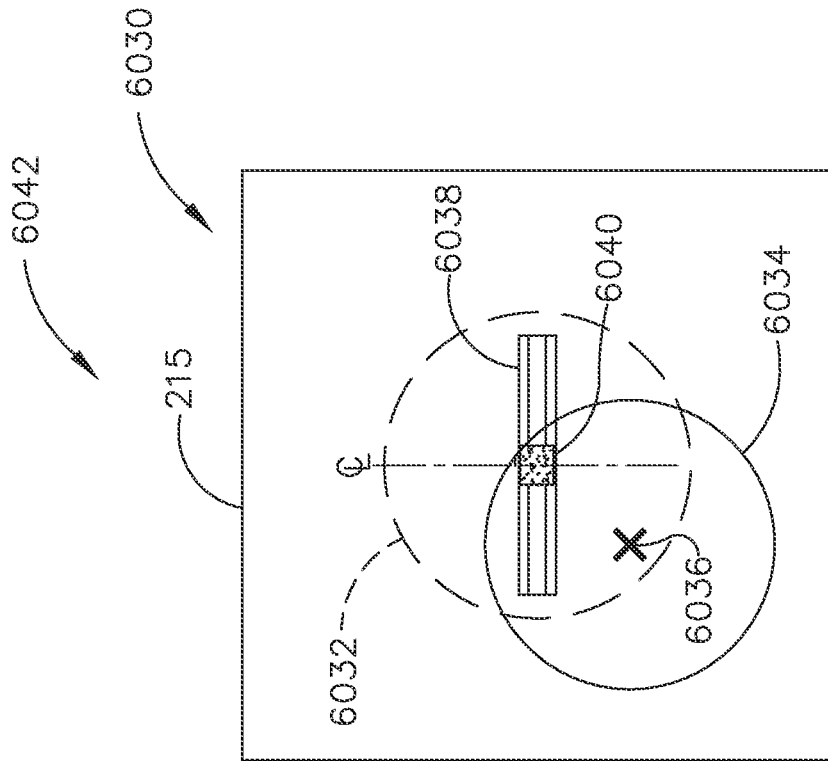


FIG. 111

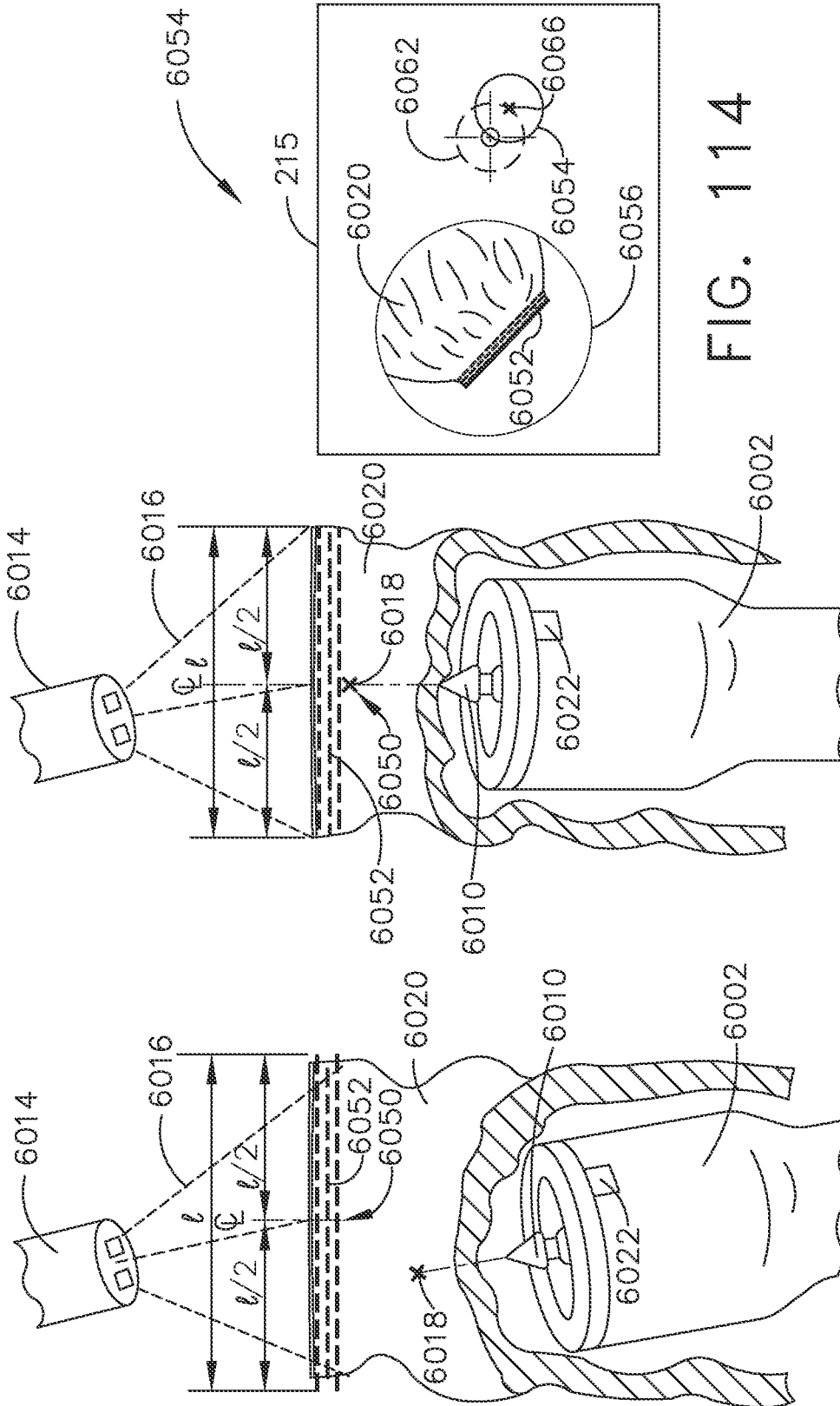


FIG. 112

FIG. 113

FIG. 114



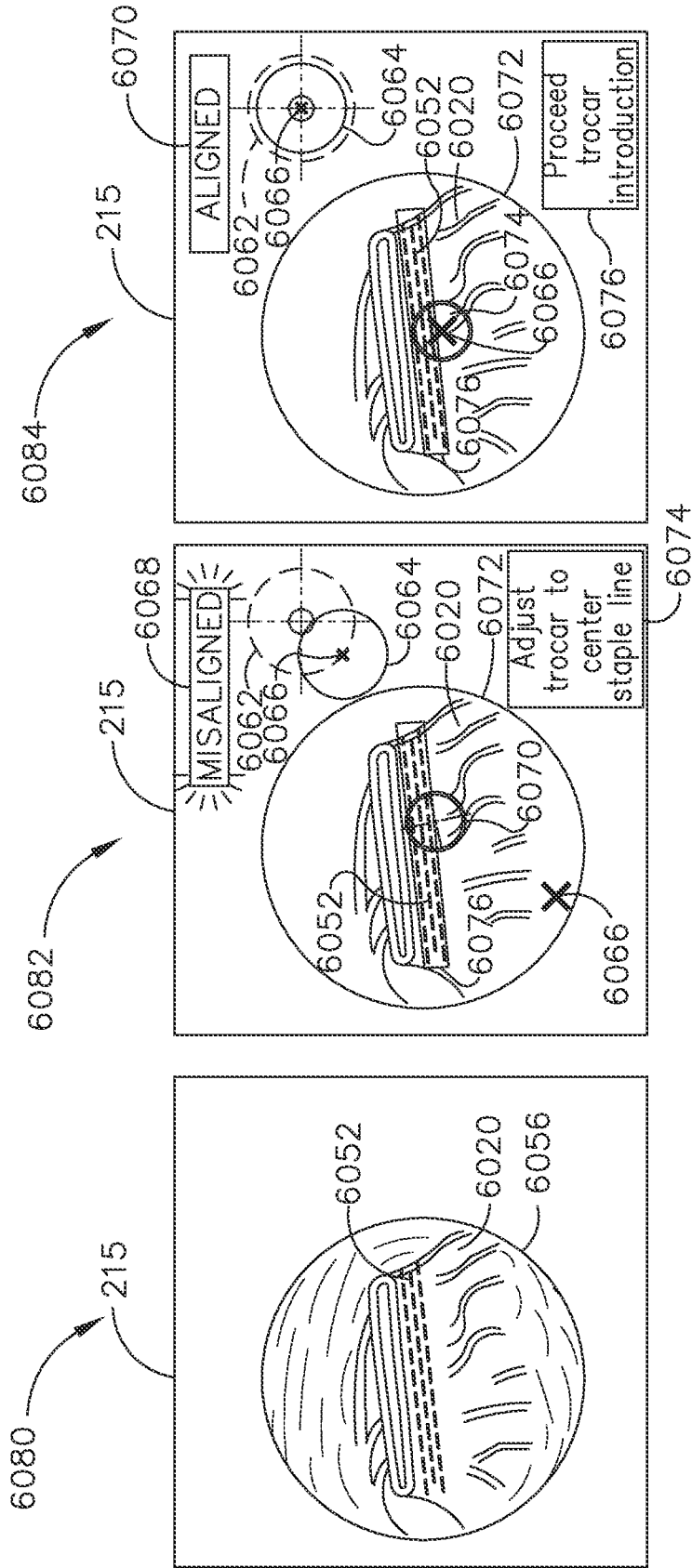


FIG. 117

FIG. 116

FIG. 115

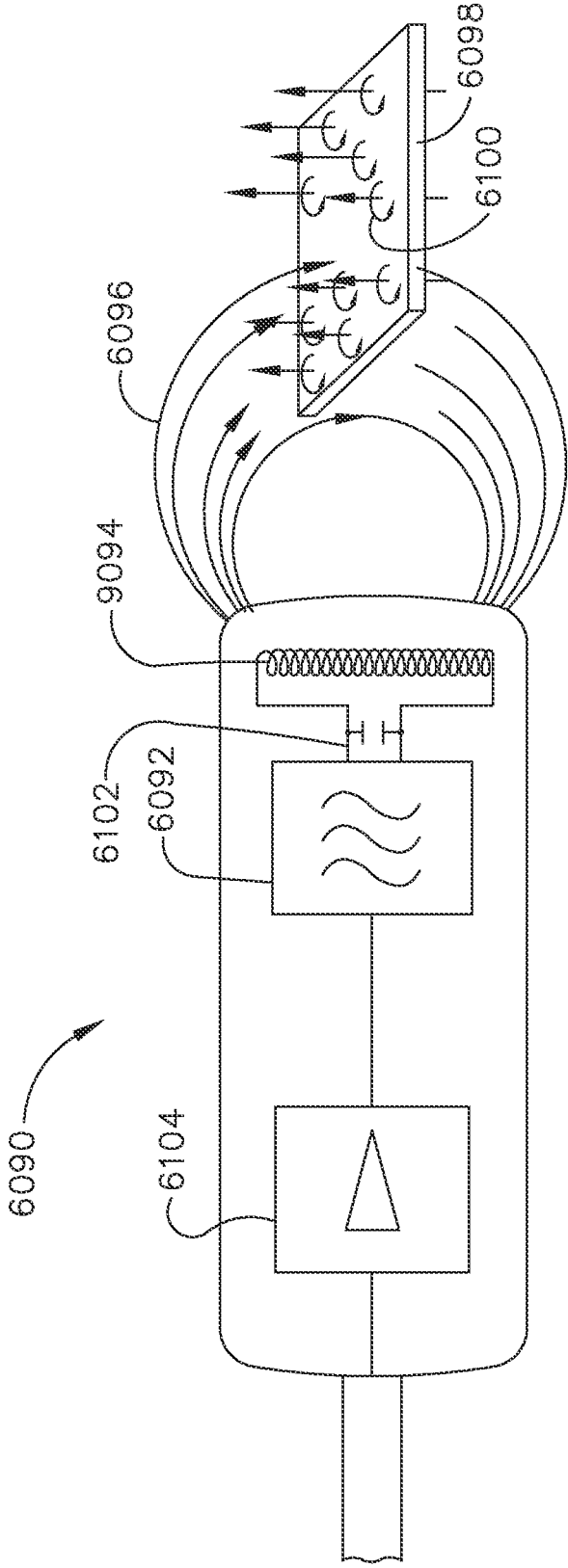


FIG. 118

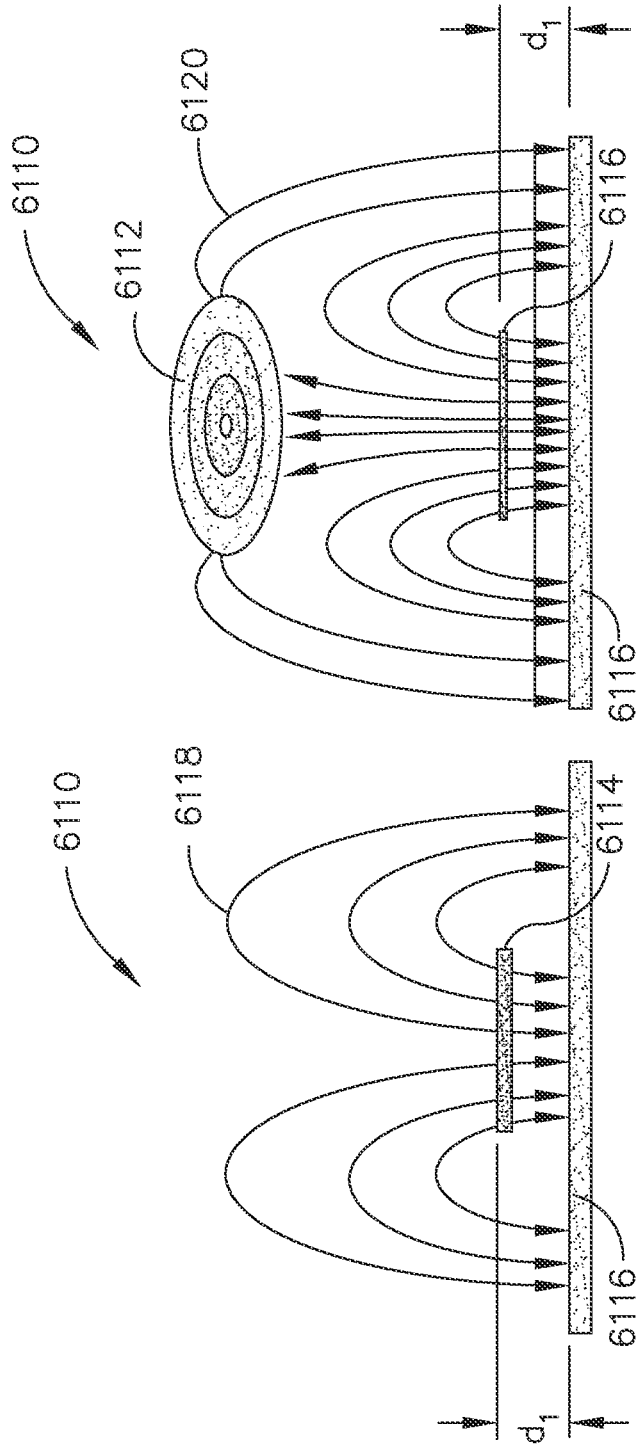


FIG. 119B

FIG. 119A

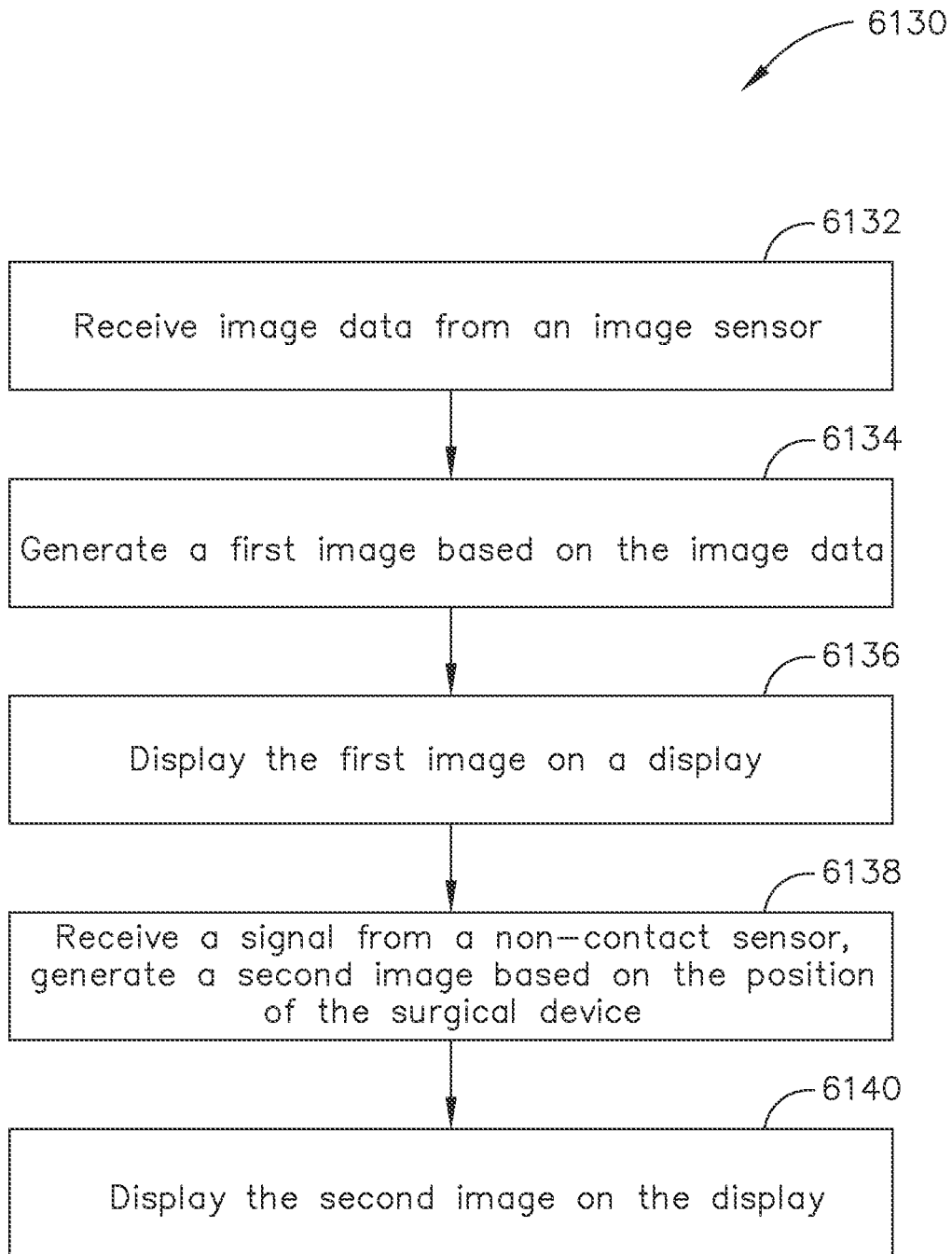


FIG. 120

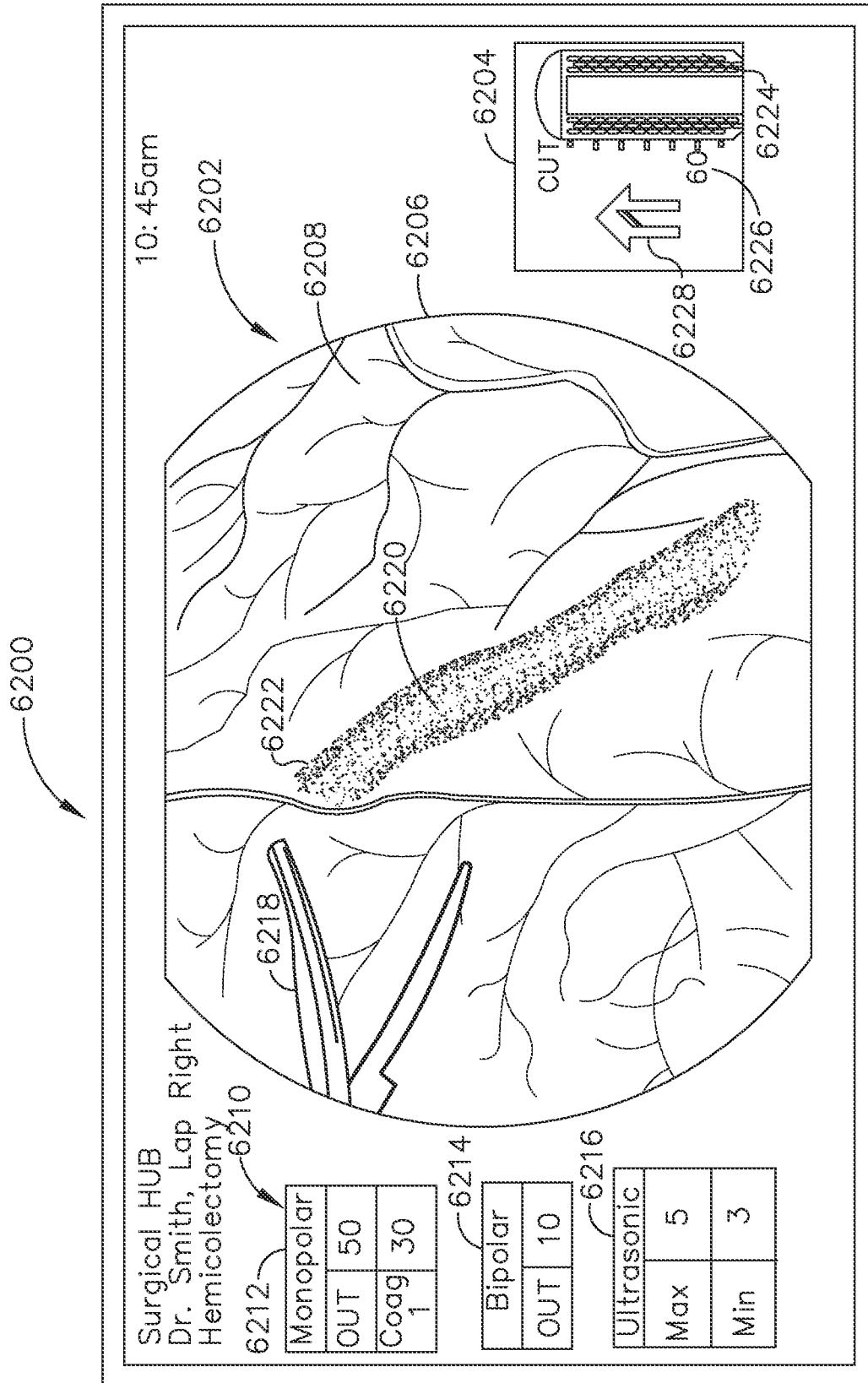


FIG. 121

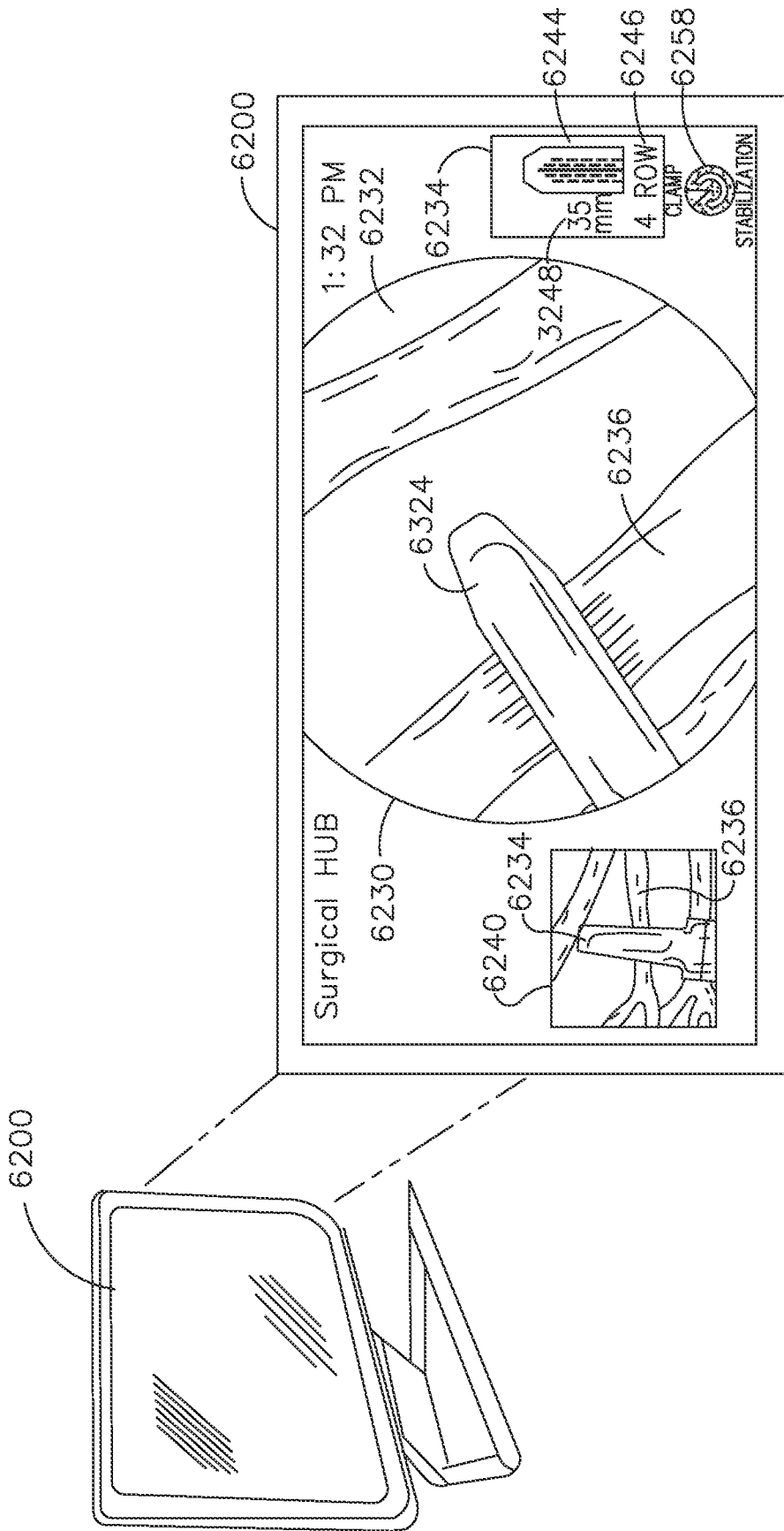


FIG. 122

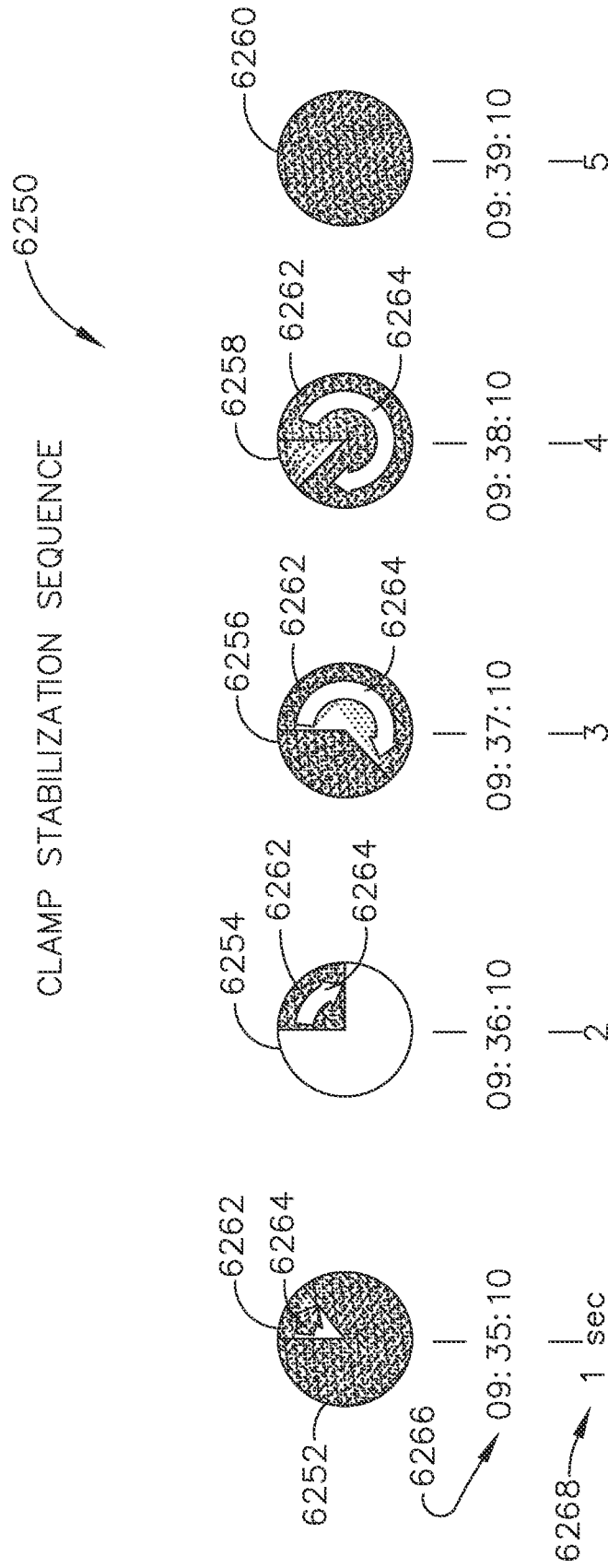


FIG. 123

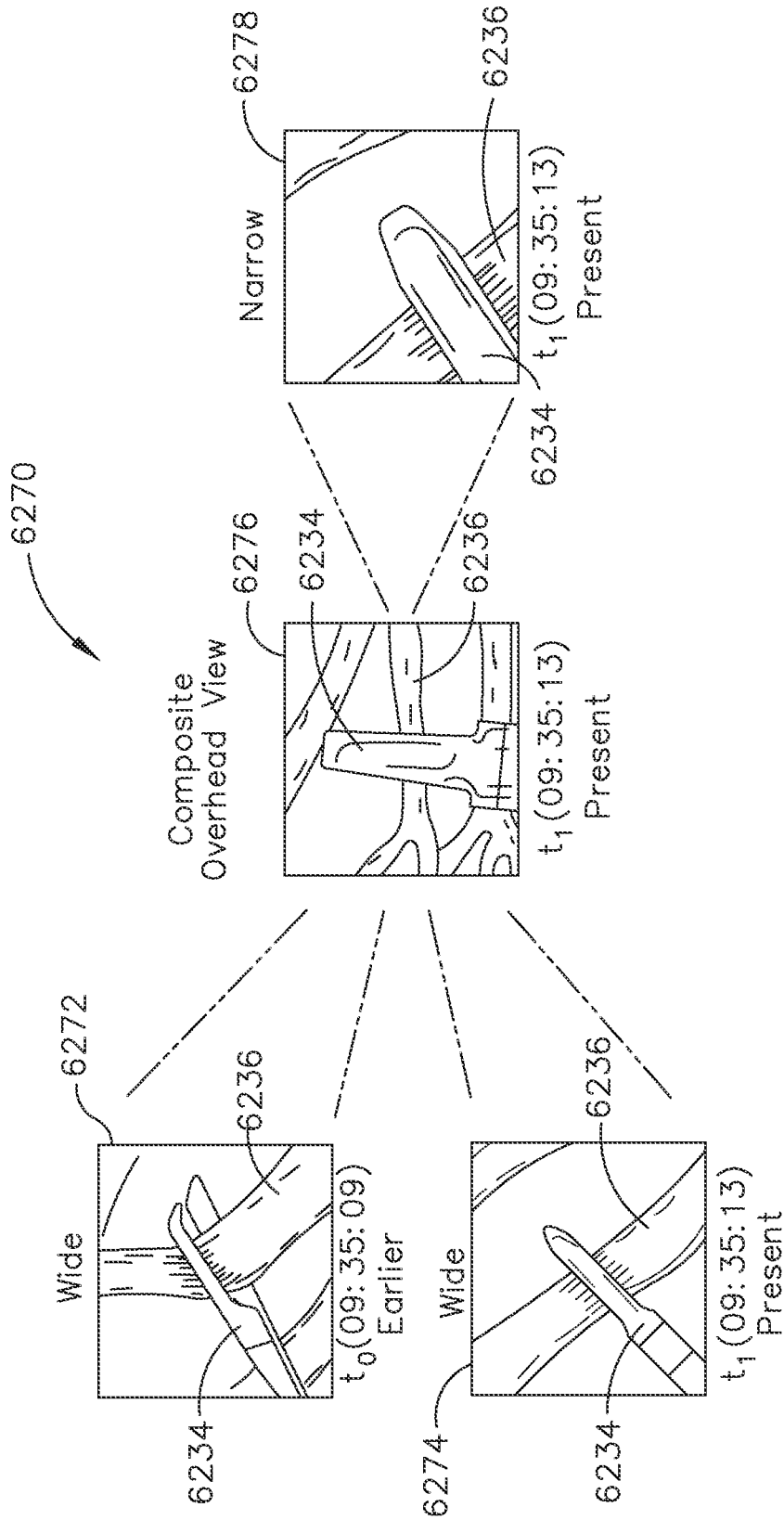


FIG. 124



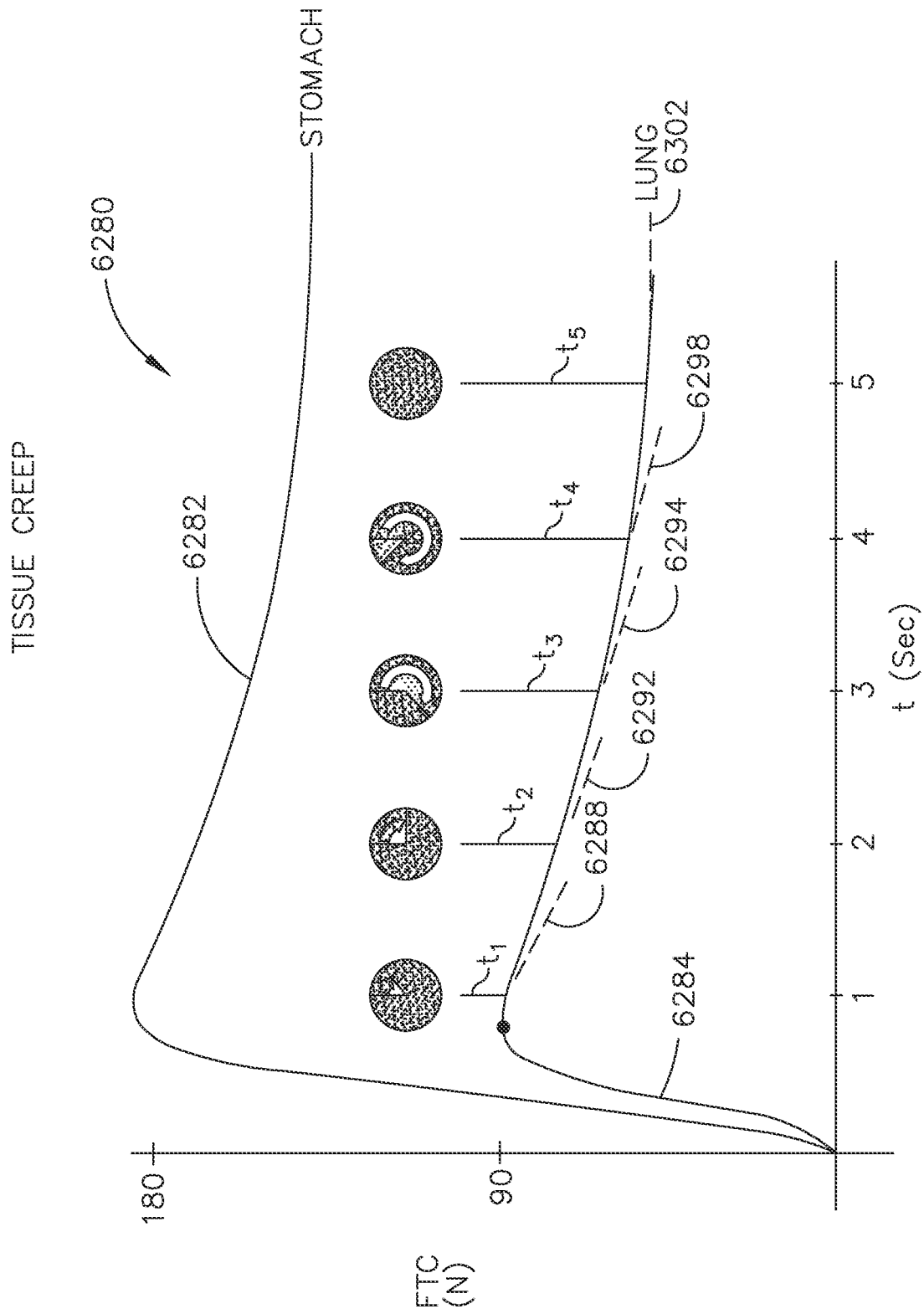


FIG. 125

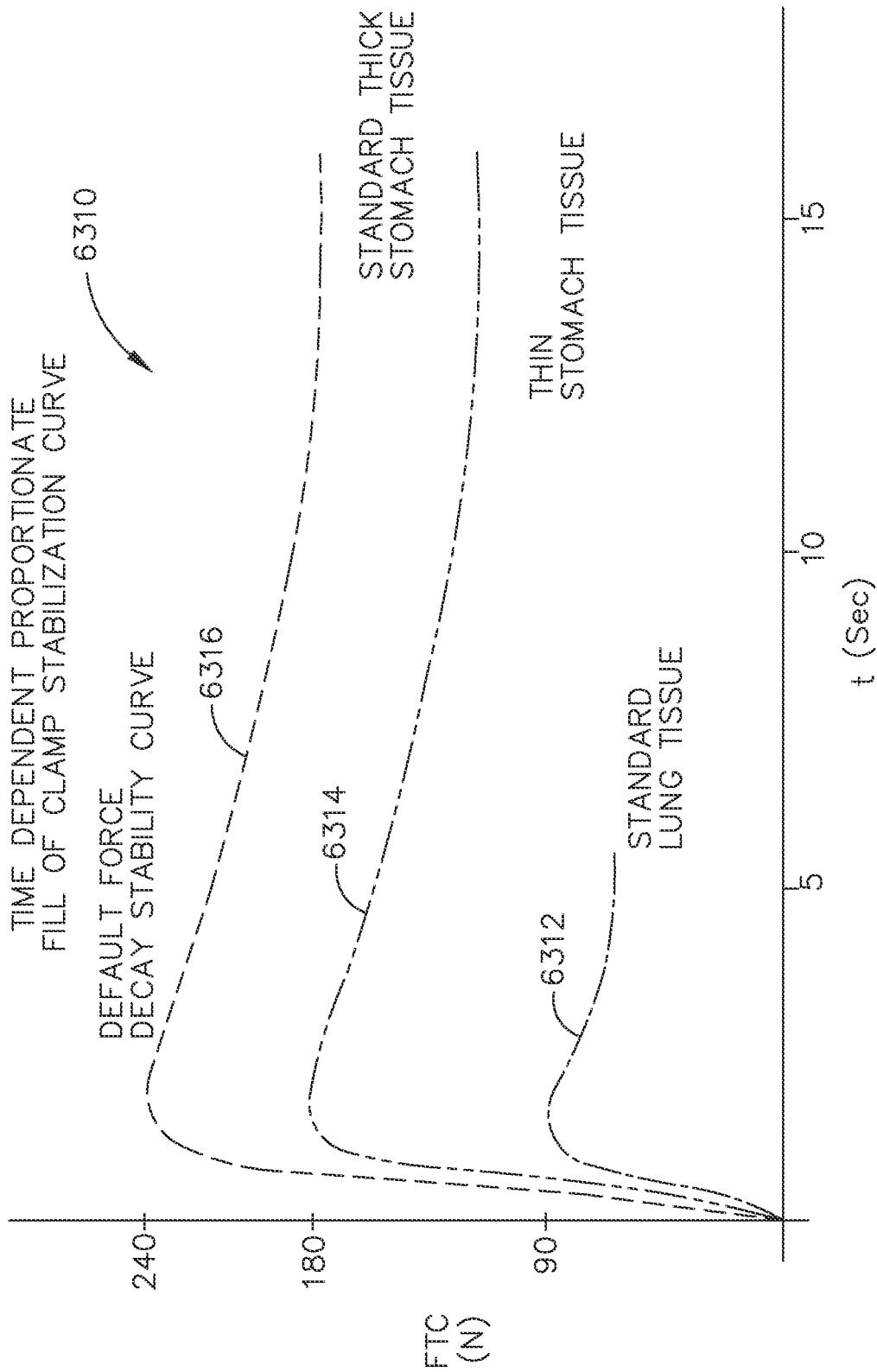


FIG. 126

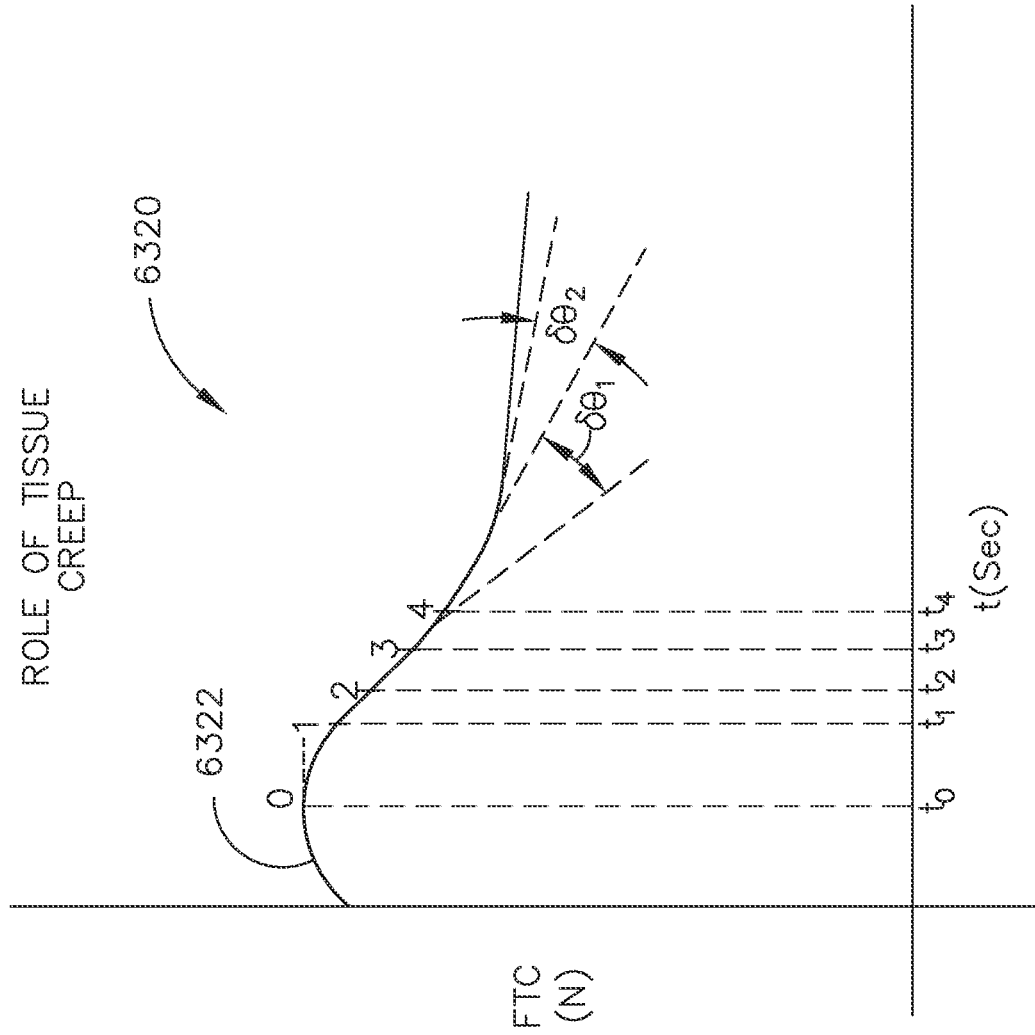
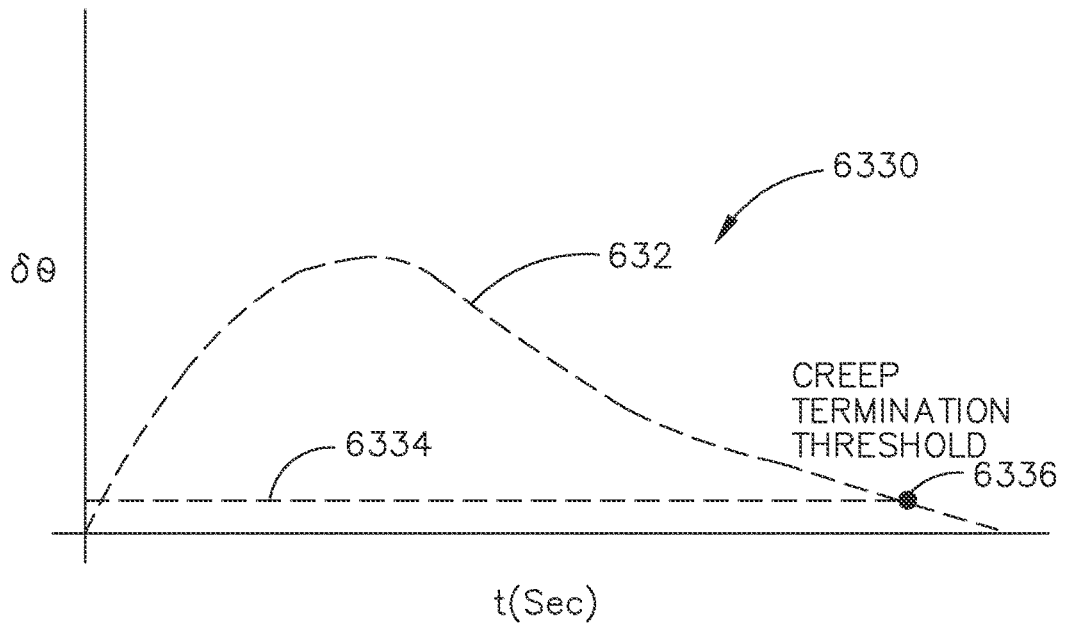


FIG. 127



t(Sec)  
FIG. 128A

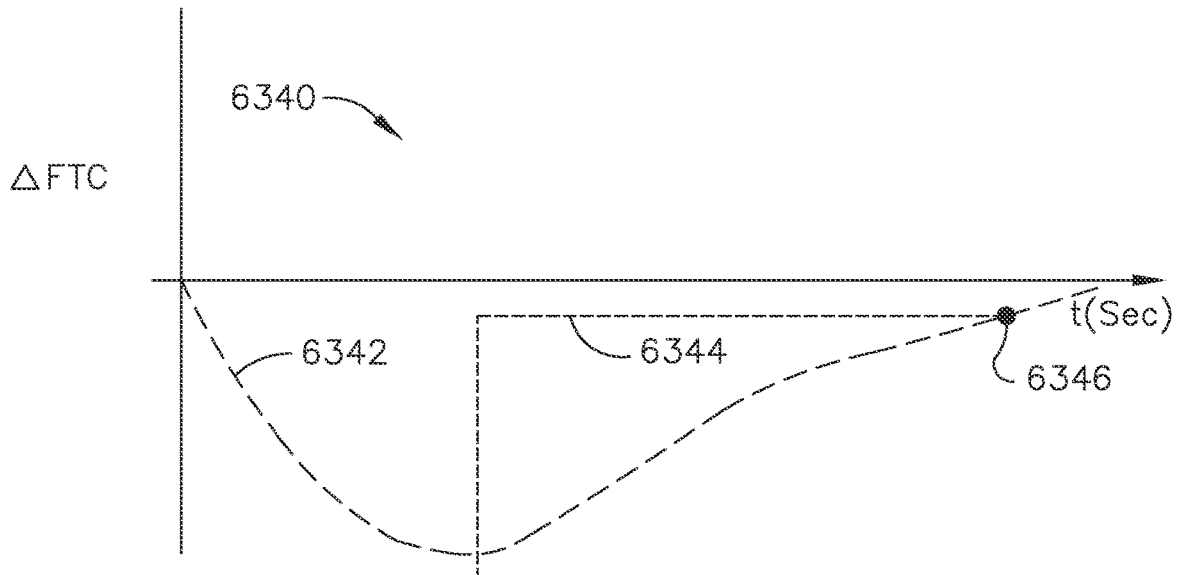


FIG. 128B

THRESHOLD TO 100%  
COMPLETE CREEP  
STABILITY METER

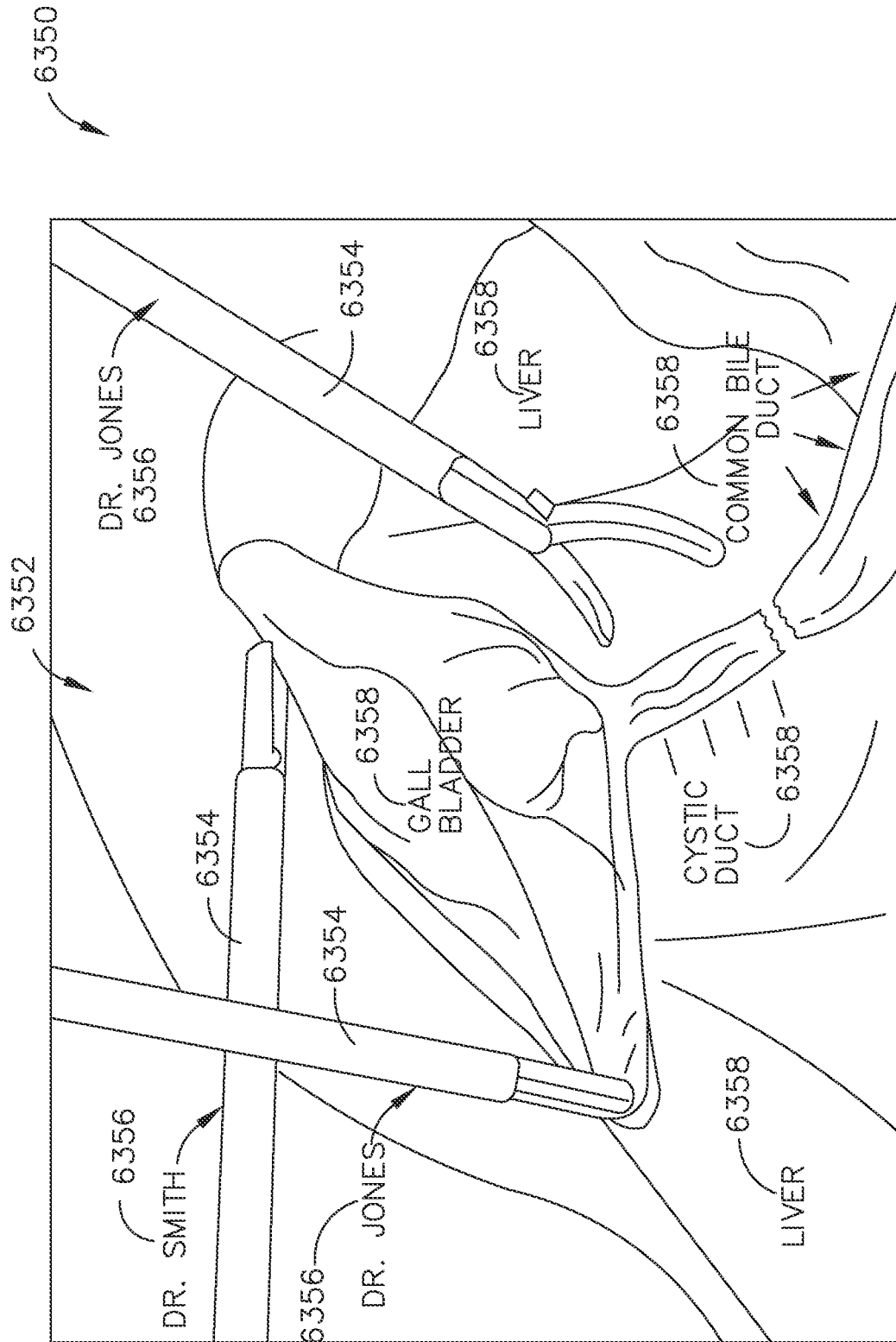


FIG. 129

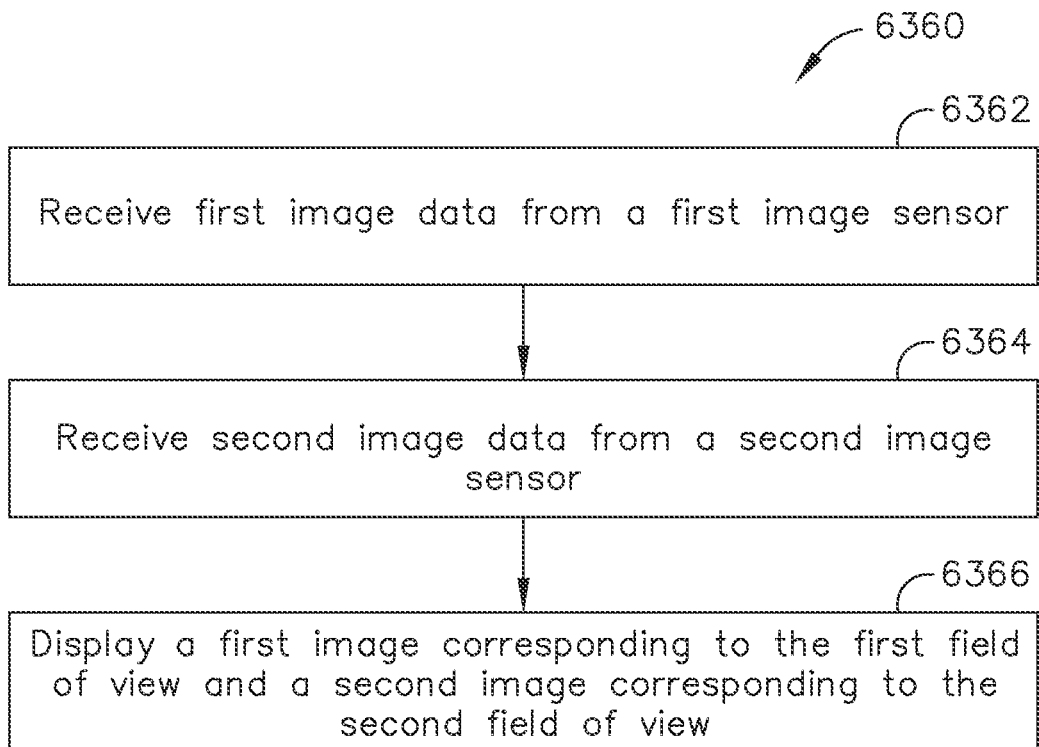


FIG. 130

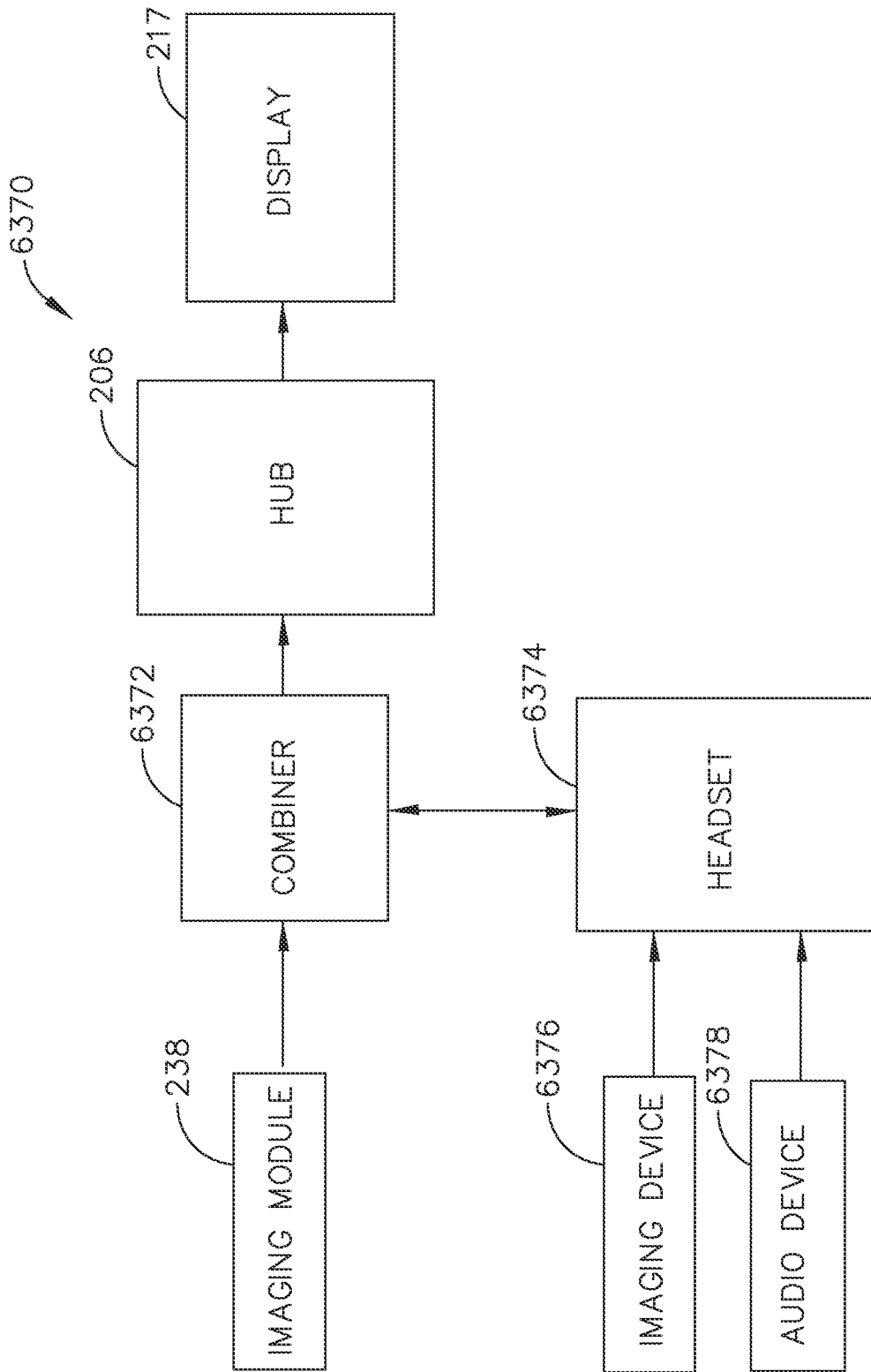
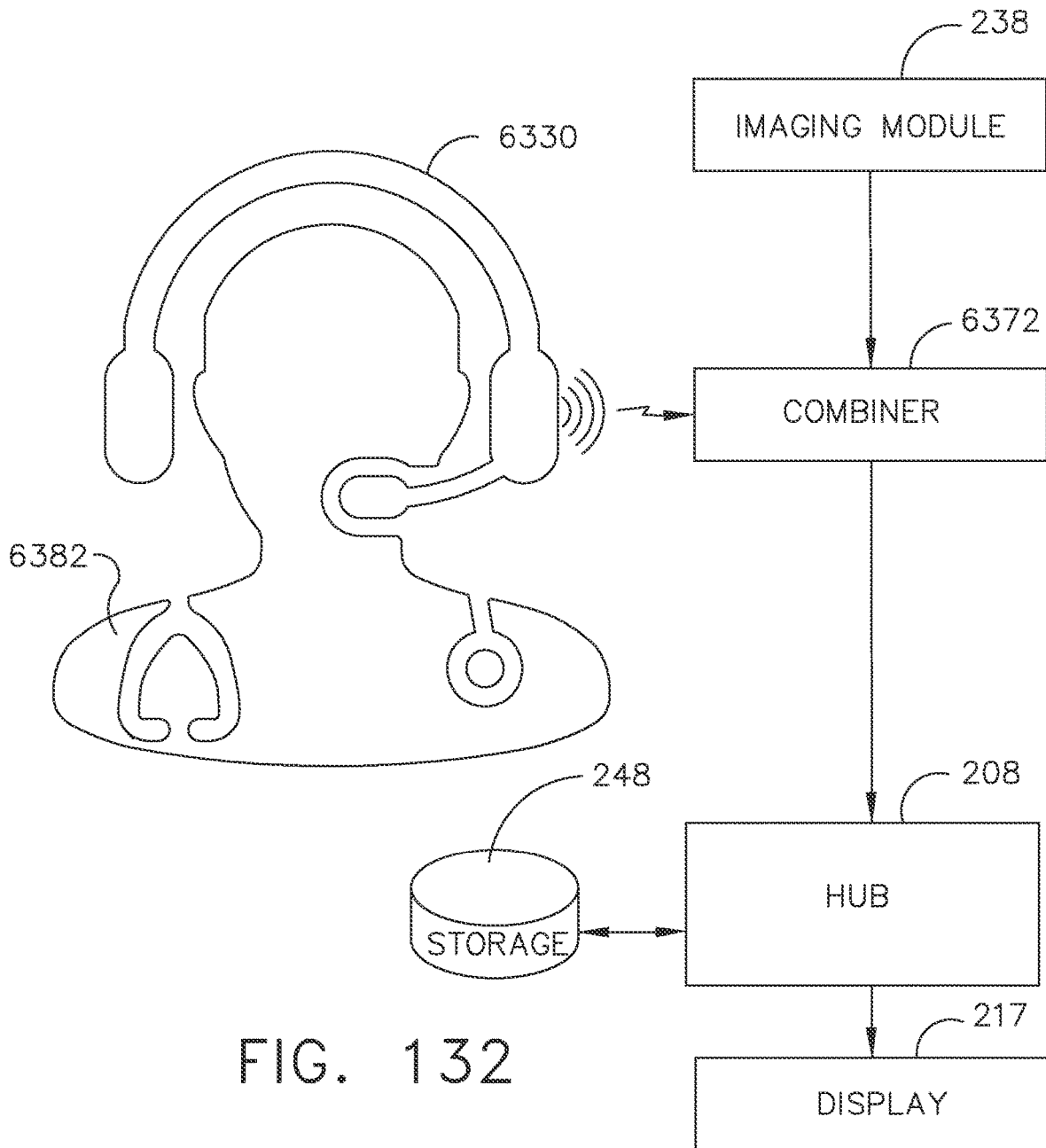


FIG. 131





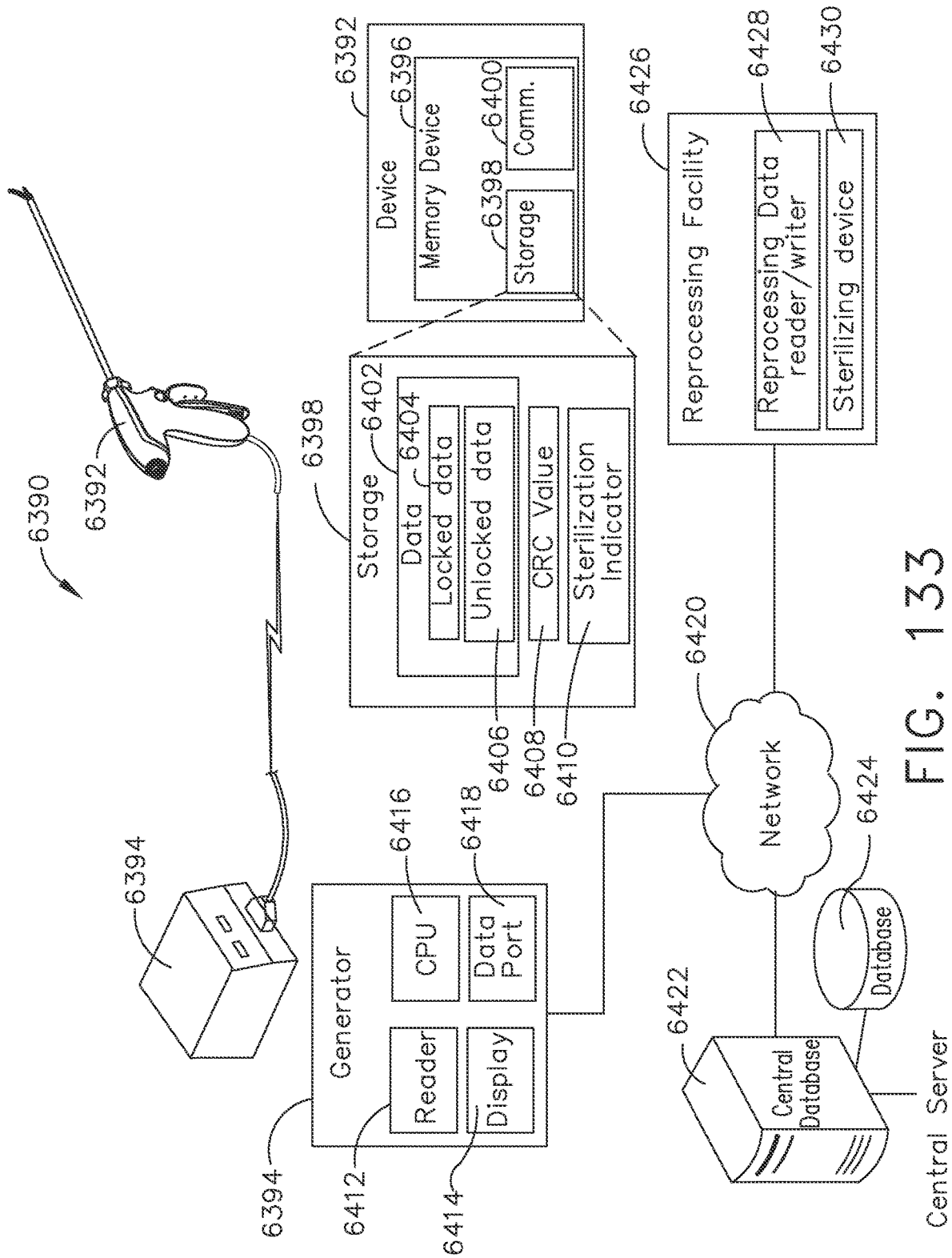


FIG. 133

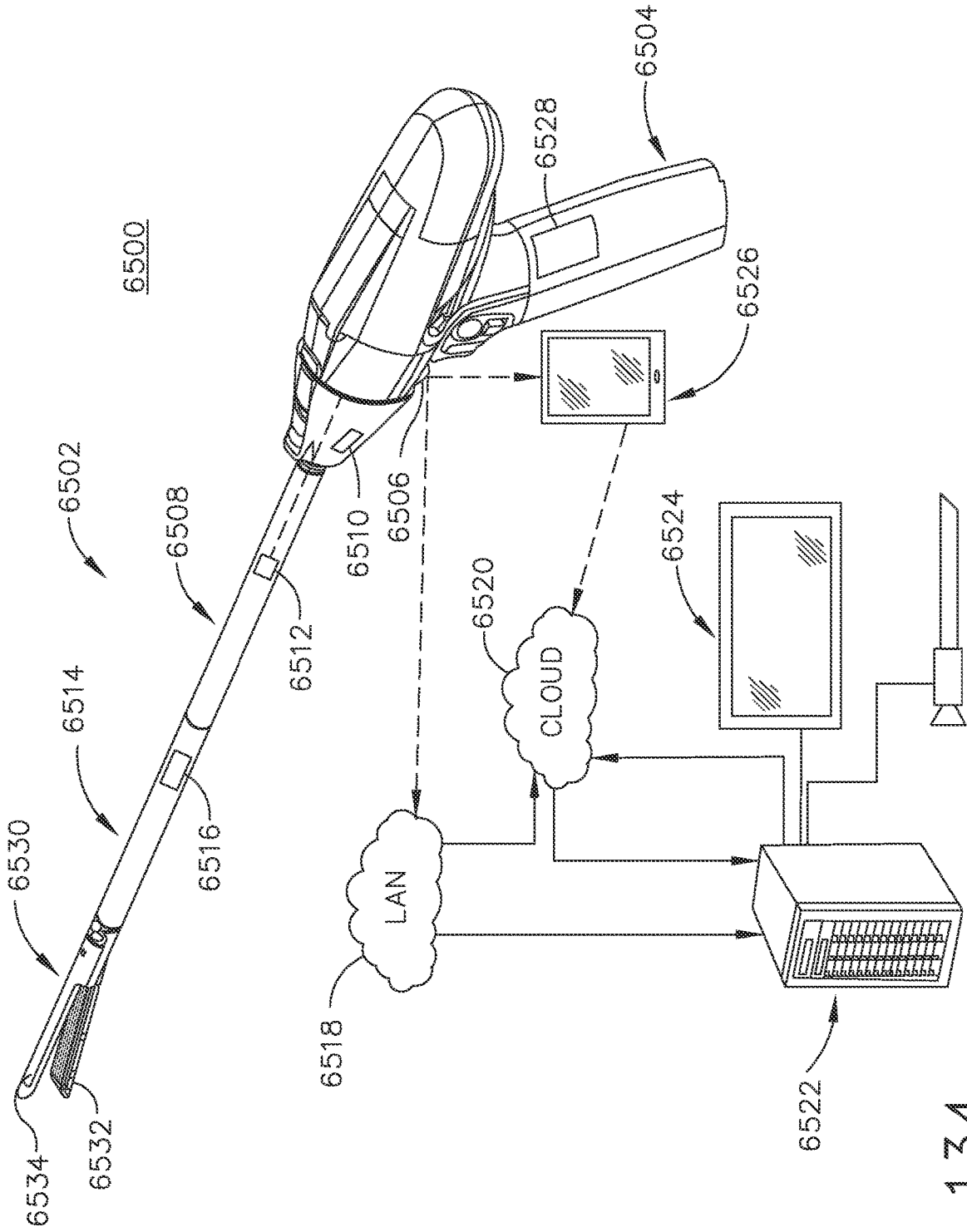


FIG. 134

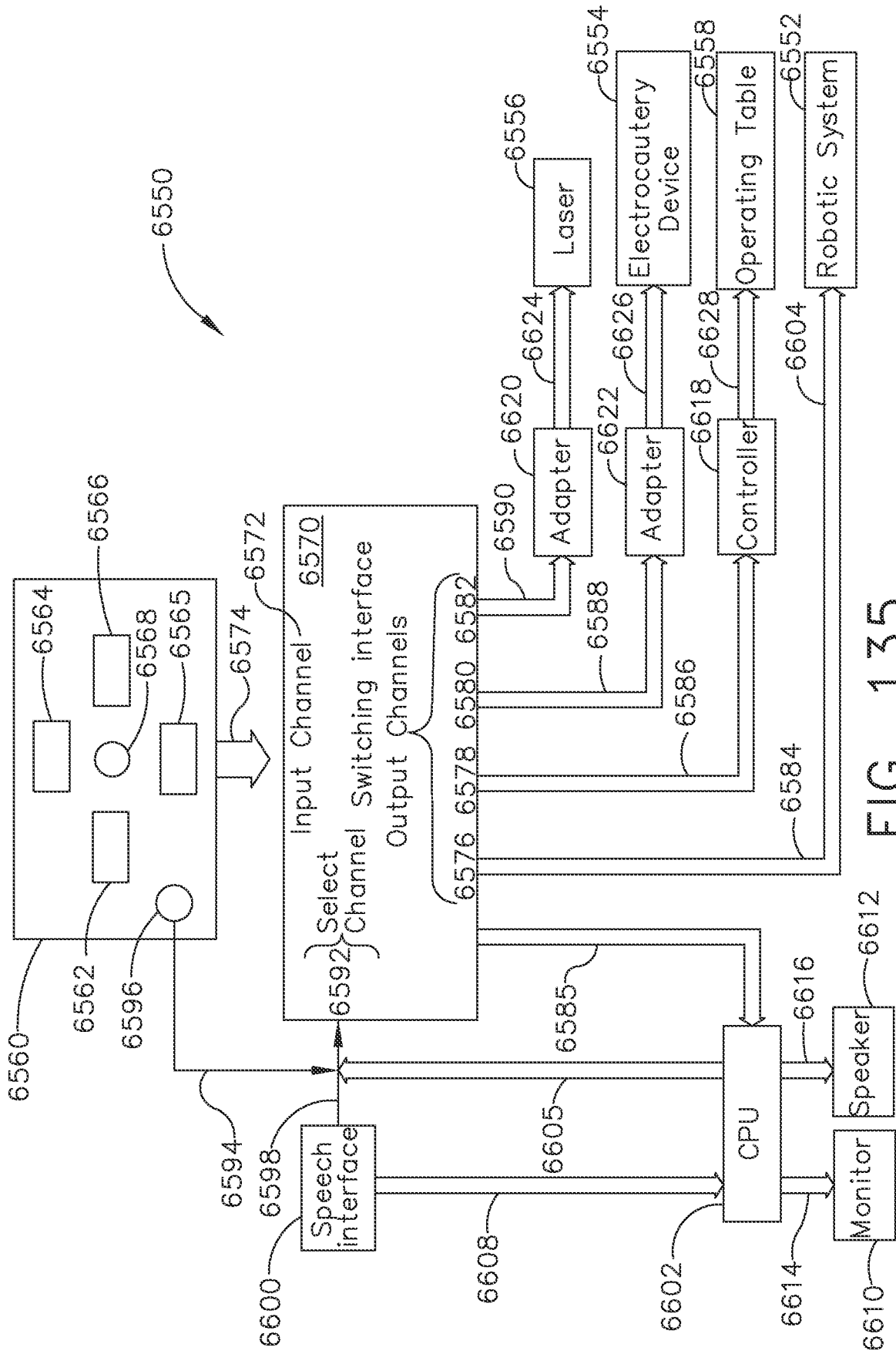
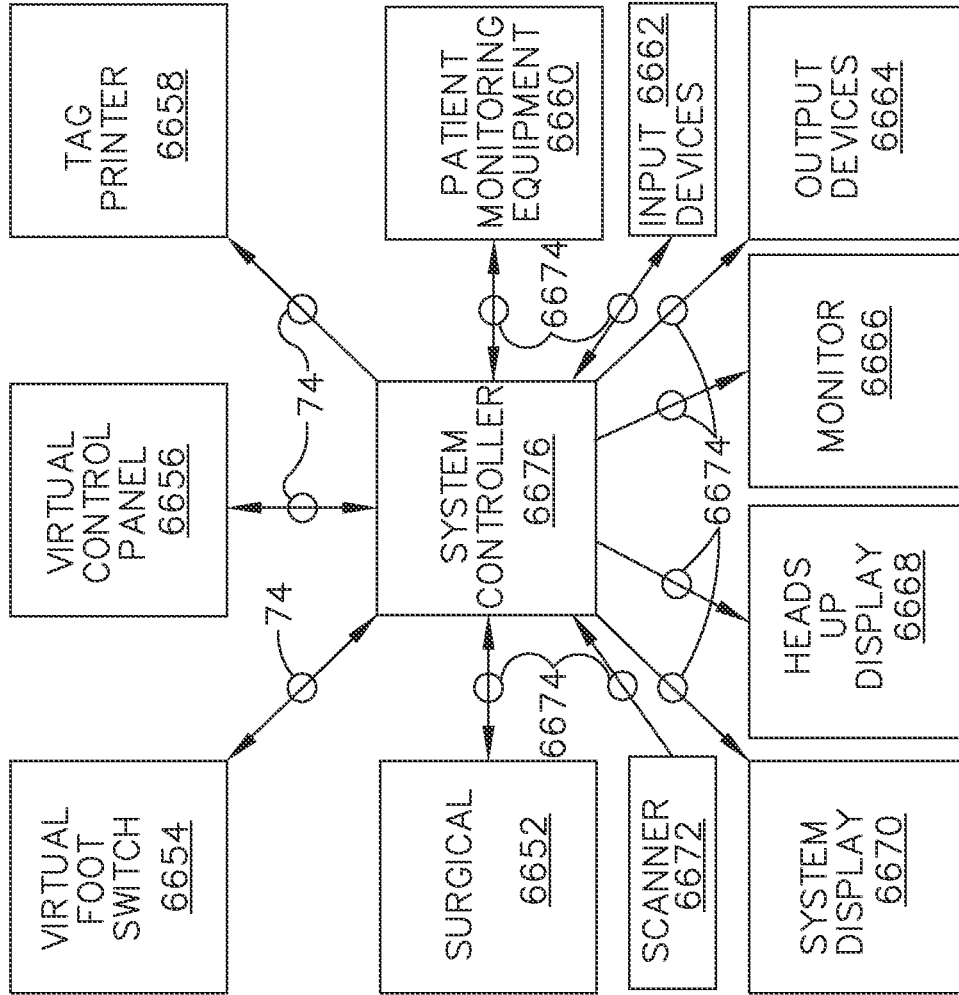


FIG. 135



6650 → FIG. 136

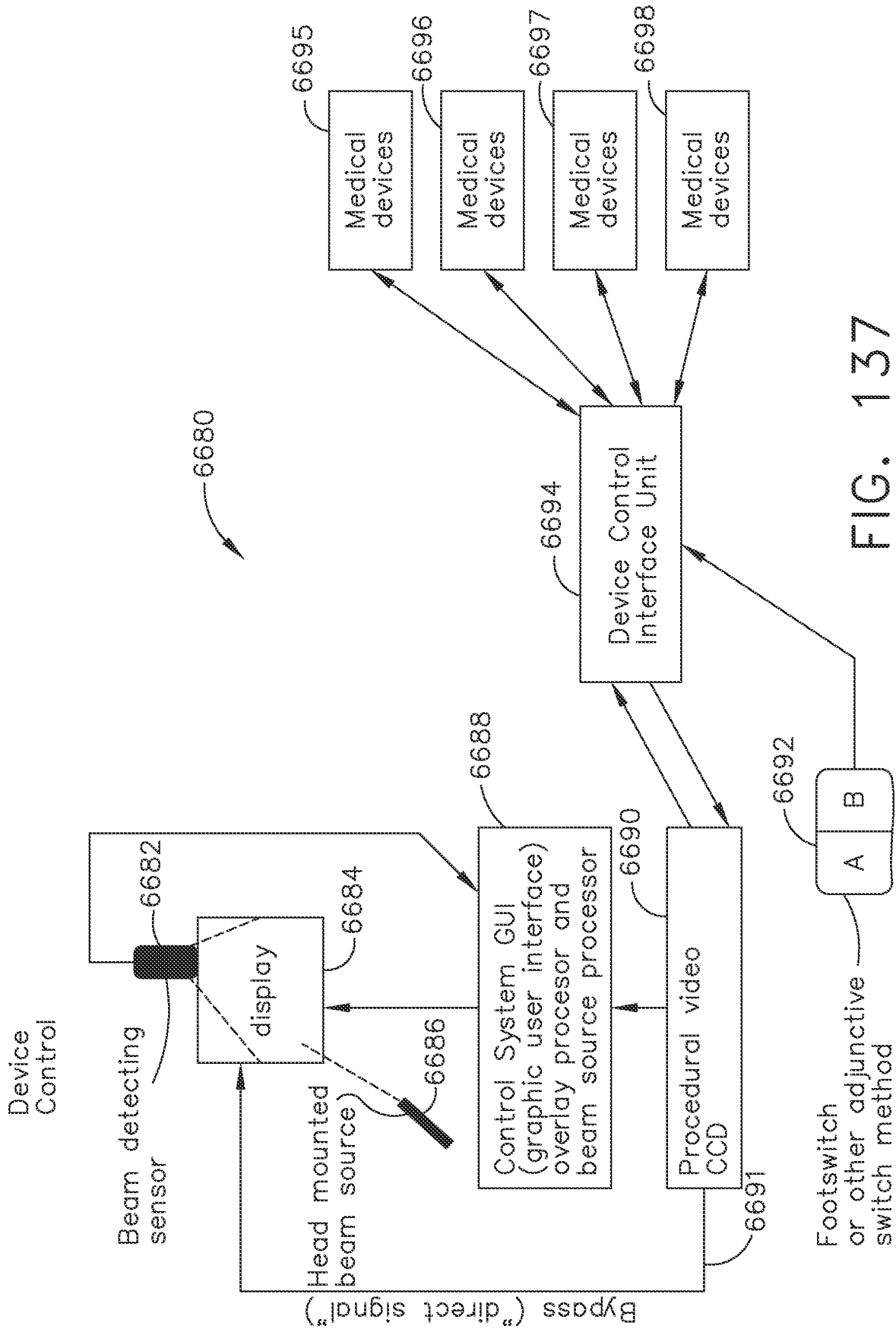


FIG. 137

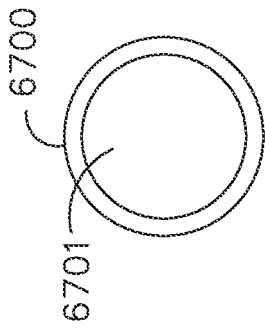


FIG. 138A

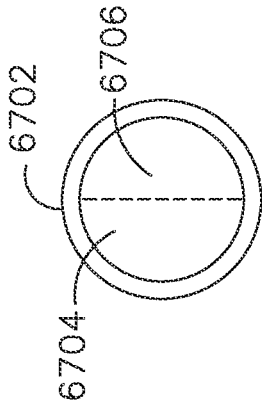


FIG. 138B

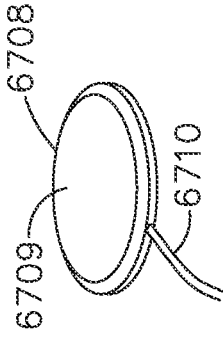


FIG. 138C

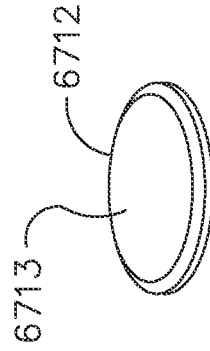


FIG. 138D

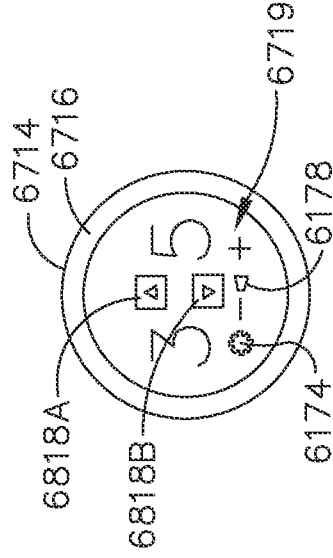


FIG. 138E

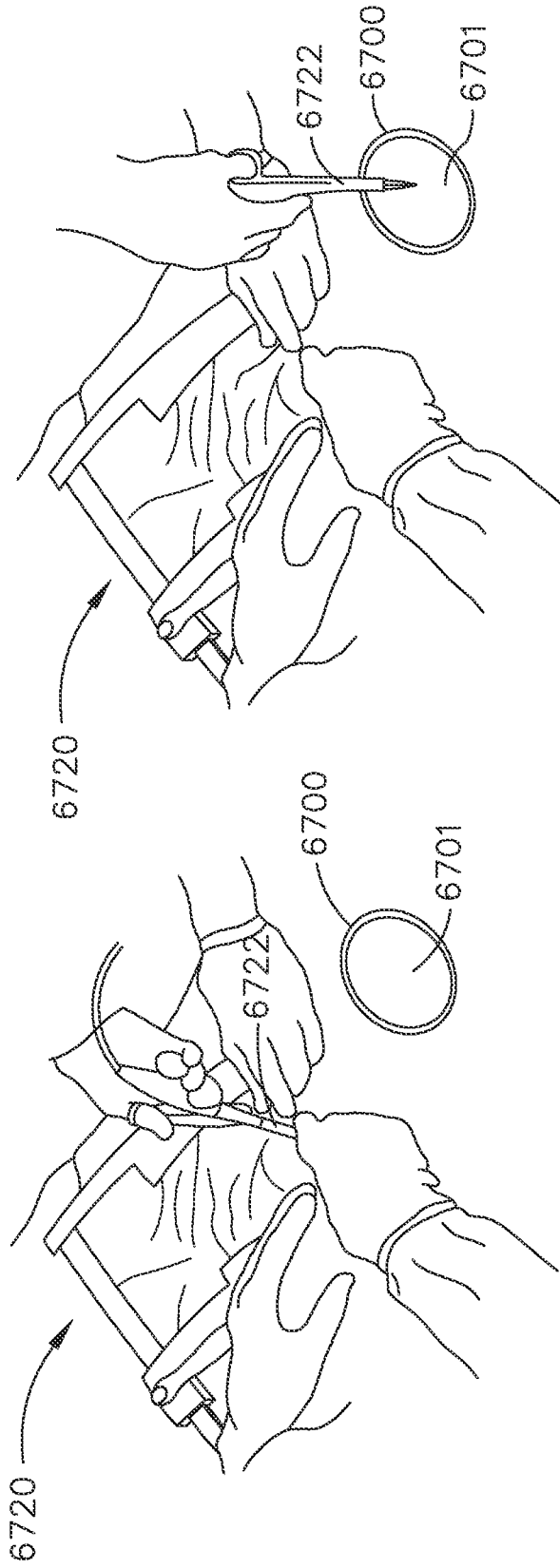


FIG. 139B

FIG. 139A

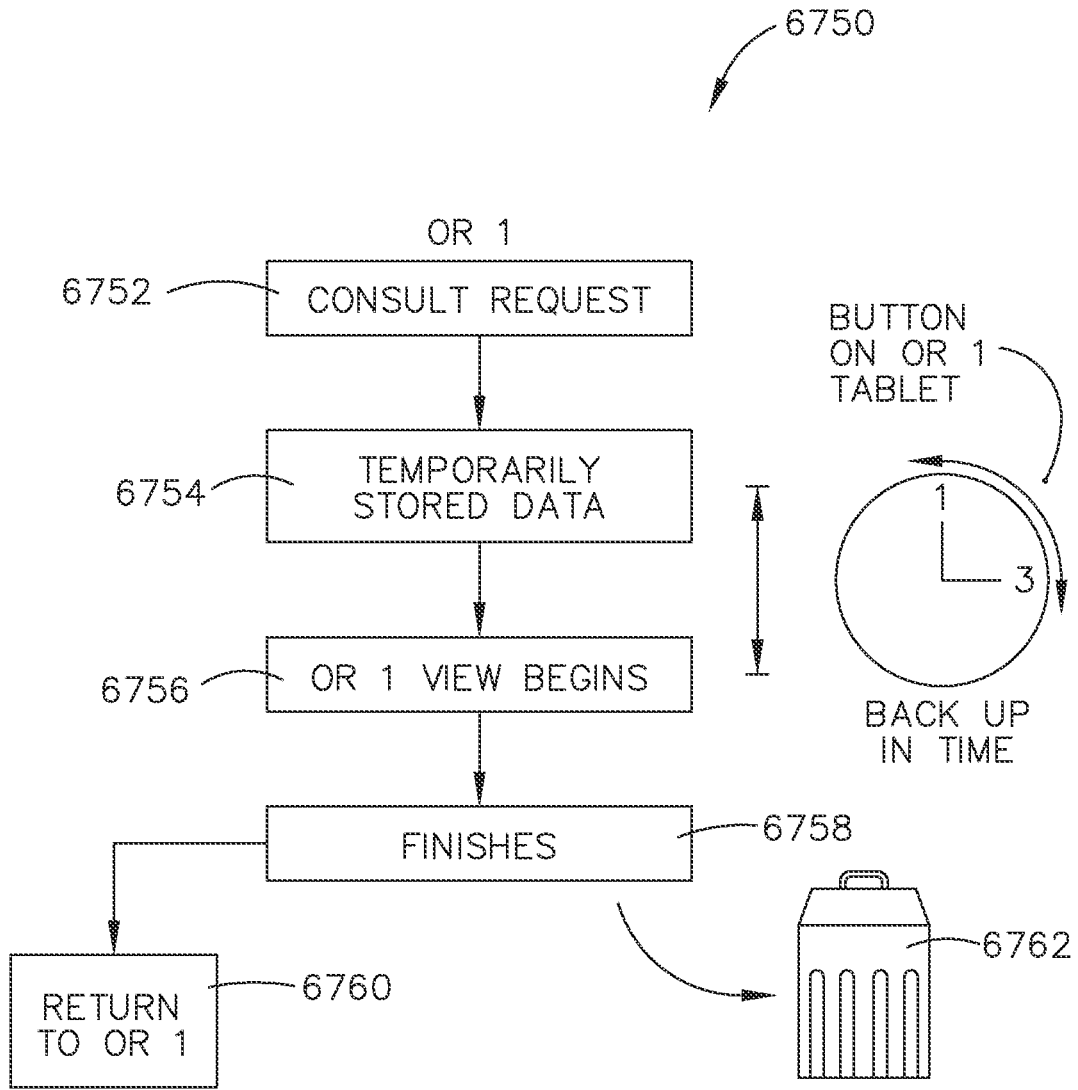


FIG. 140



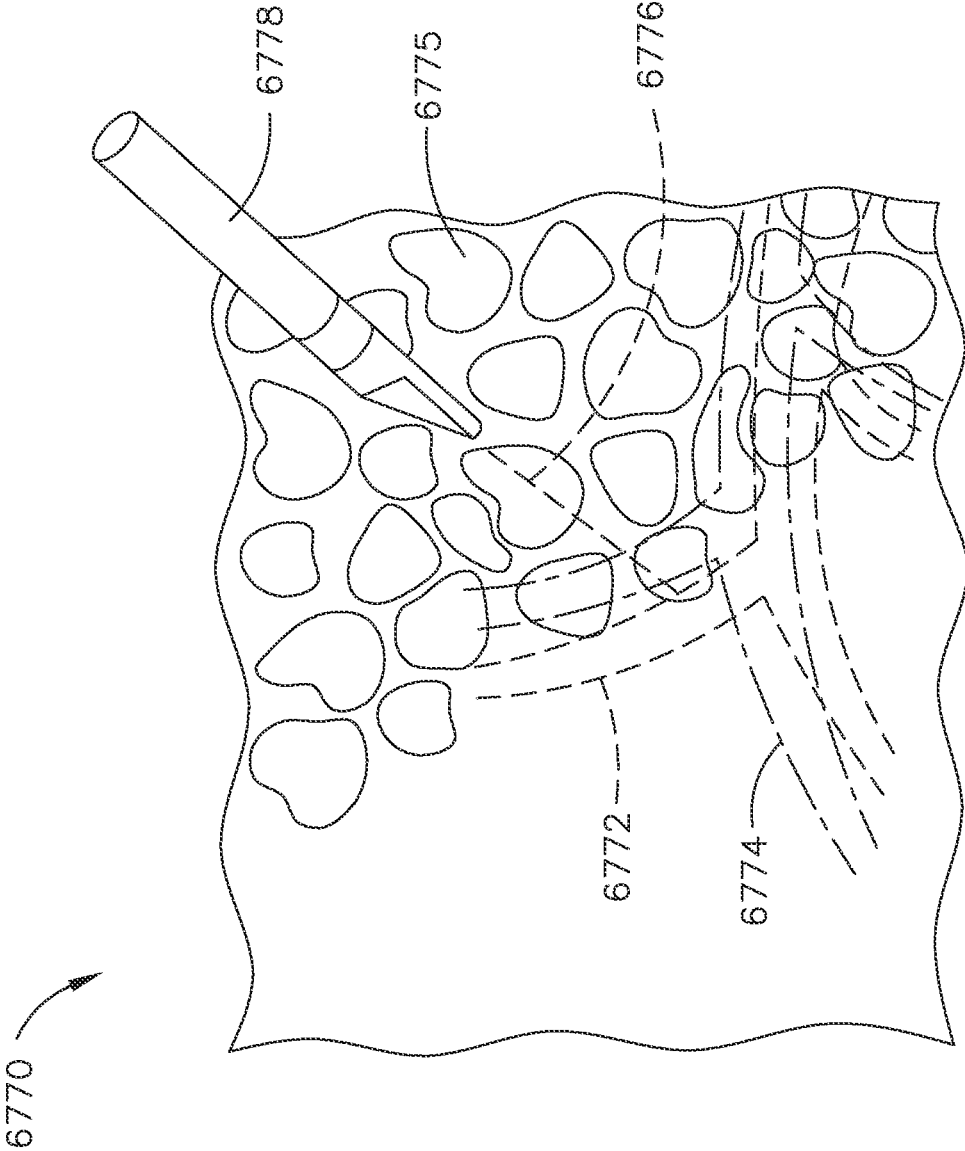


FIG. 141

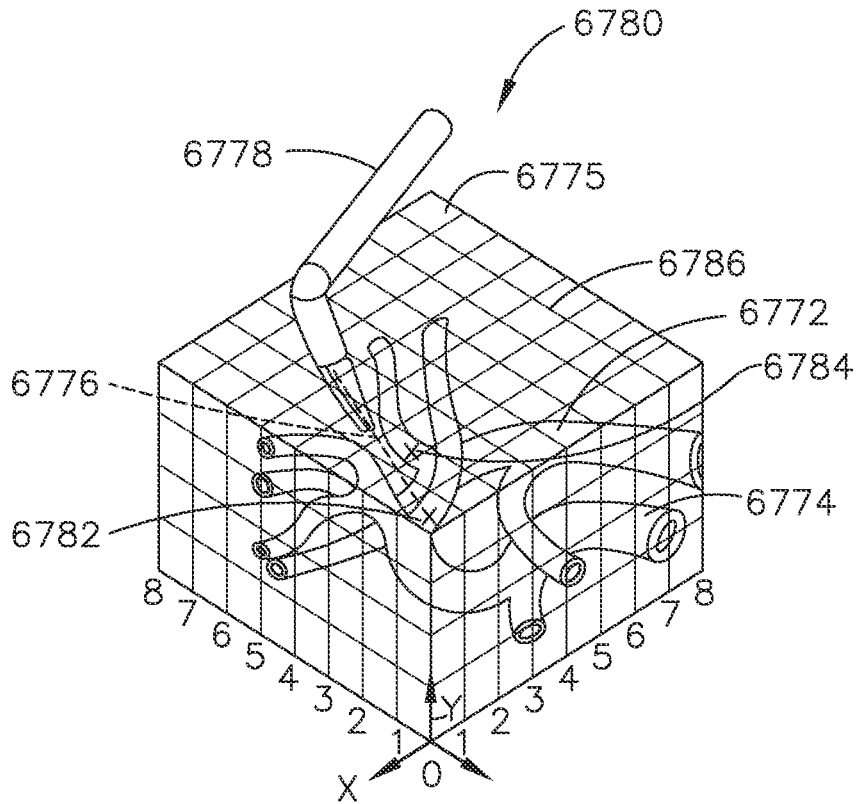


FIG. 142A

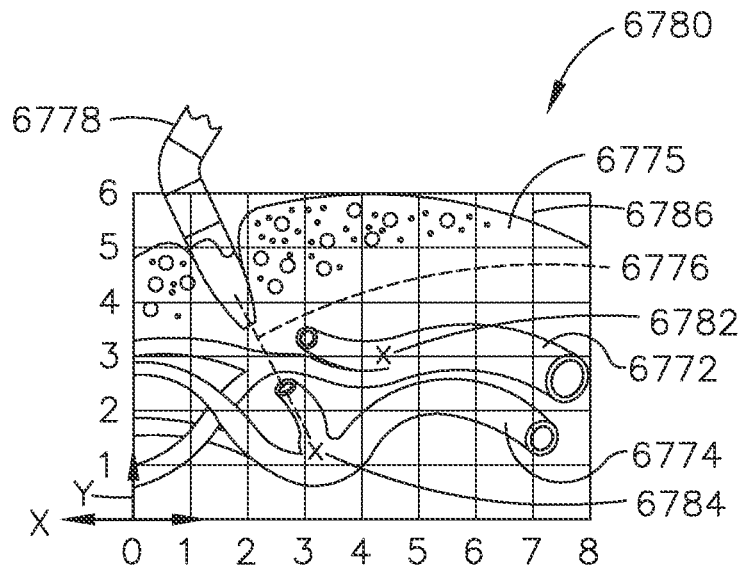


FIG. 142B

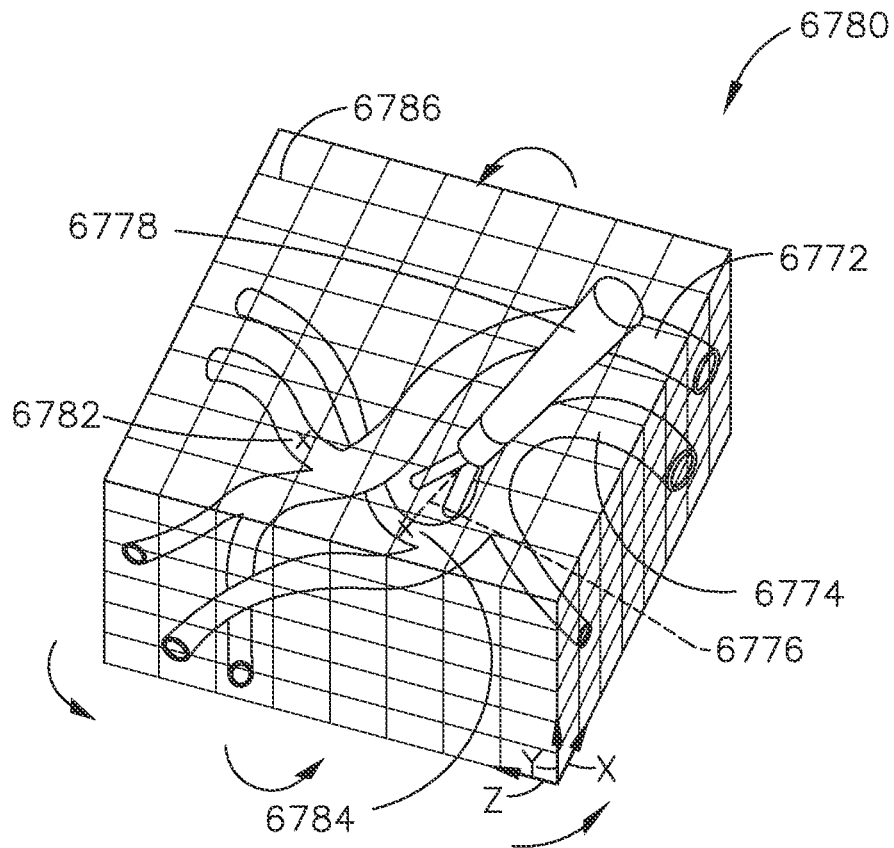


FIG. 142C

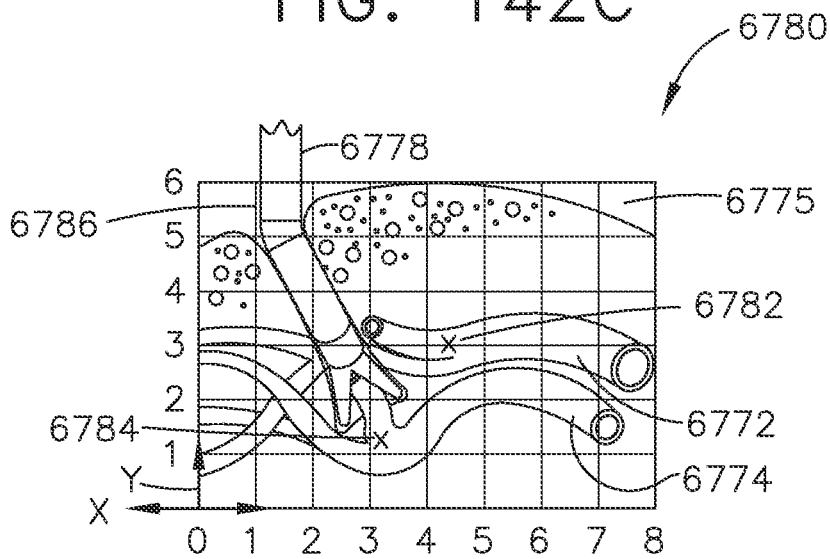


FIG. 142D

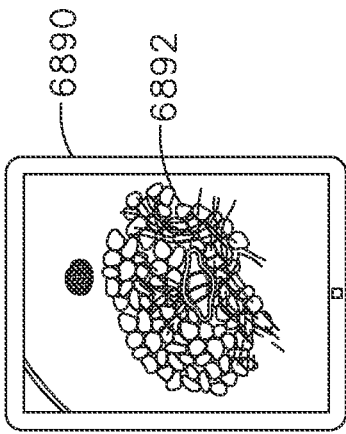


FIG. 143A

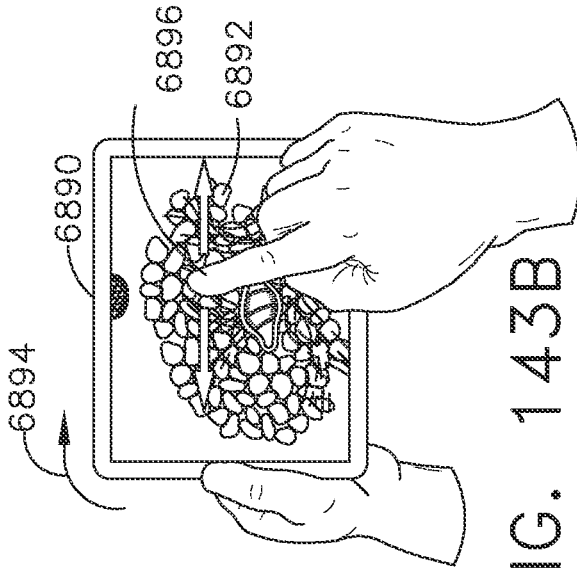


FIG. 143B

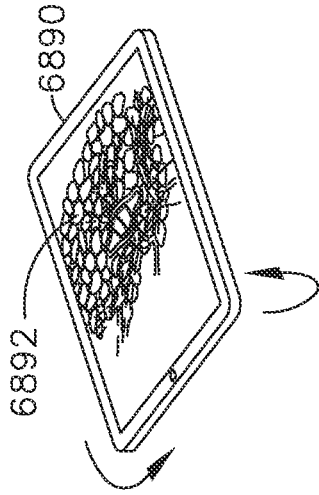


FIG. 143E

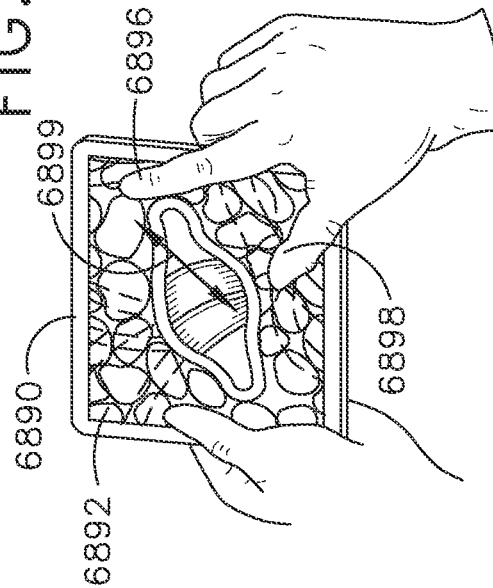


FIG. 143C

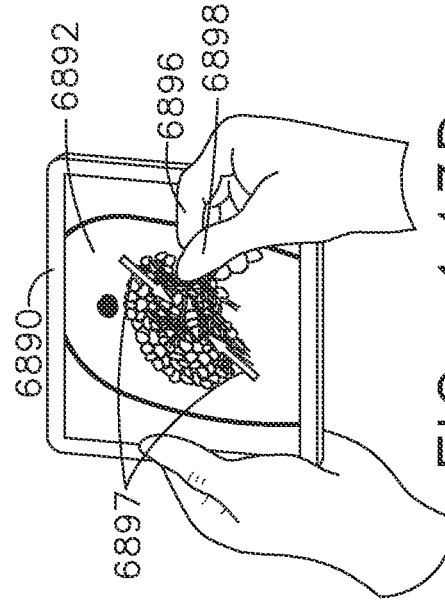


FIG. 143D

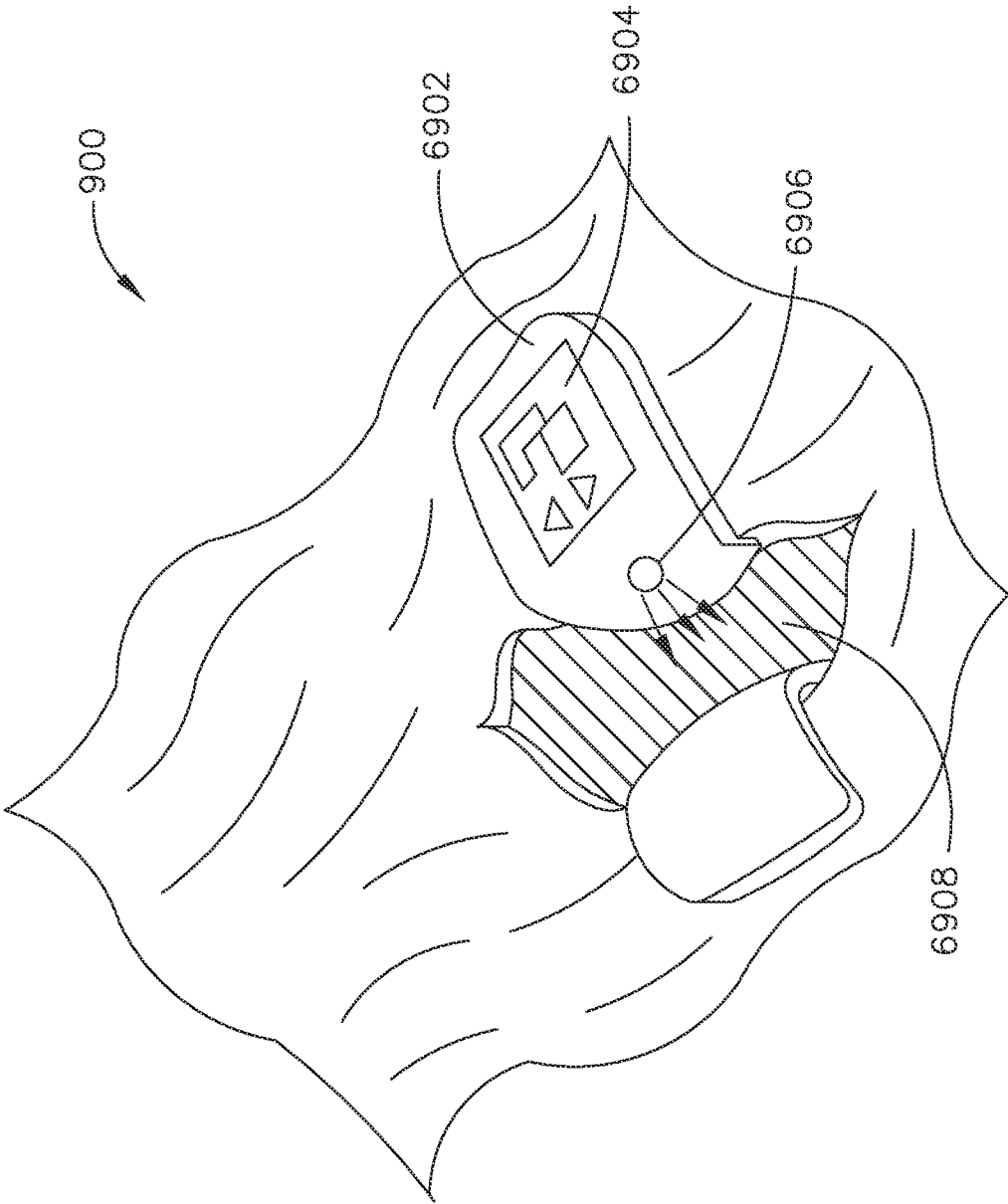


FIG. 144

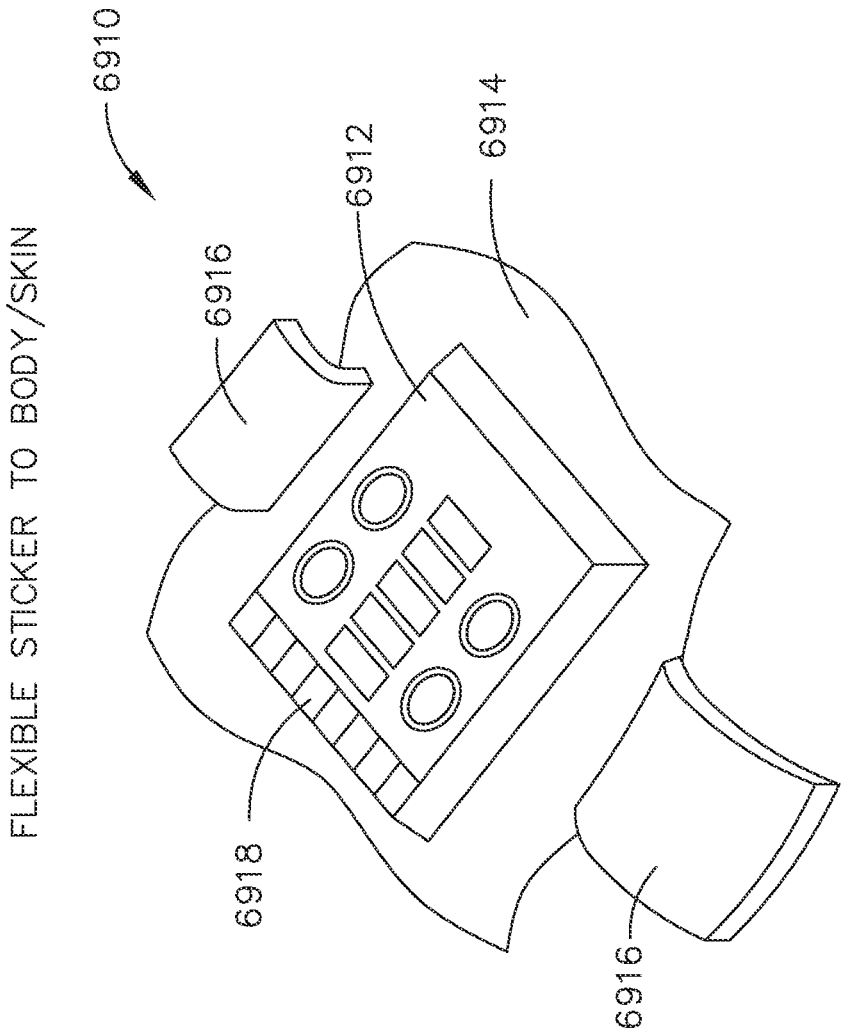


FIG. 145

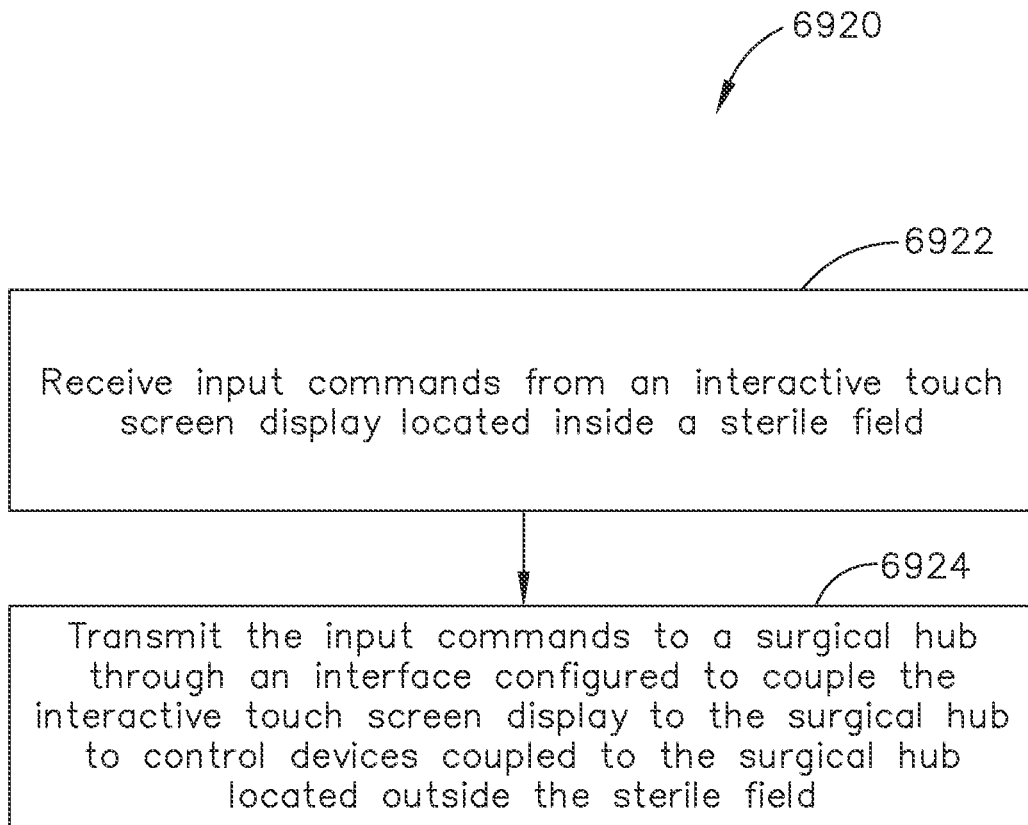


FIG. 146

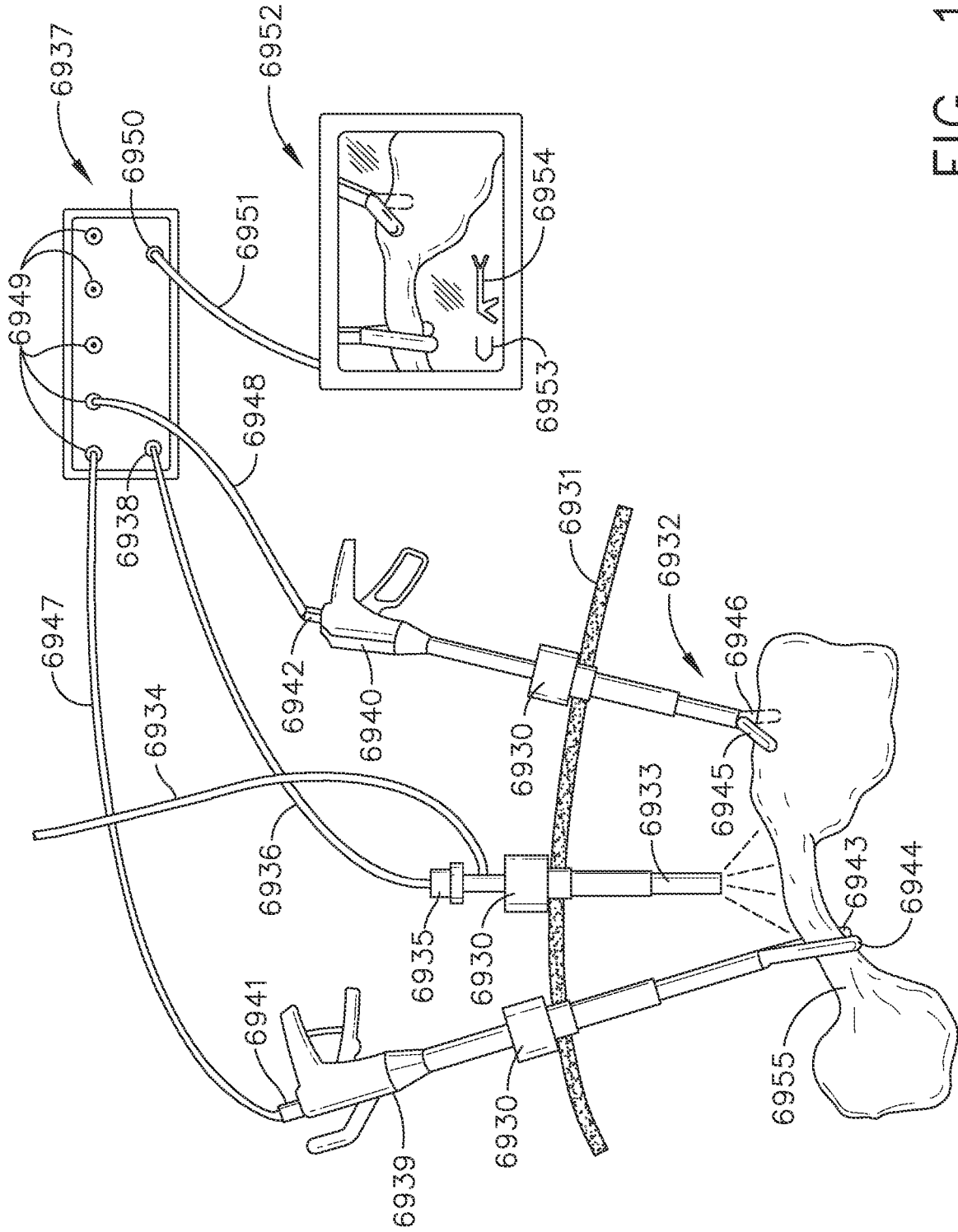


FIG. 147



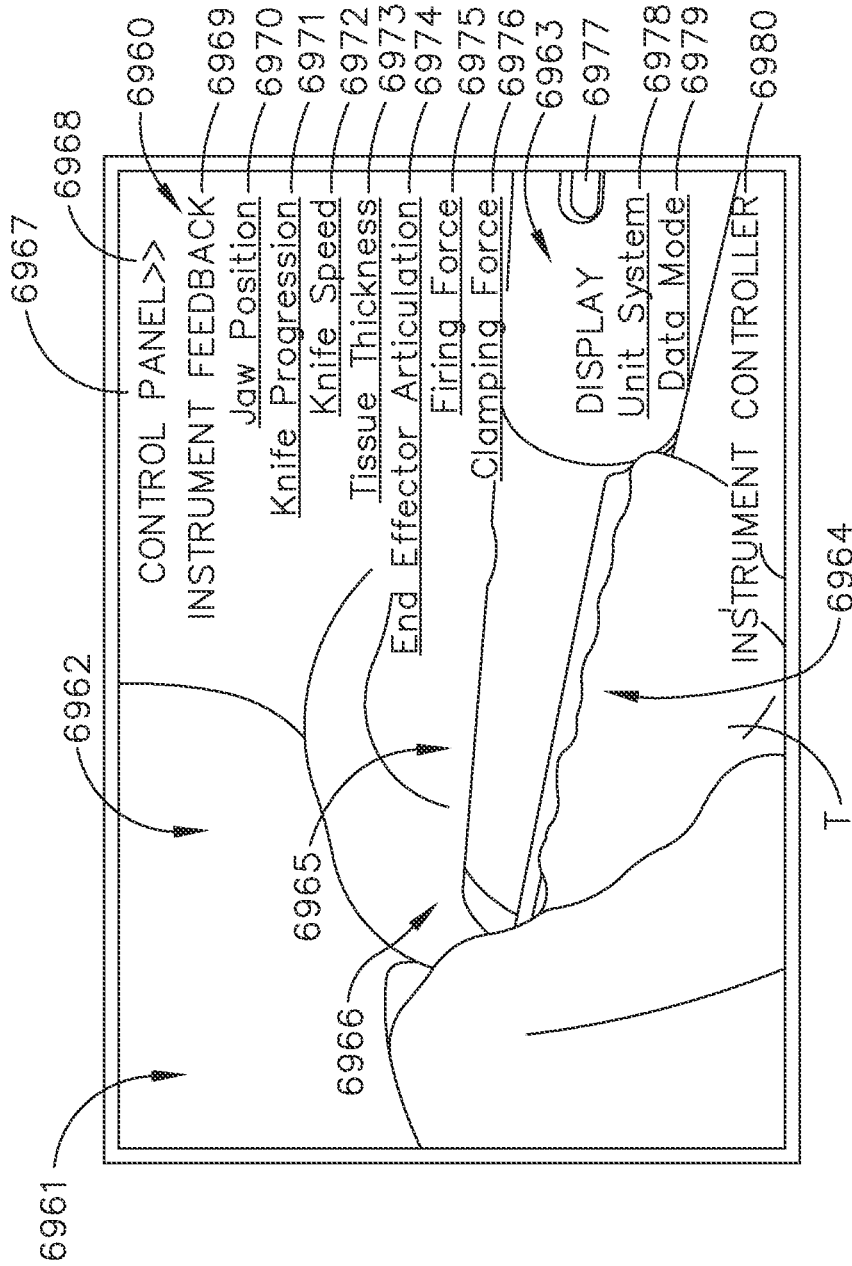


FIG. 148

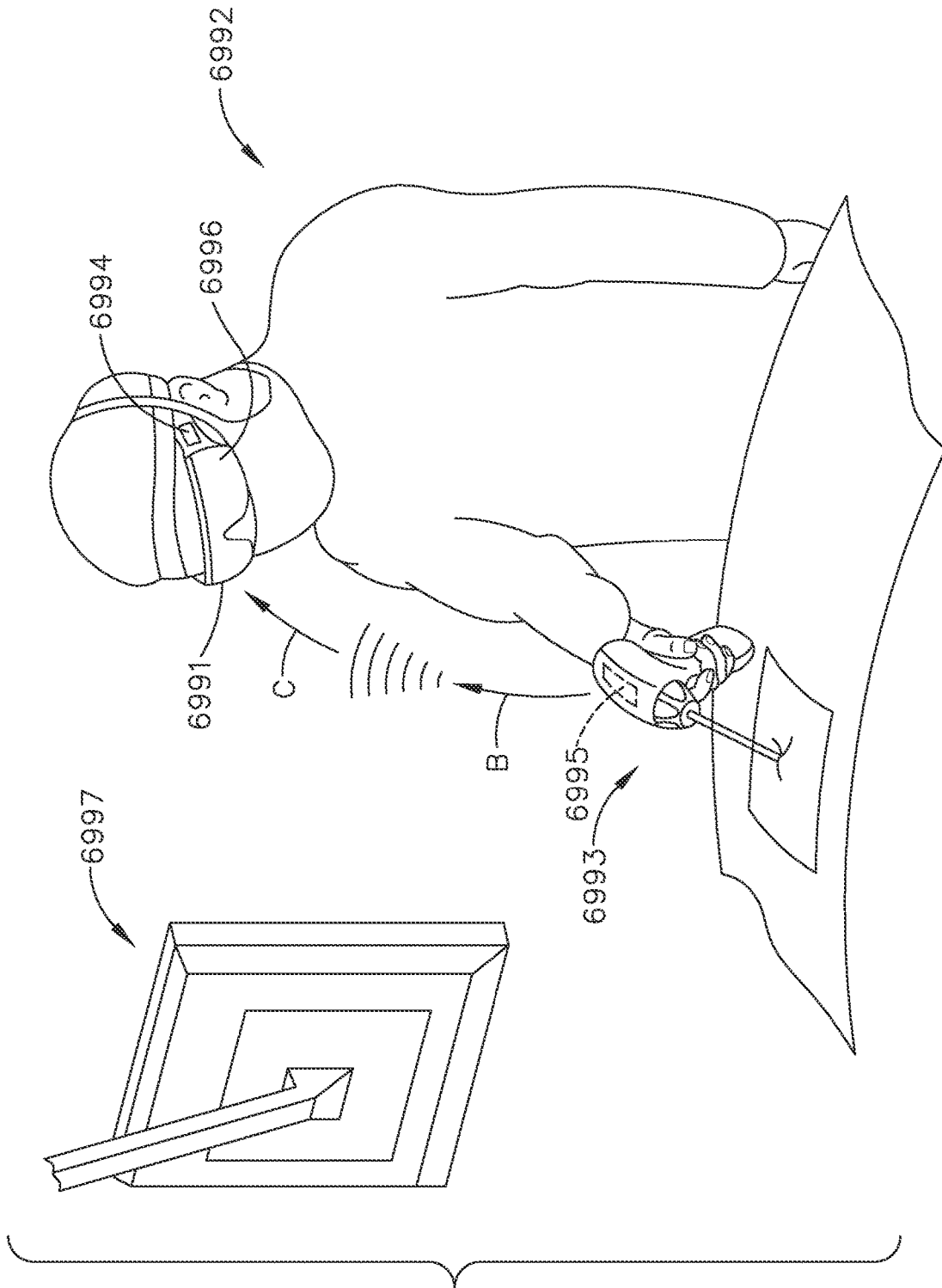


FIG. 149

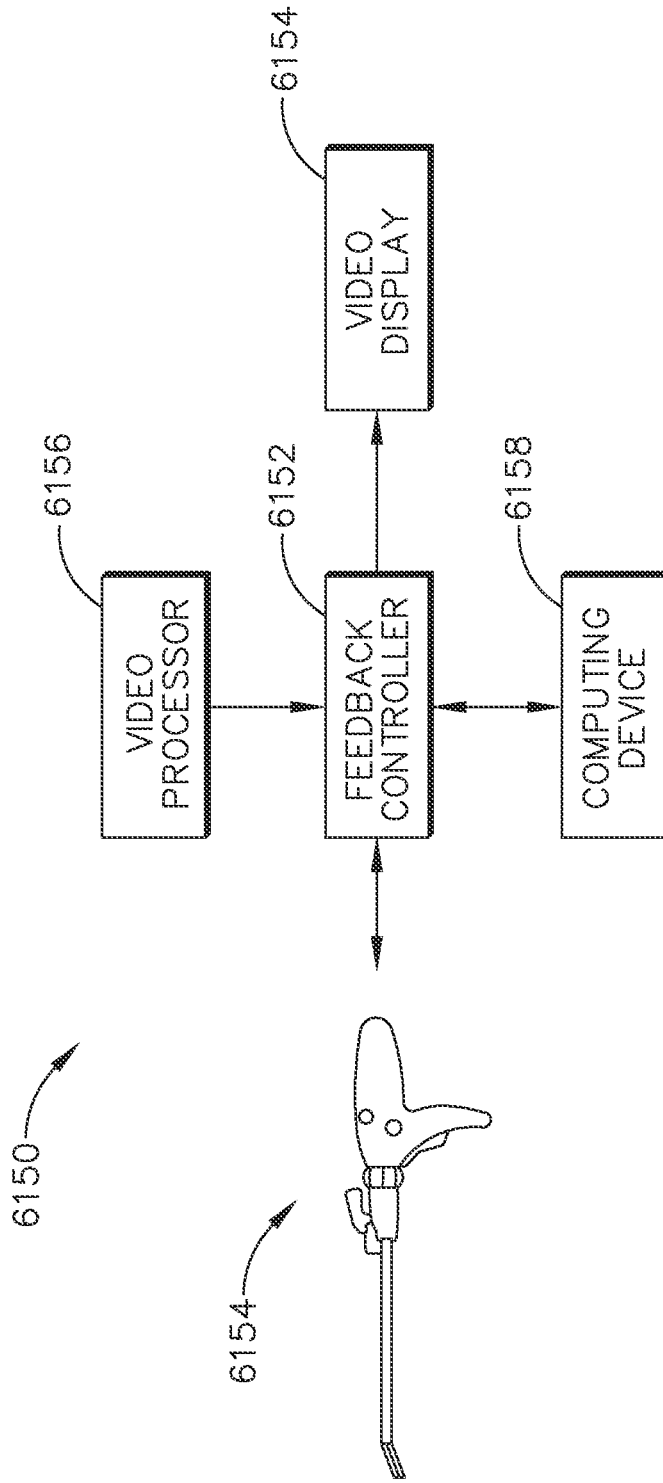


FIG. 150

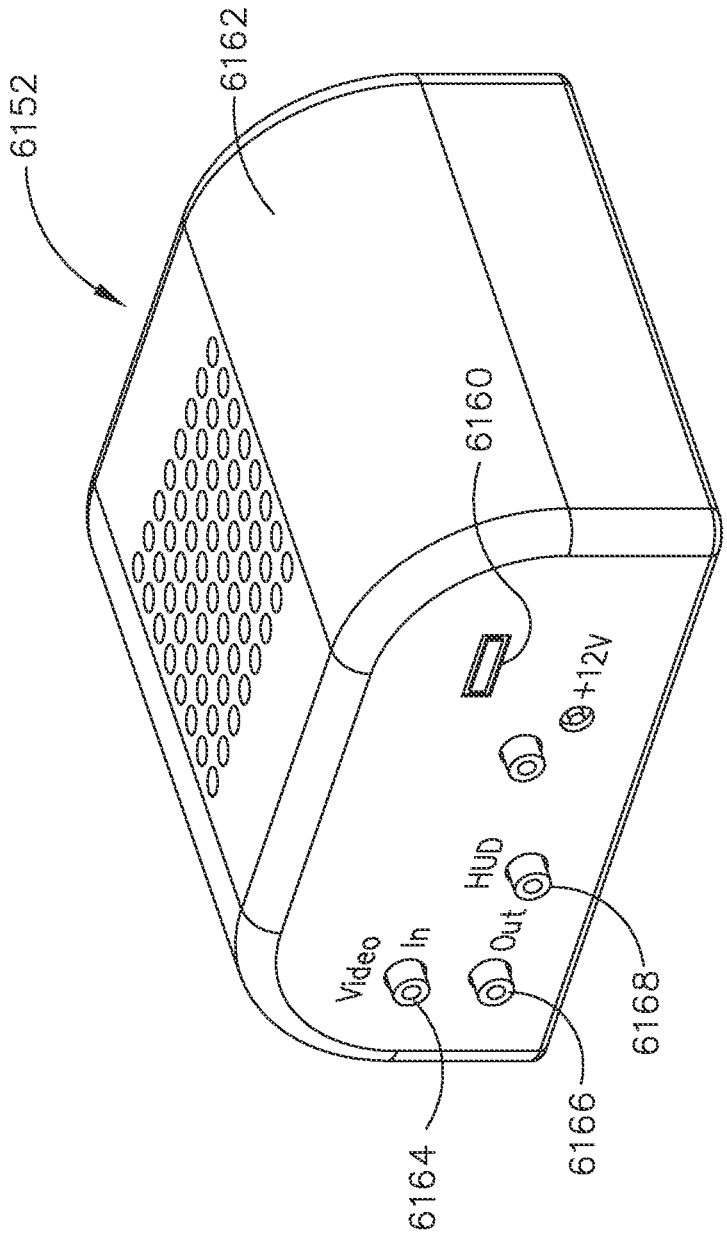


FIG. 151

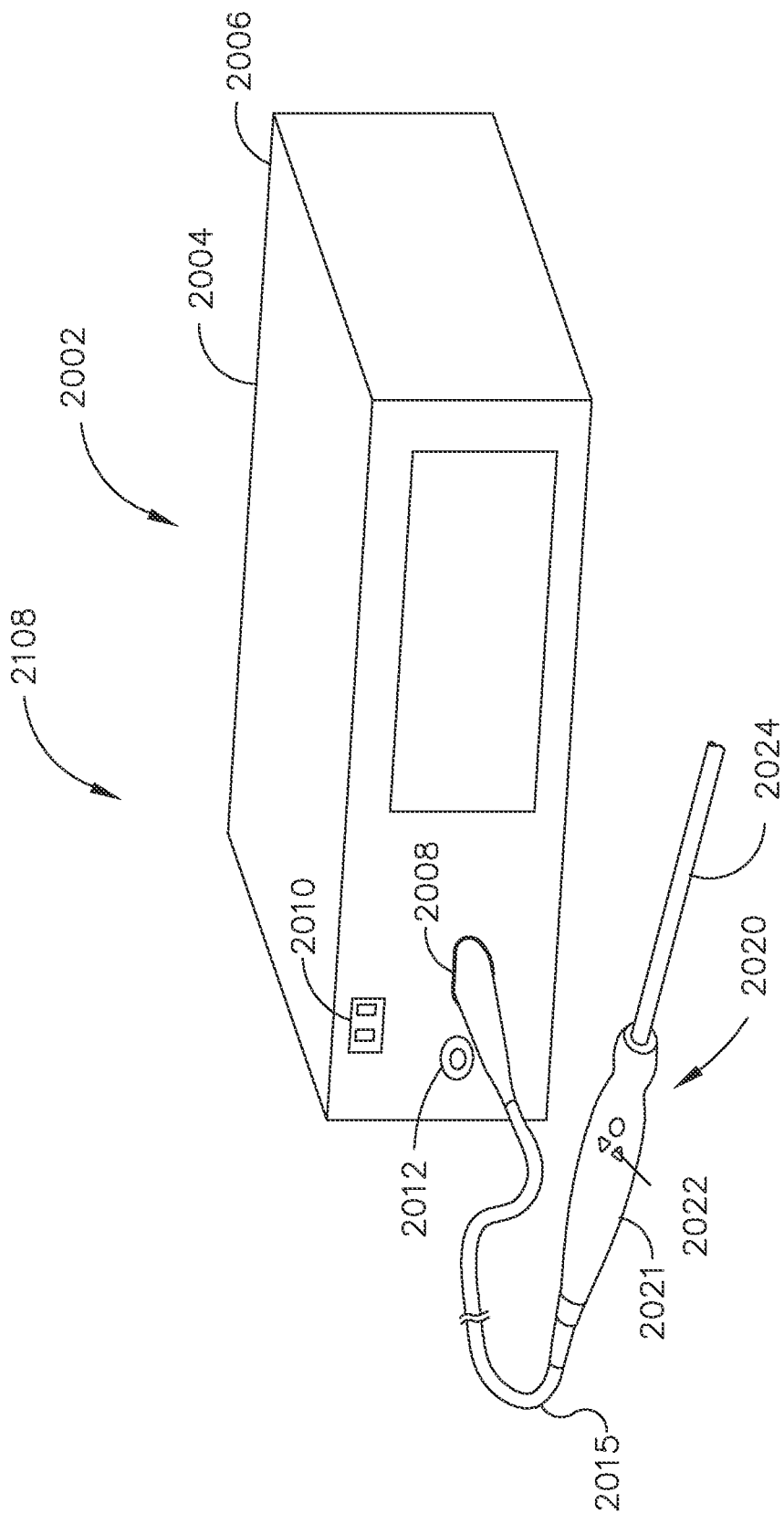


FIG. 152A

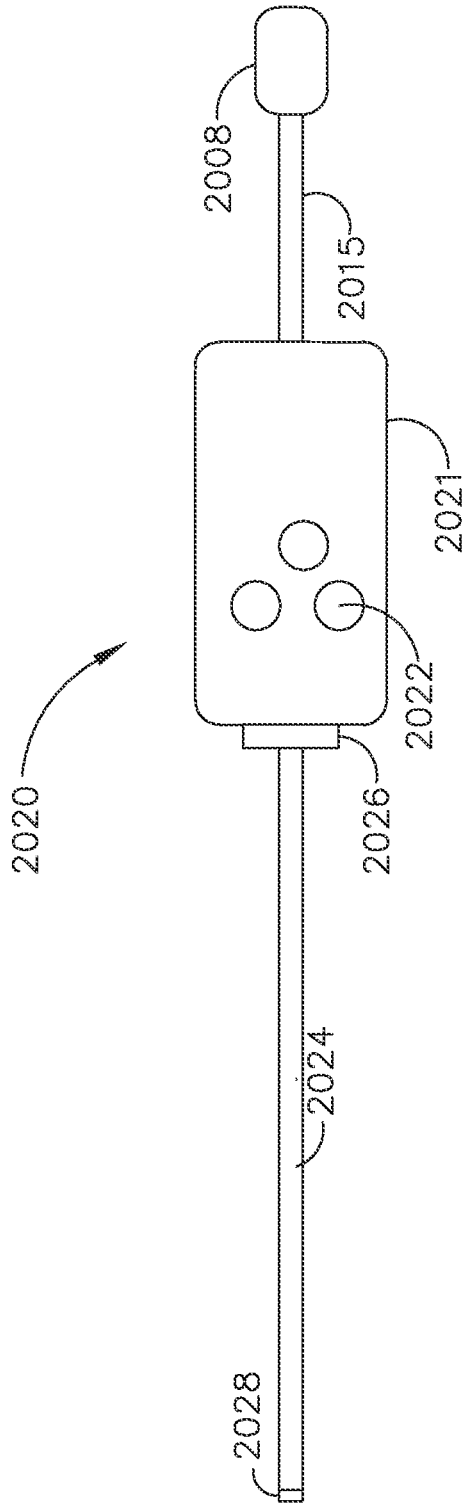


FIG. 152B

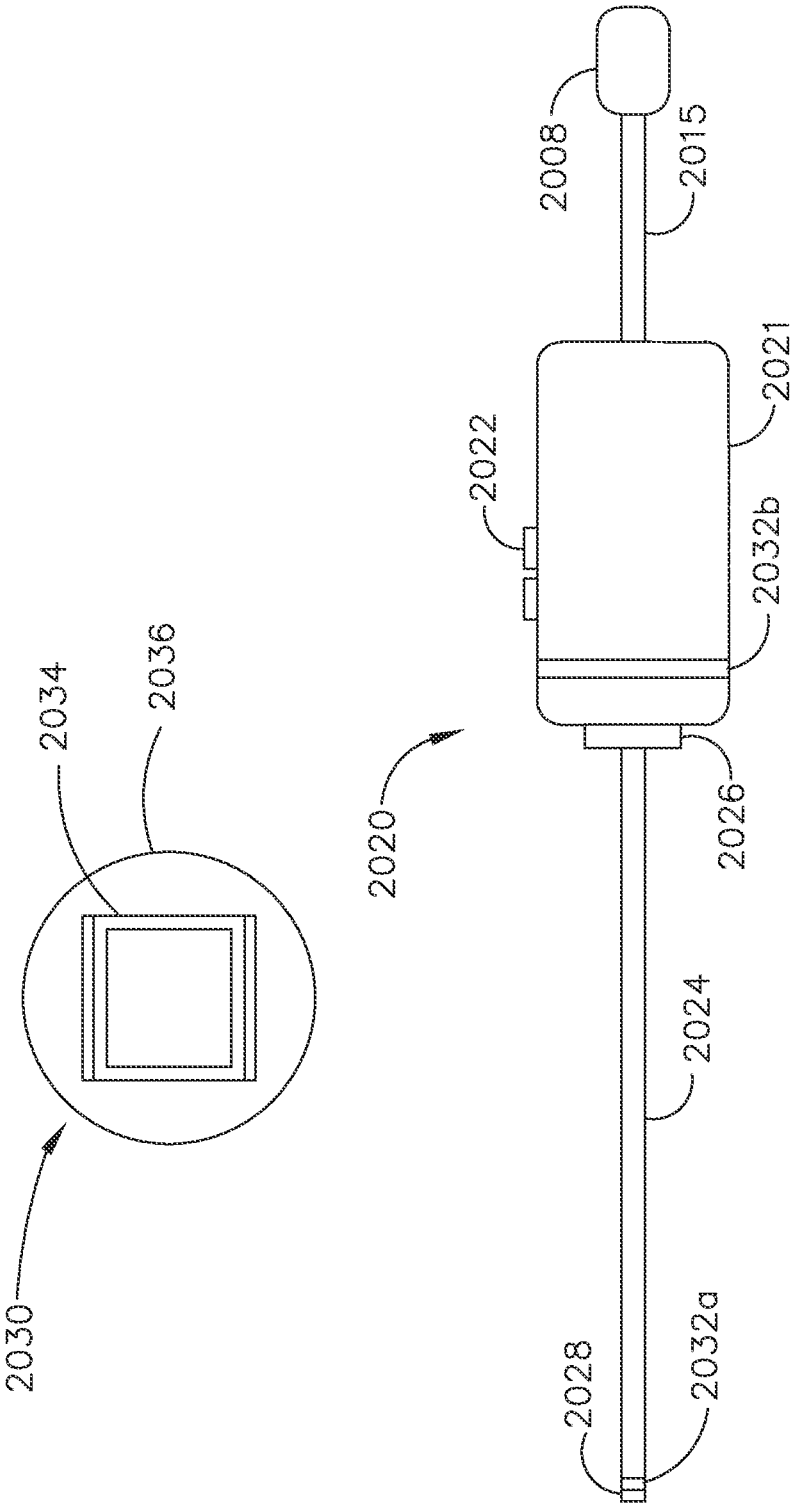


FIG. 152C

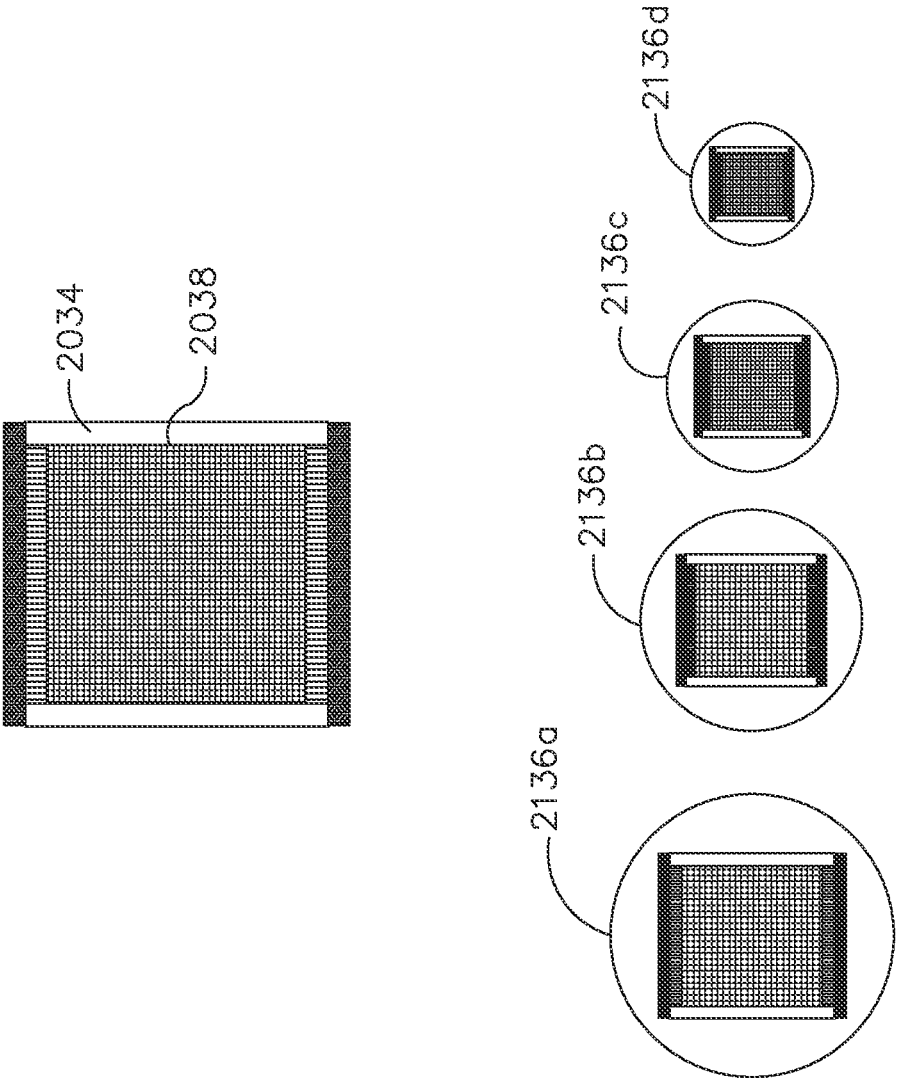


FIG. 152D



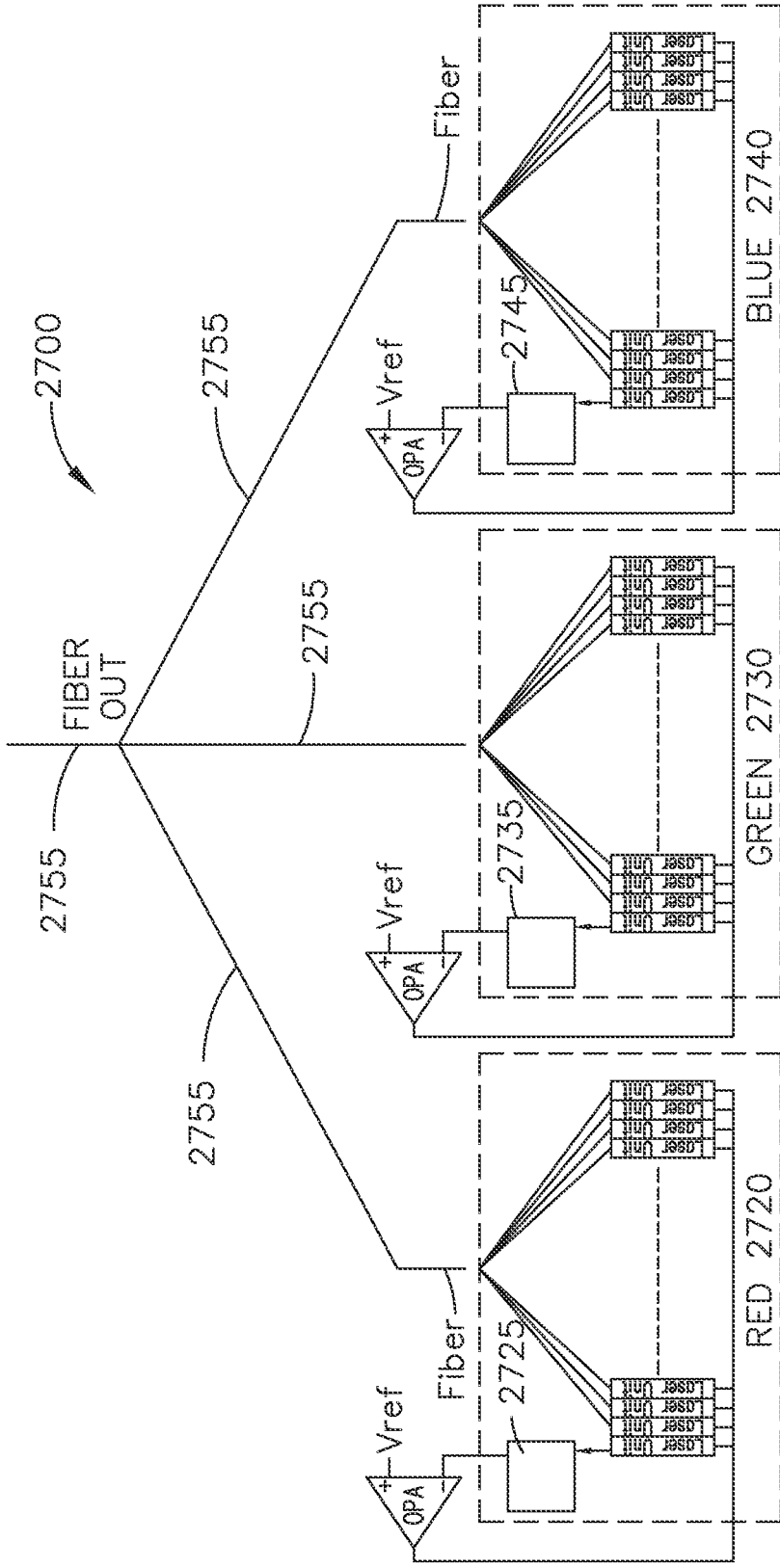


FIG. 153A

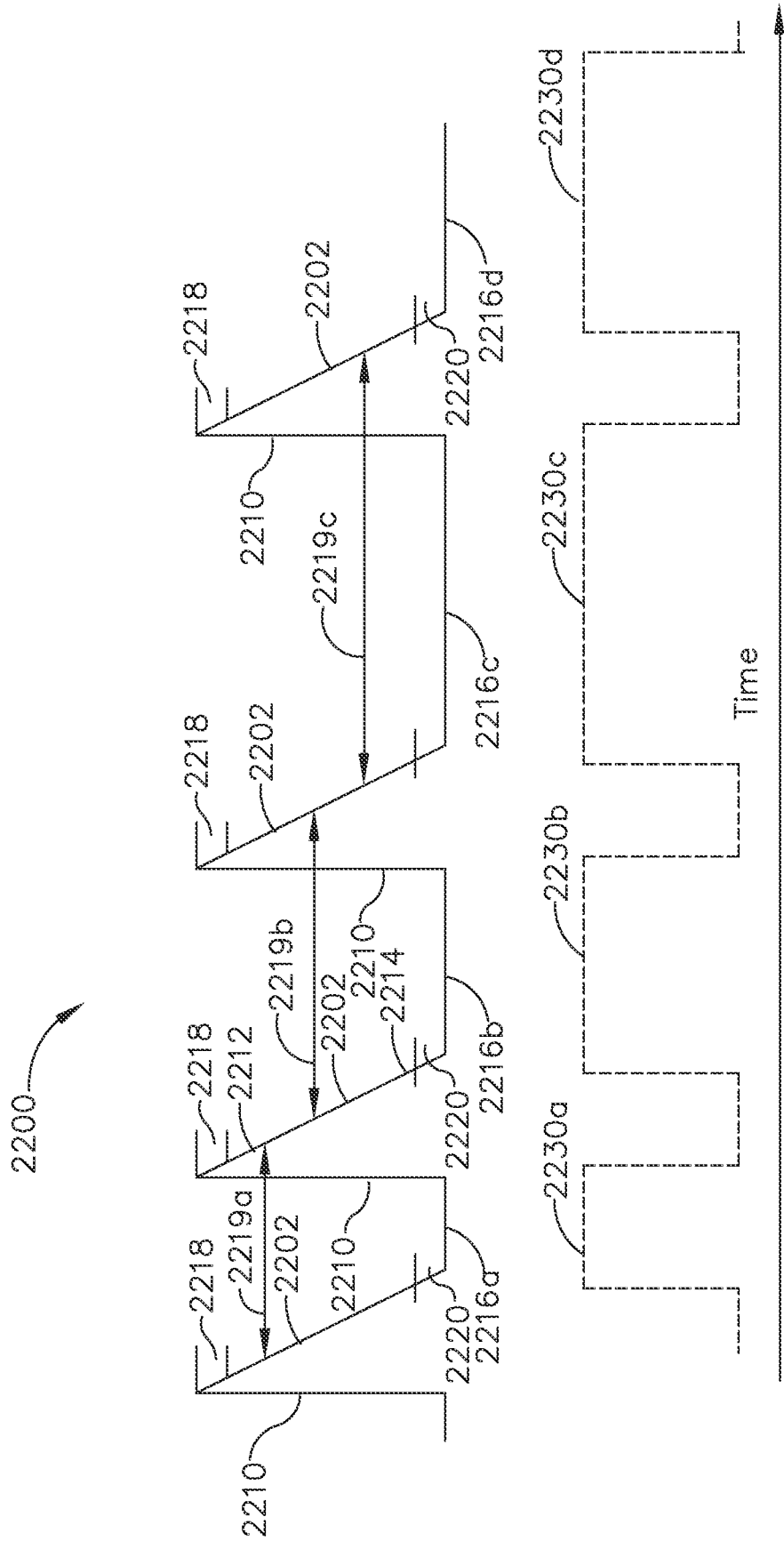


FIG. 153B

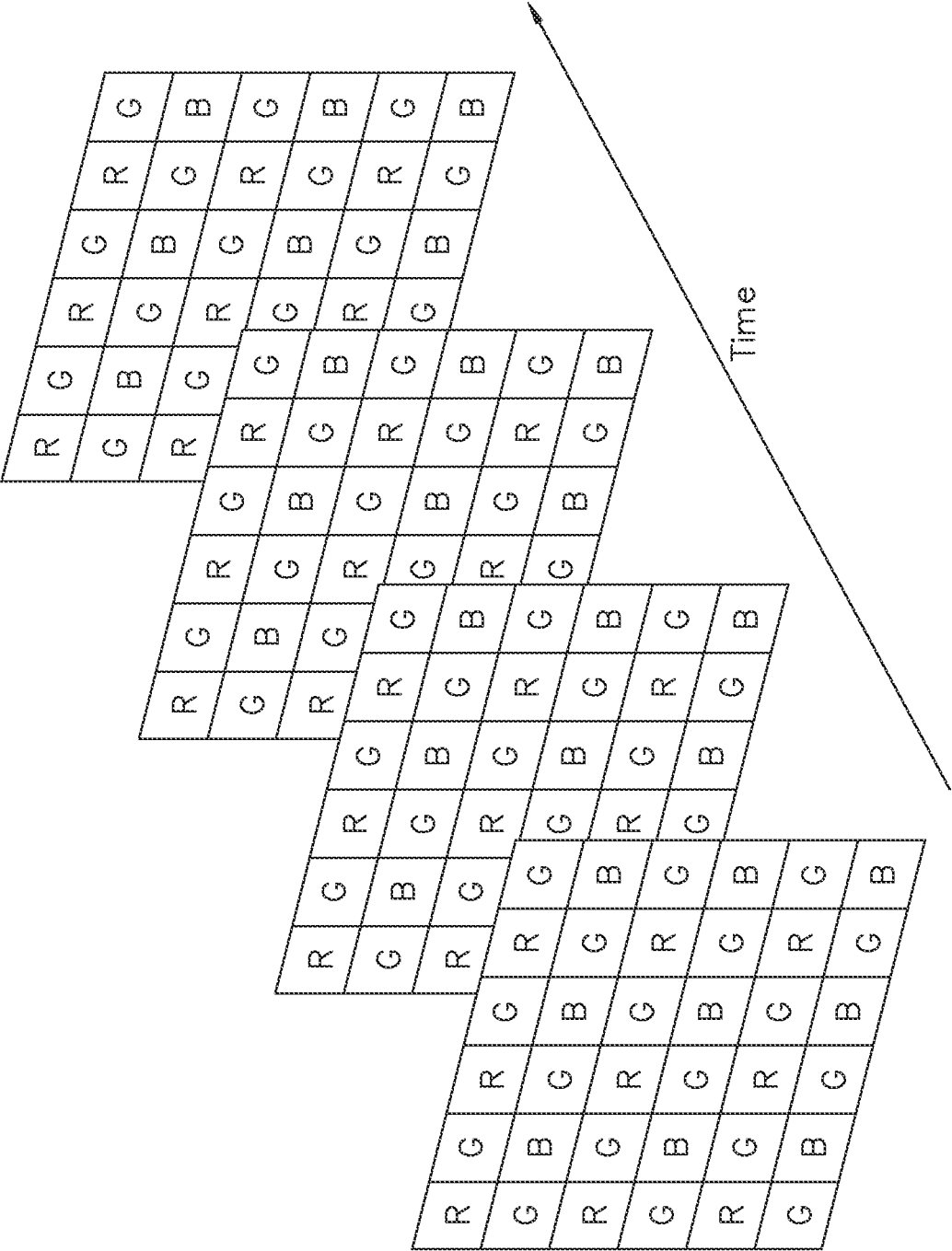


FIG. 153C

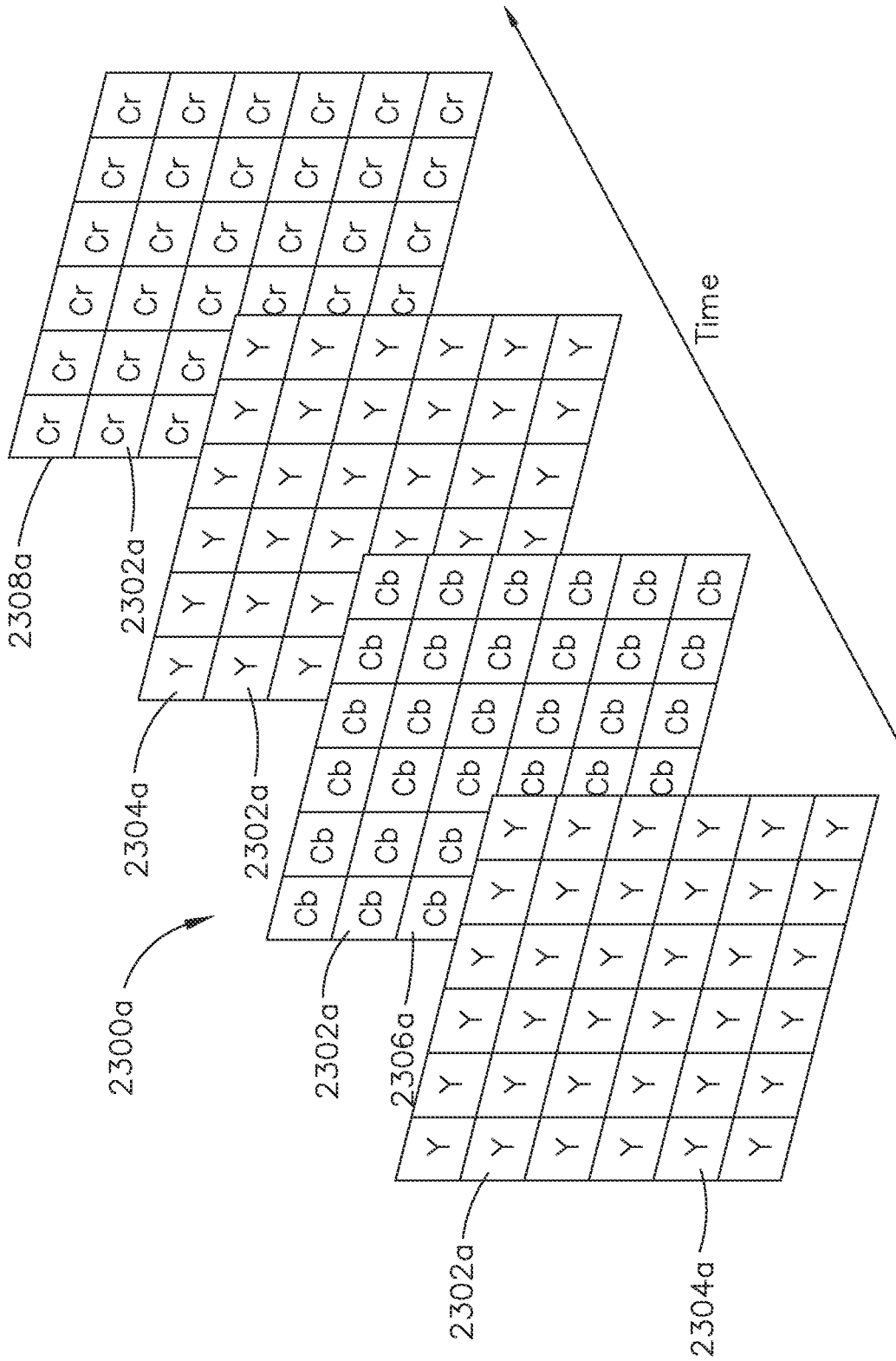


FIG. 153D

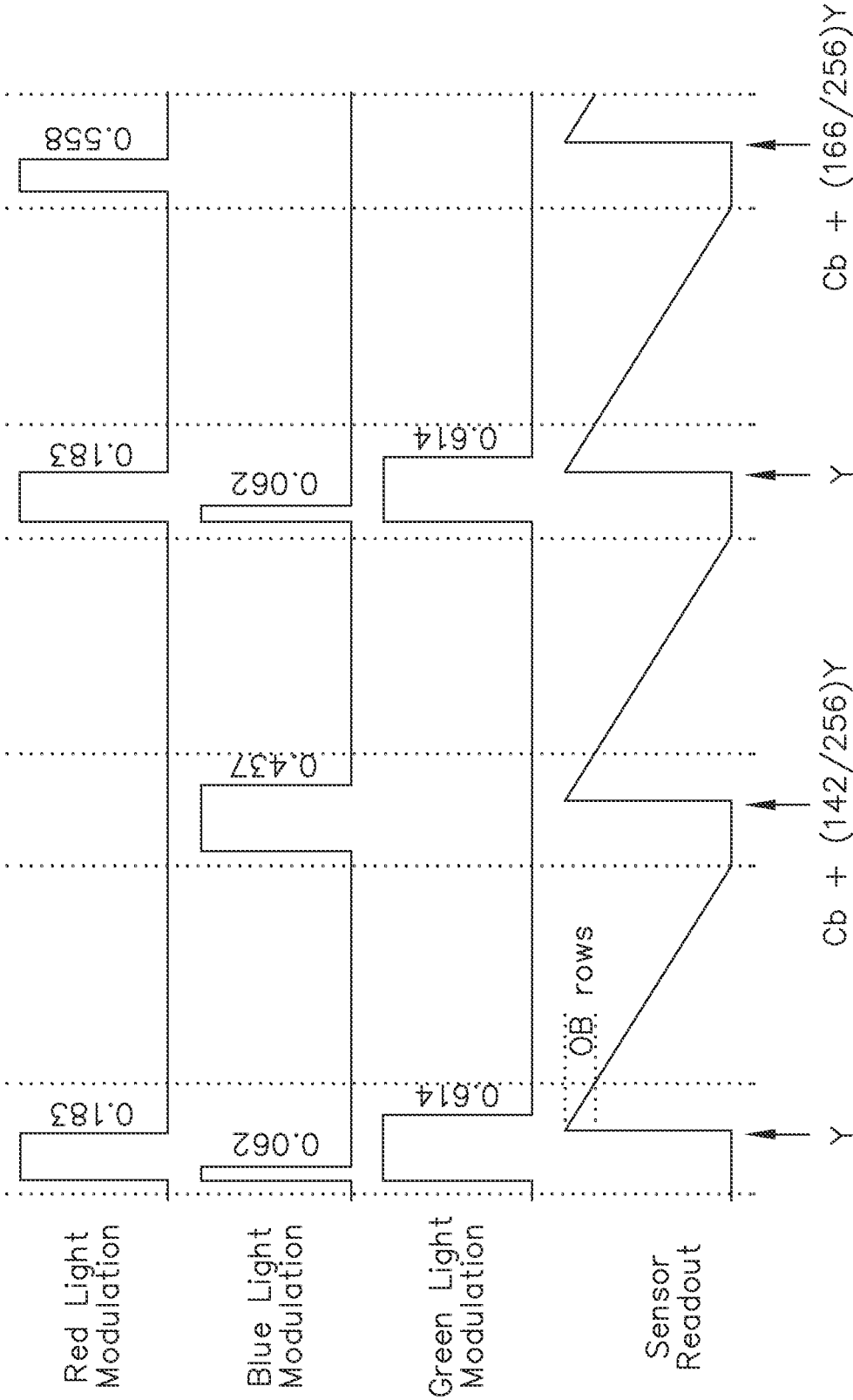


FIG. 153E

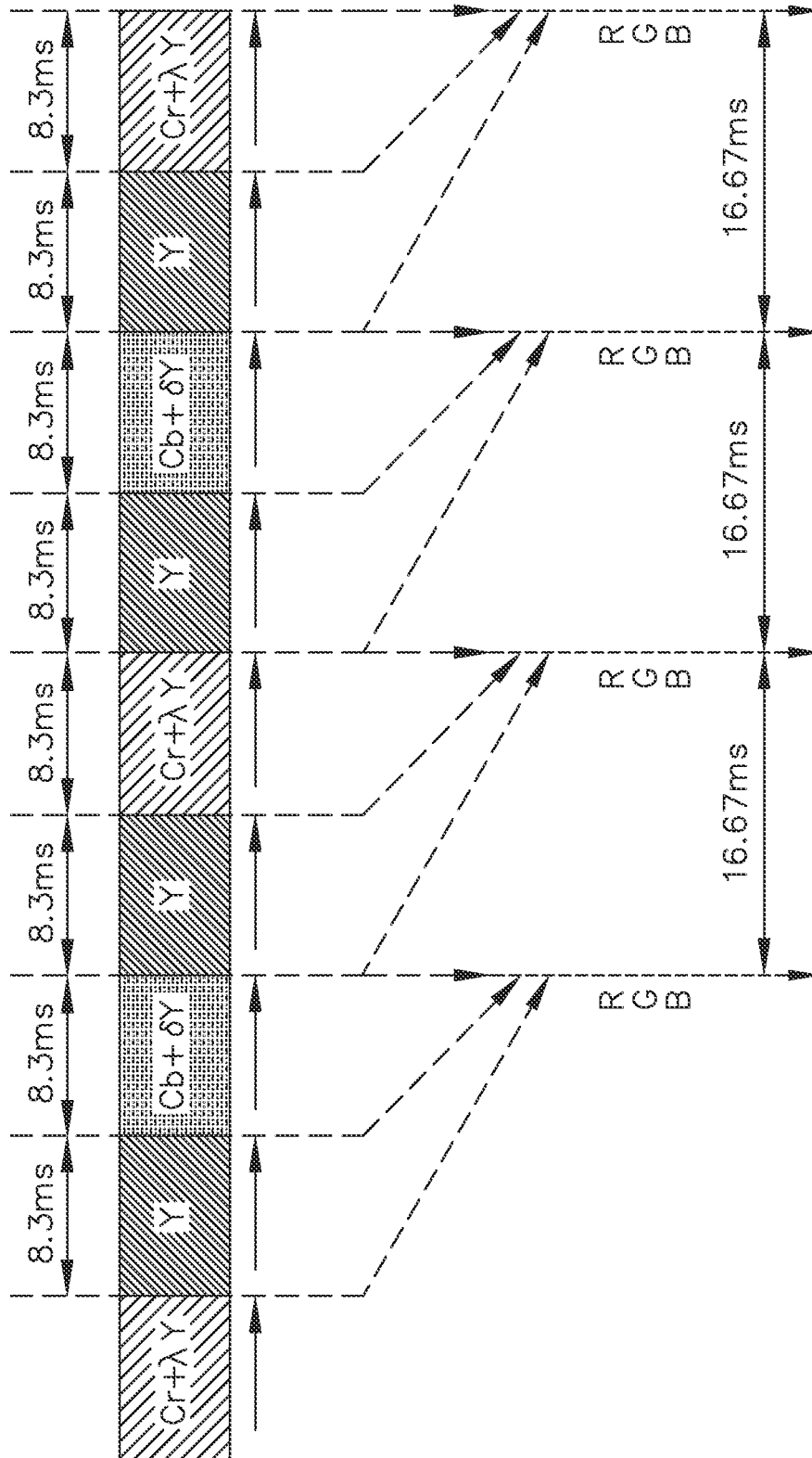


FIG. 153F

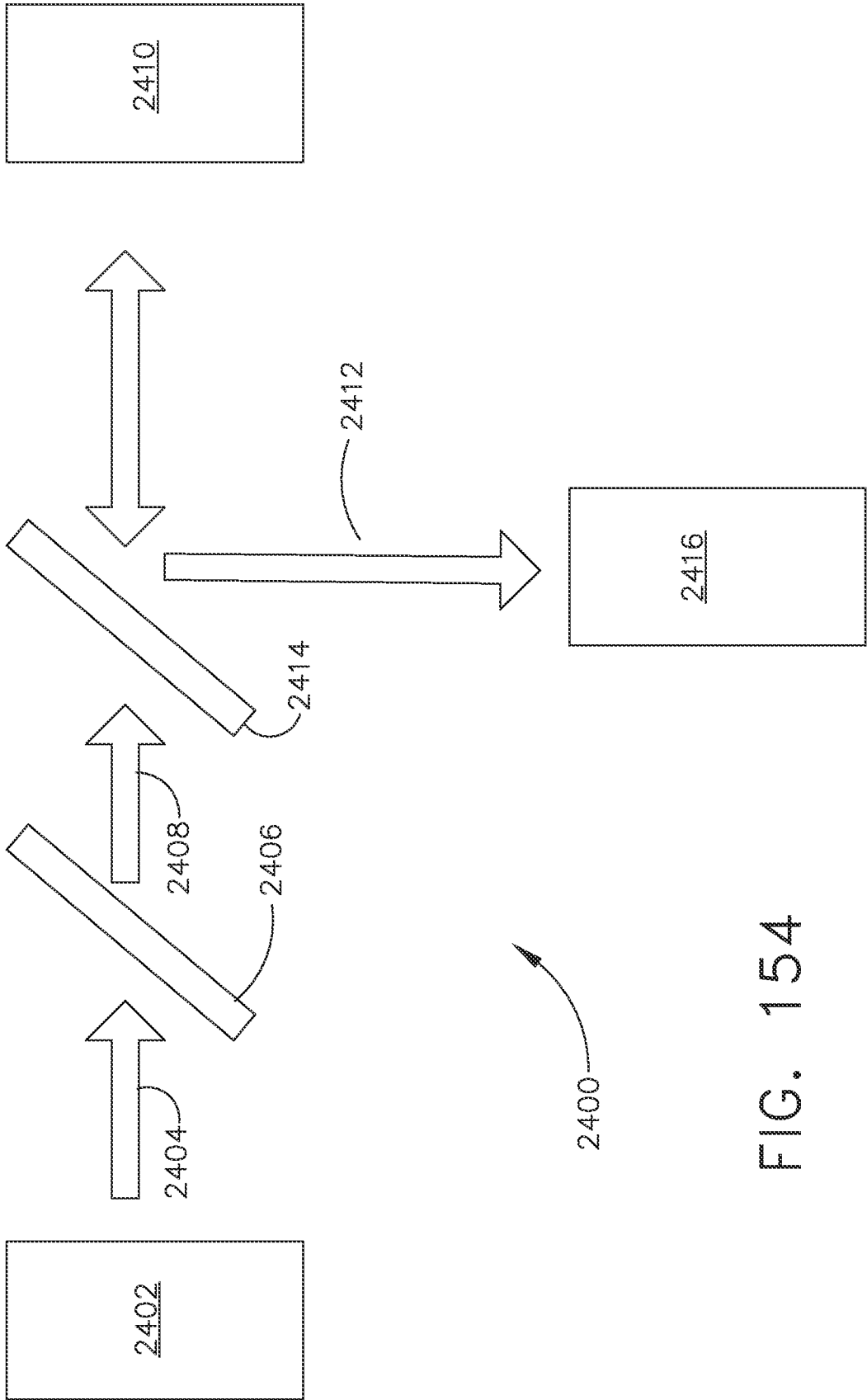


FIG. 154

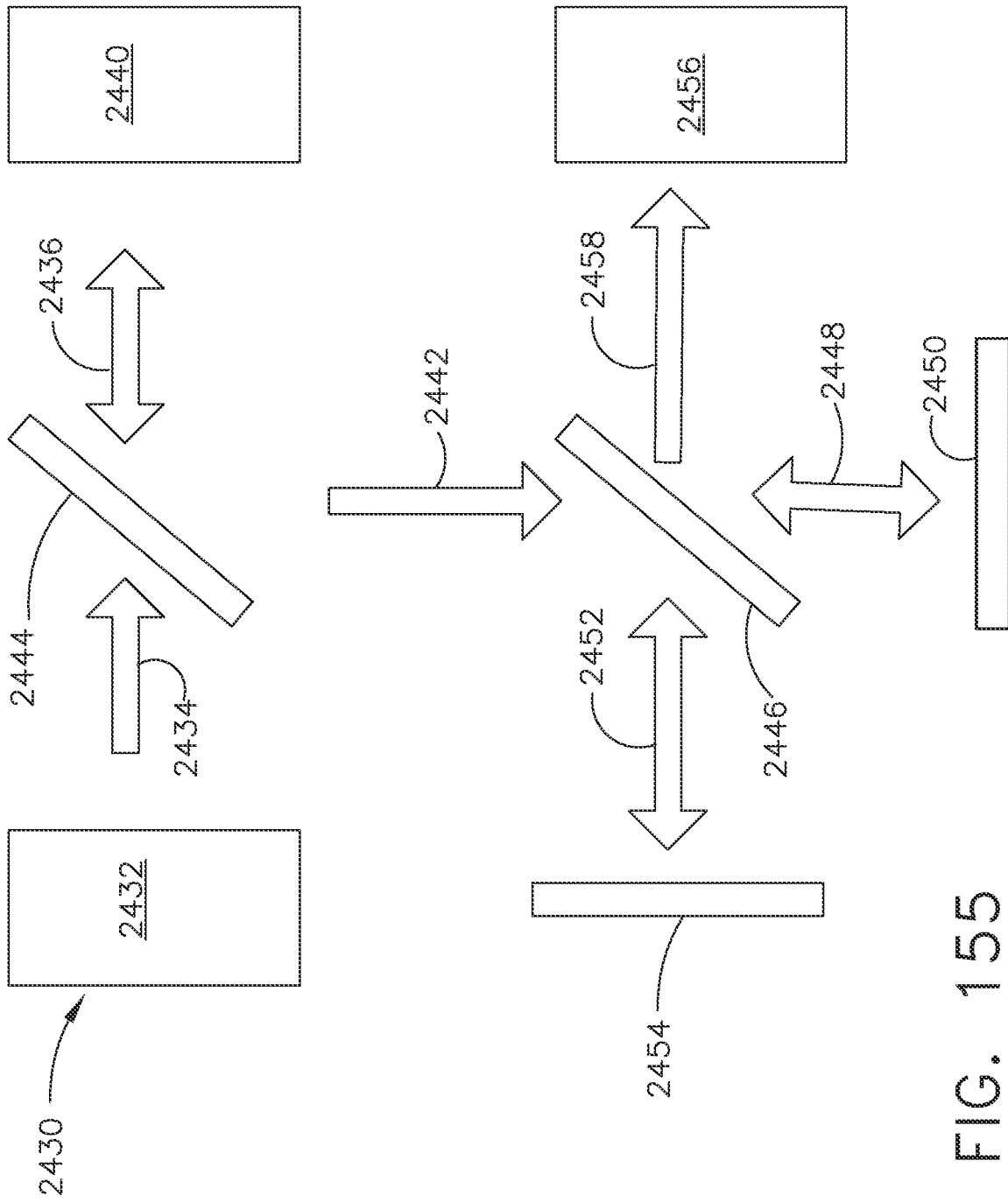


FIG. 155



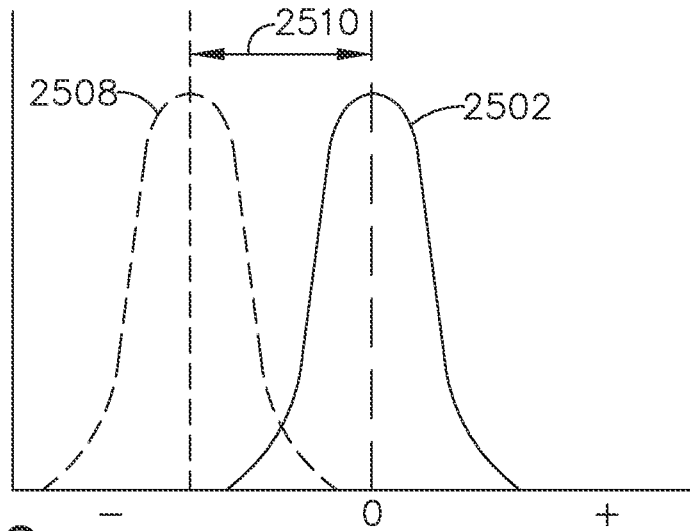


FIG. 156C

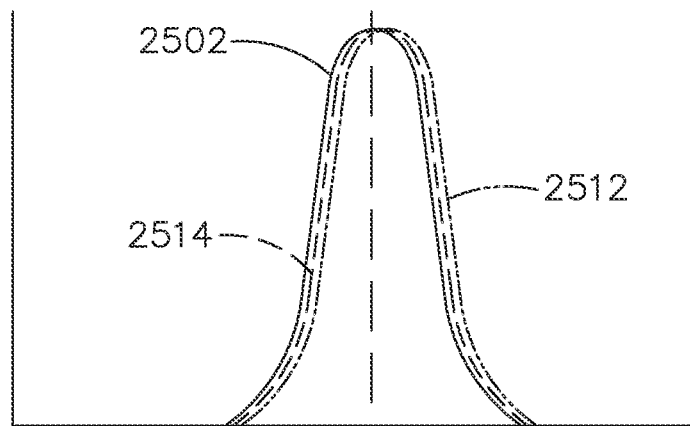


FIG. 156B

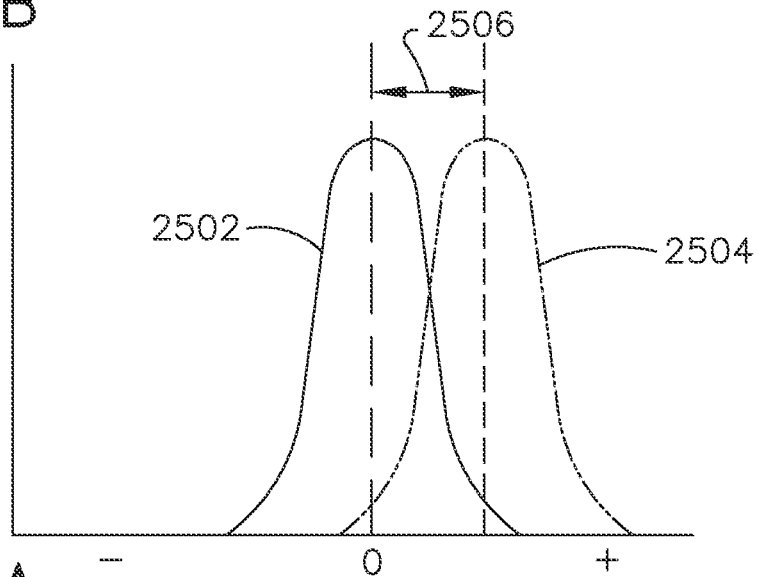


FIG. 156A

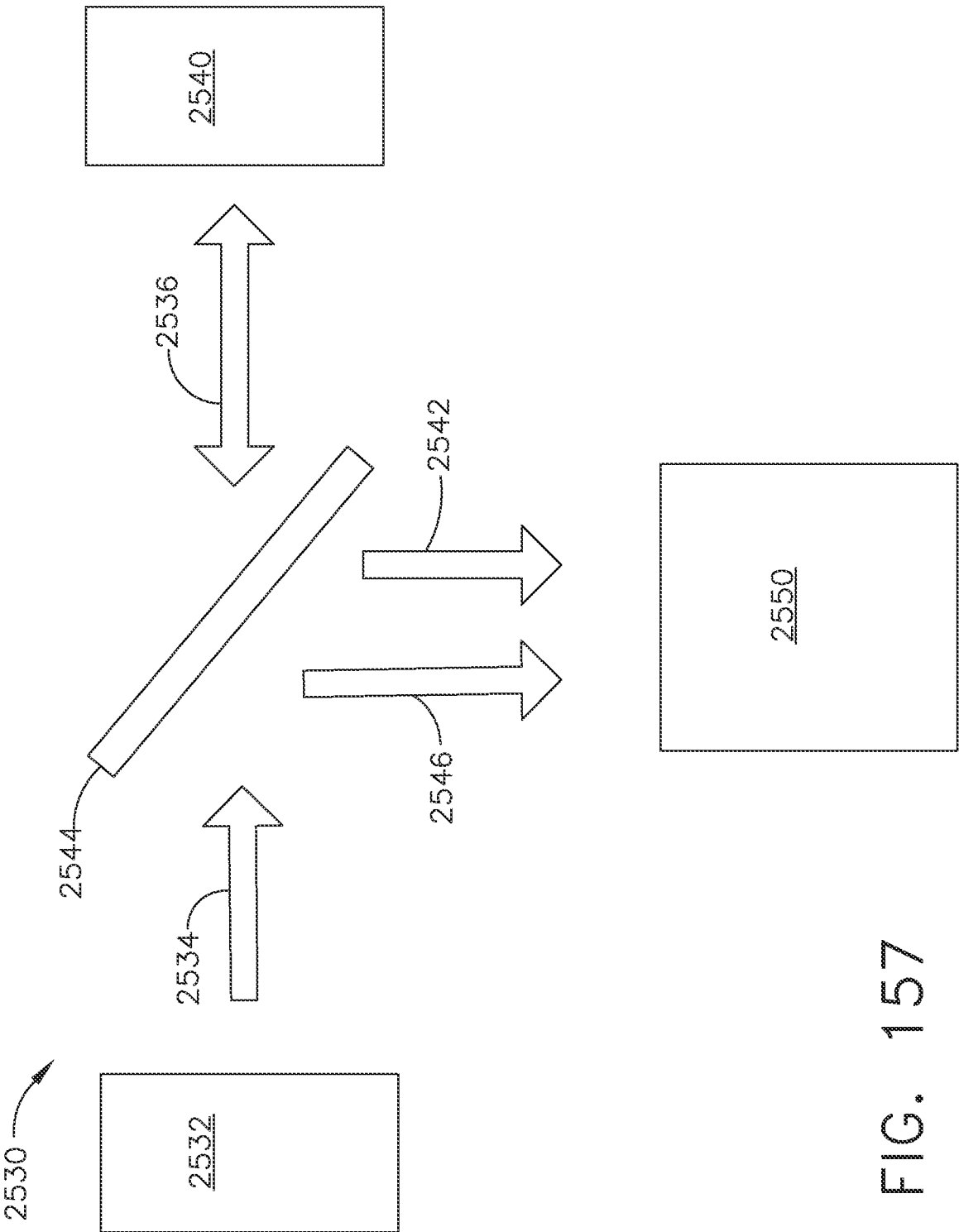


FIG. 157

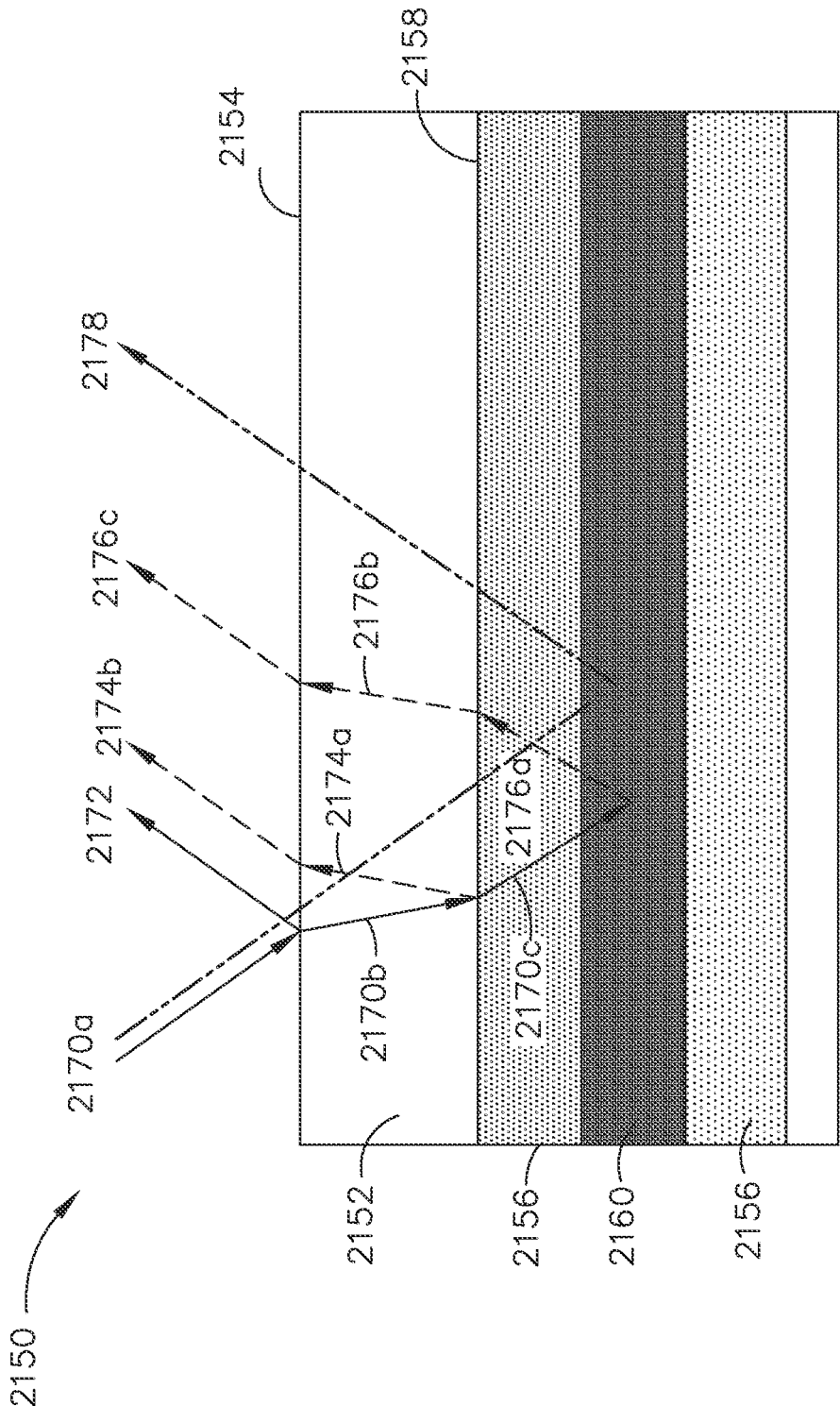


FIG. 158

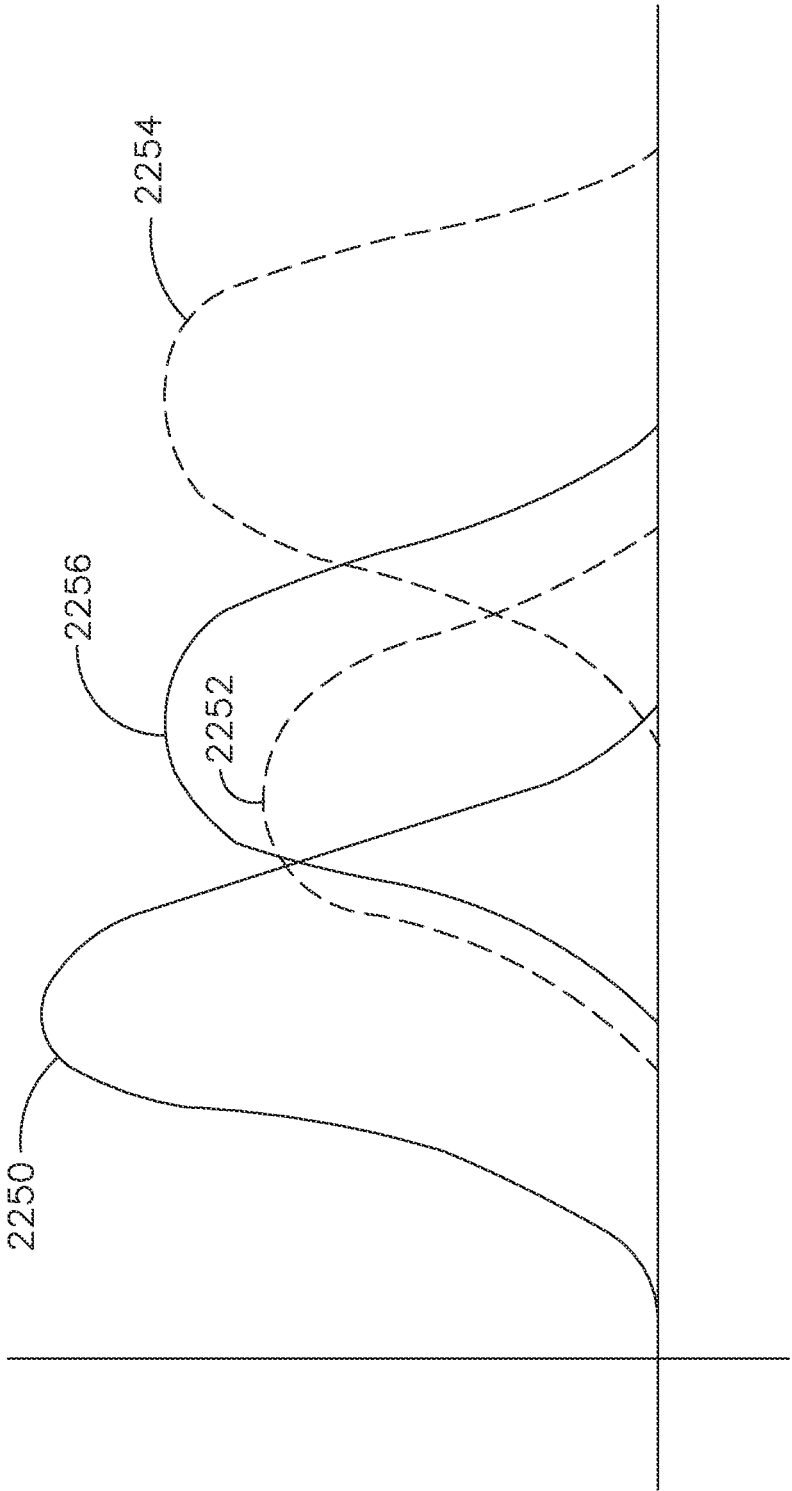
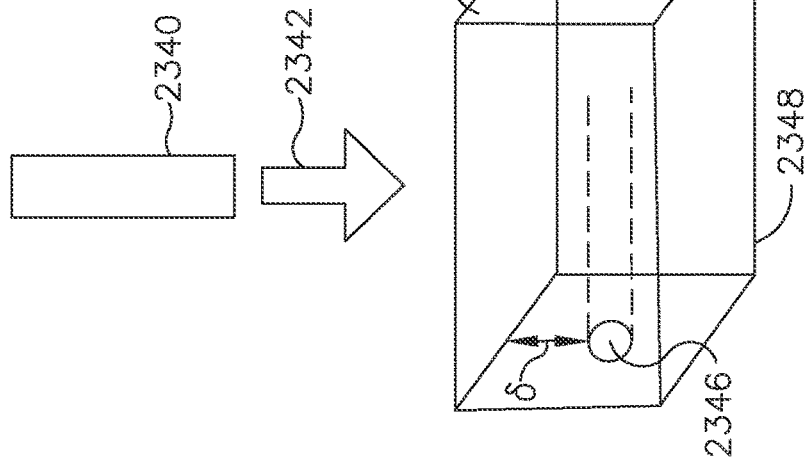
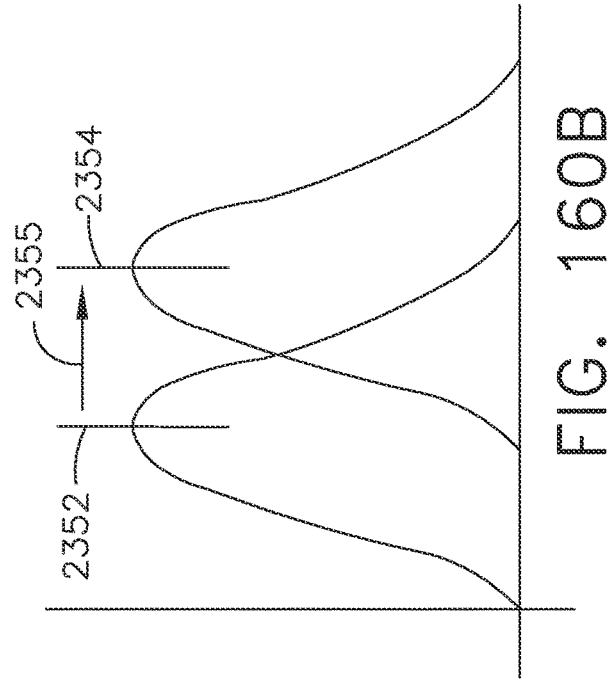
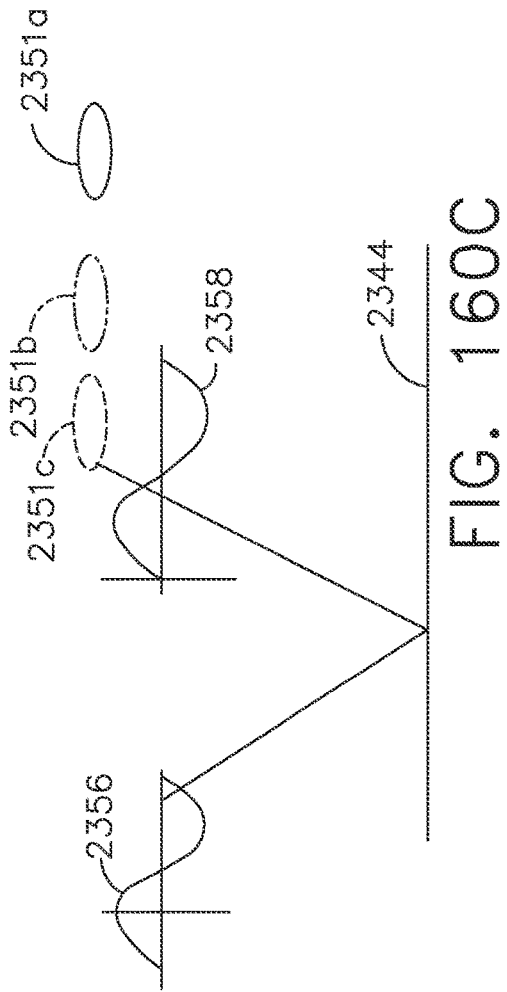


FIG. 159



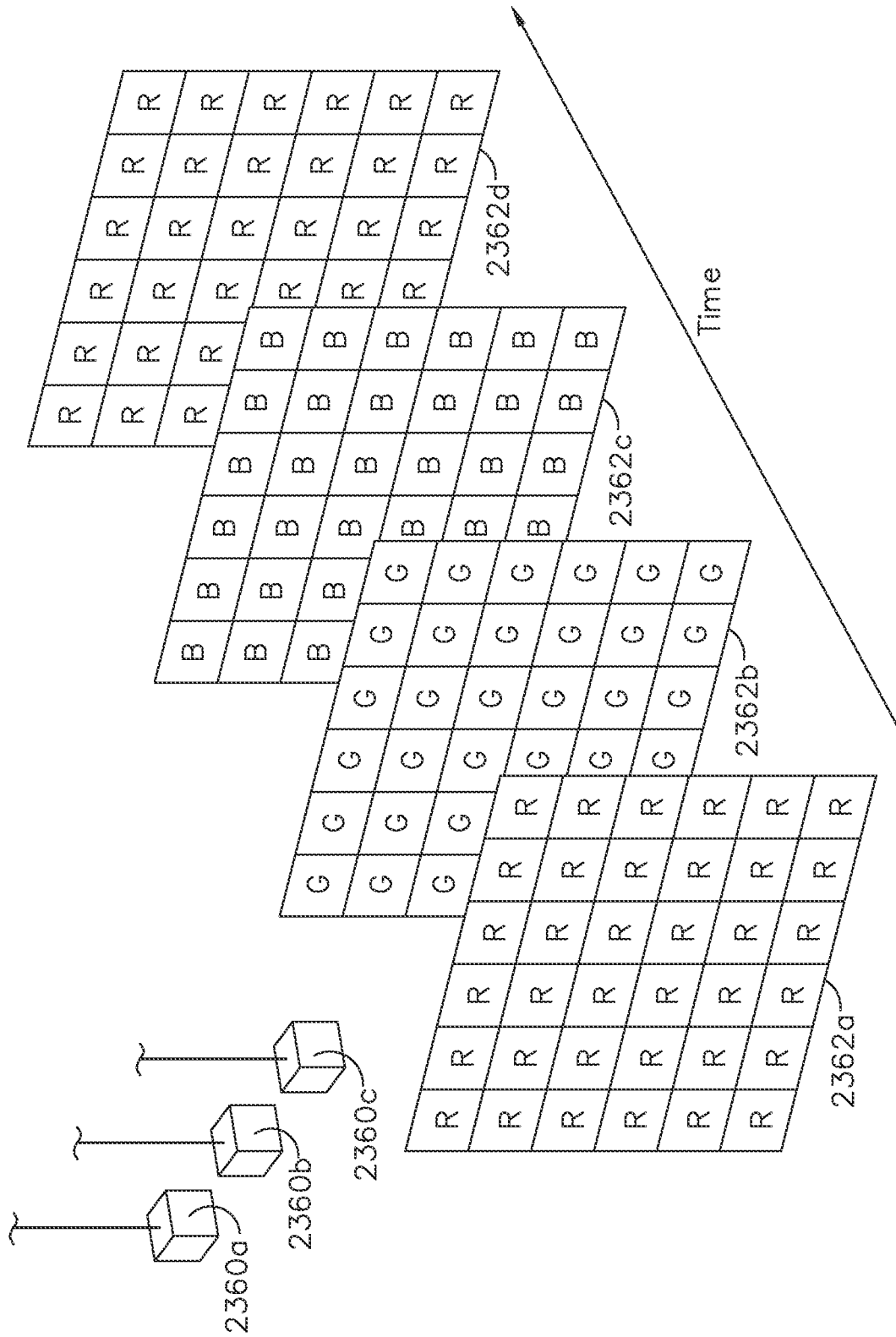


FIG. 160D

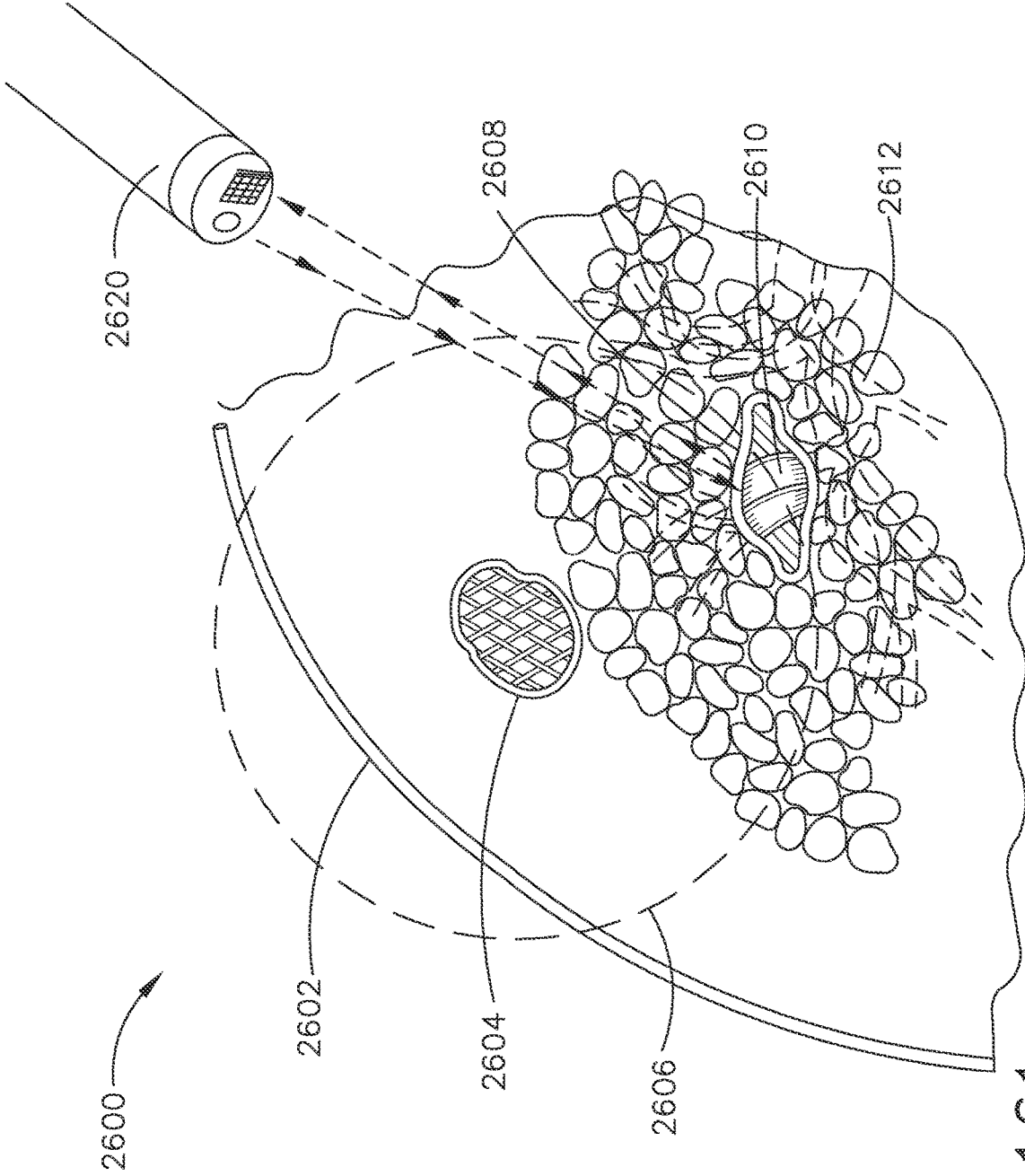


FIG. 161

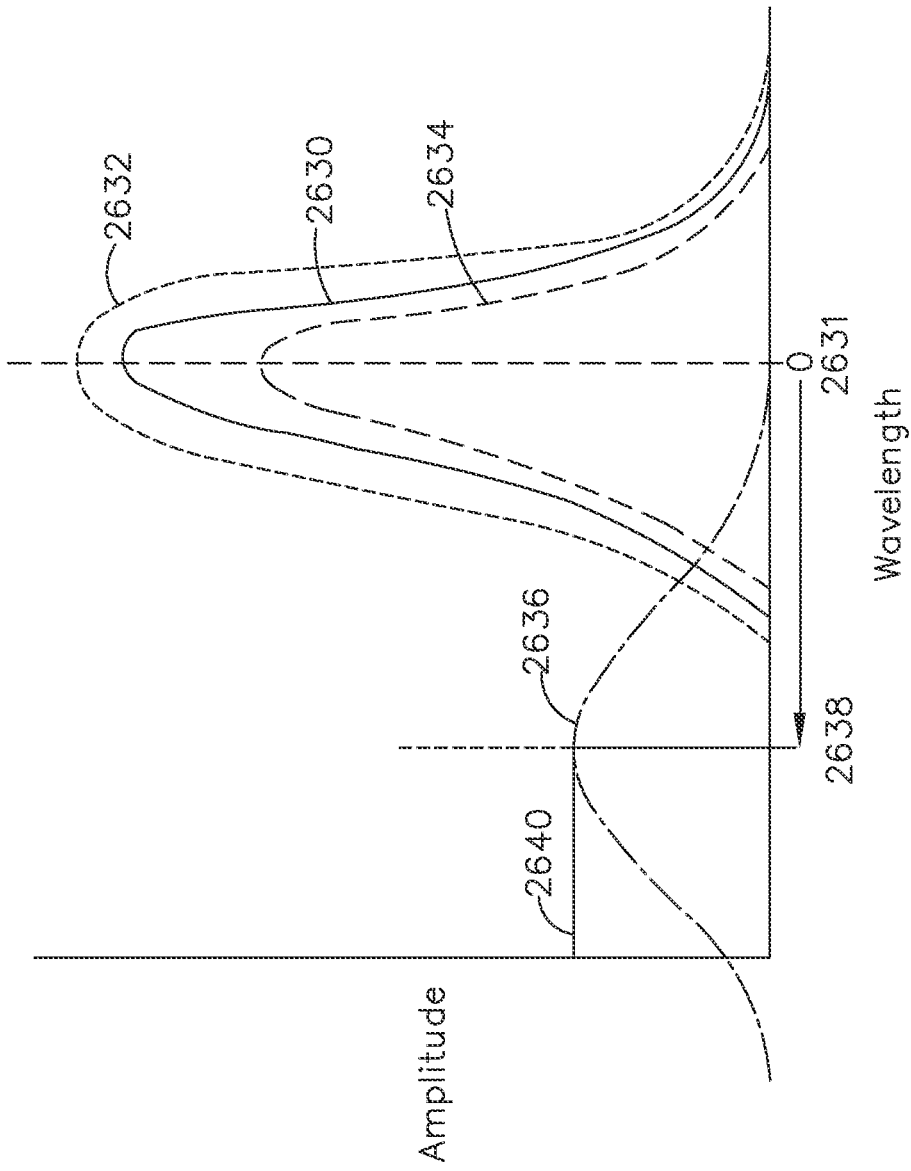


FIG. 162



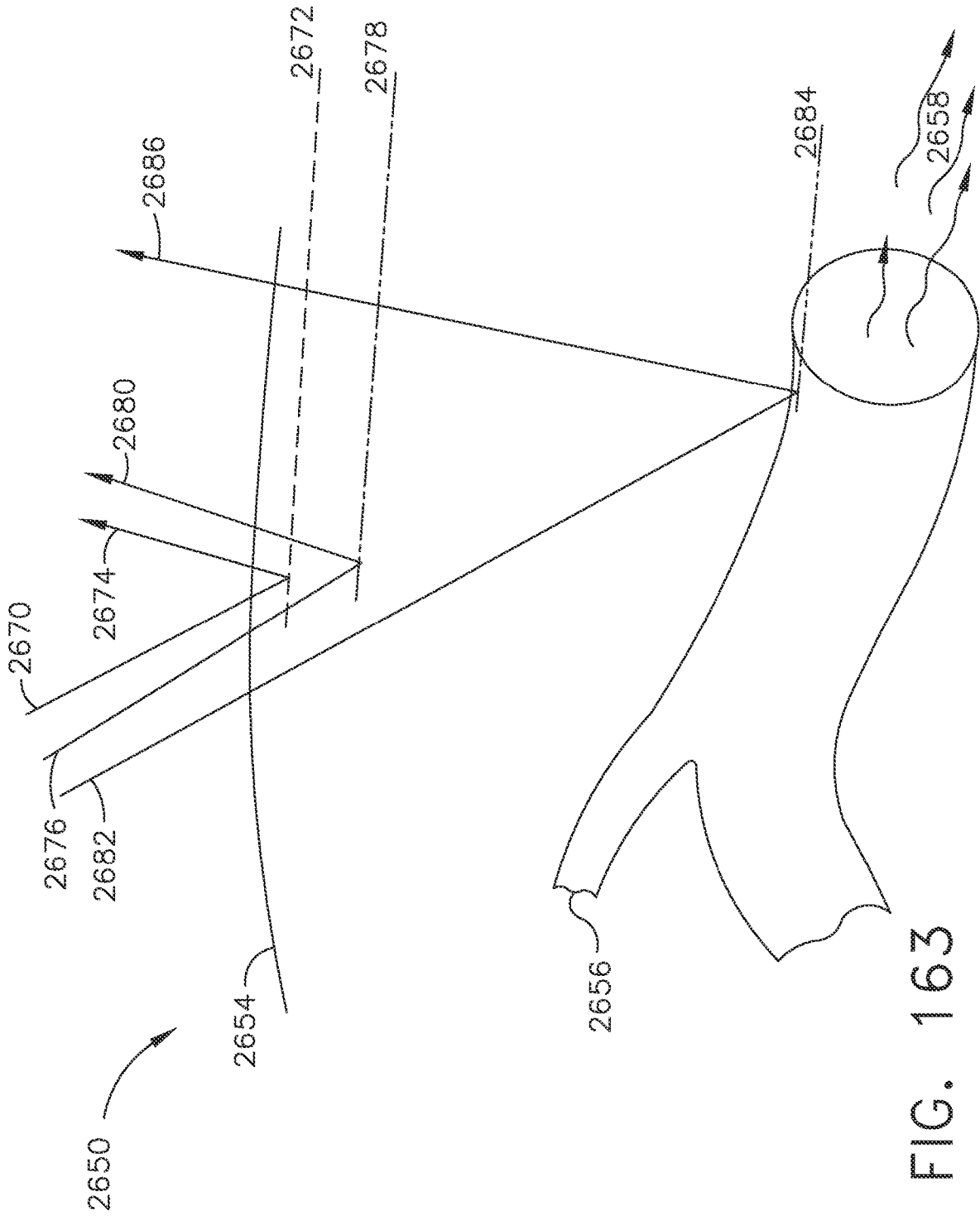


FIG. 163

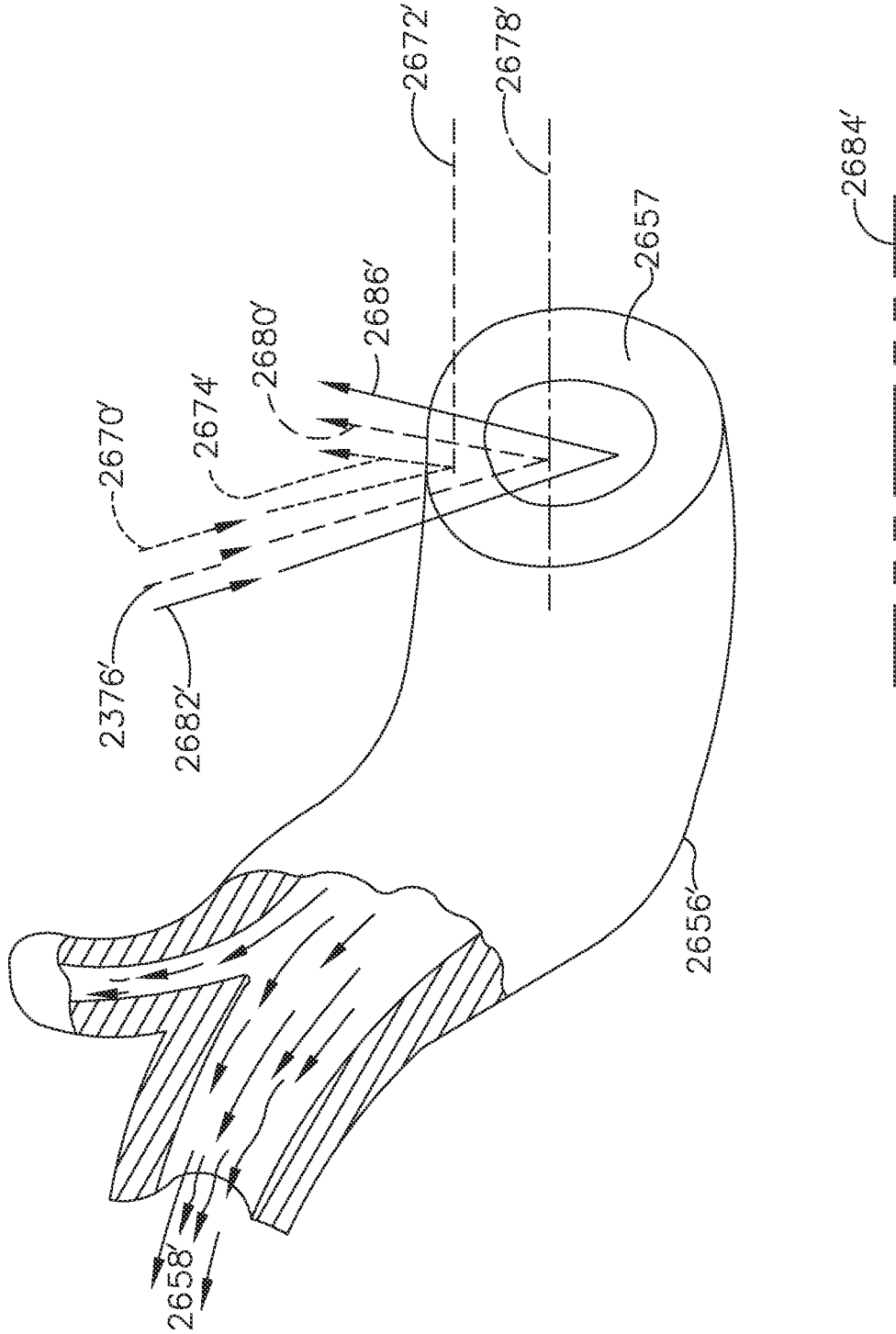


FIG. 164



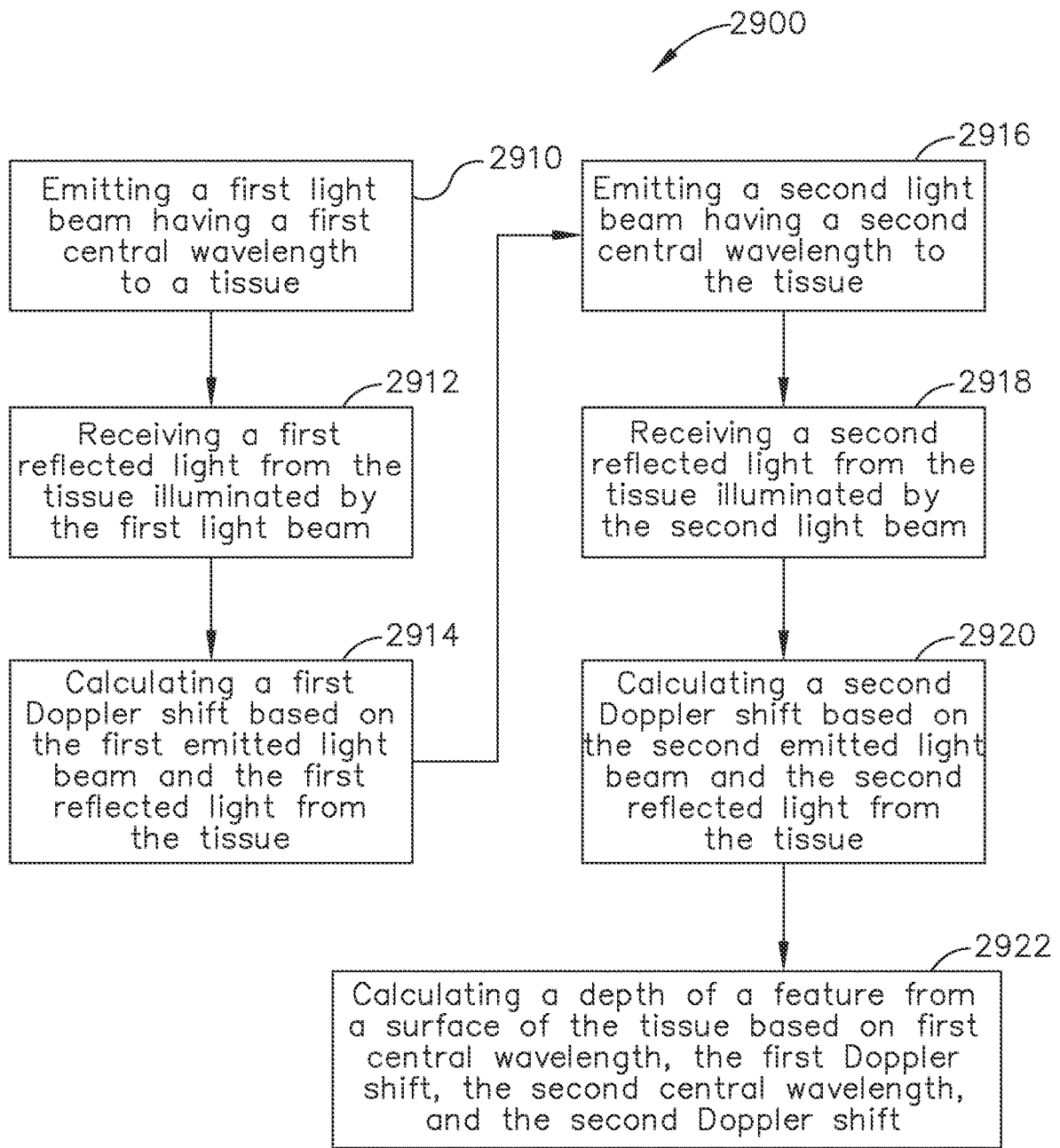


FIG. 166

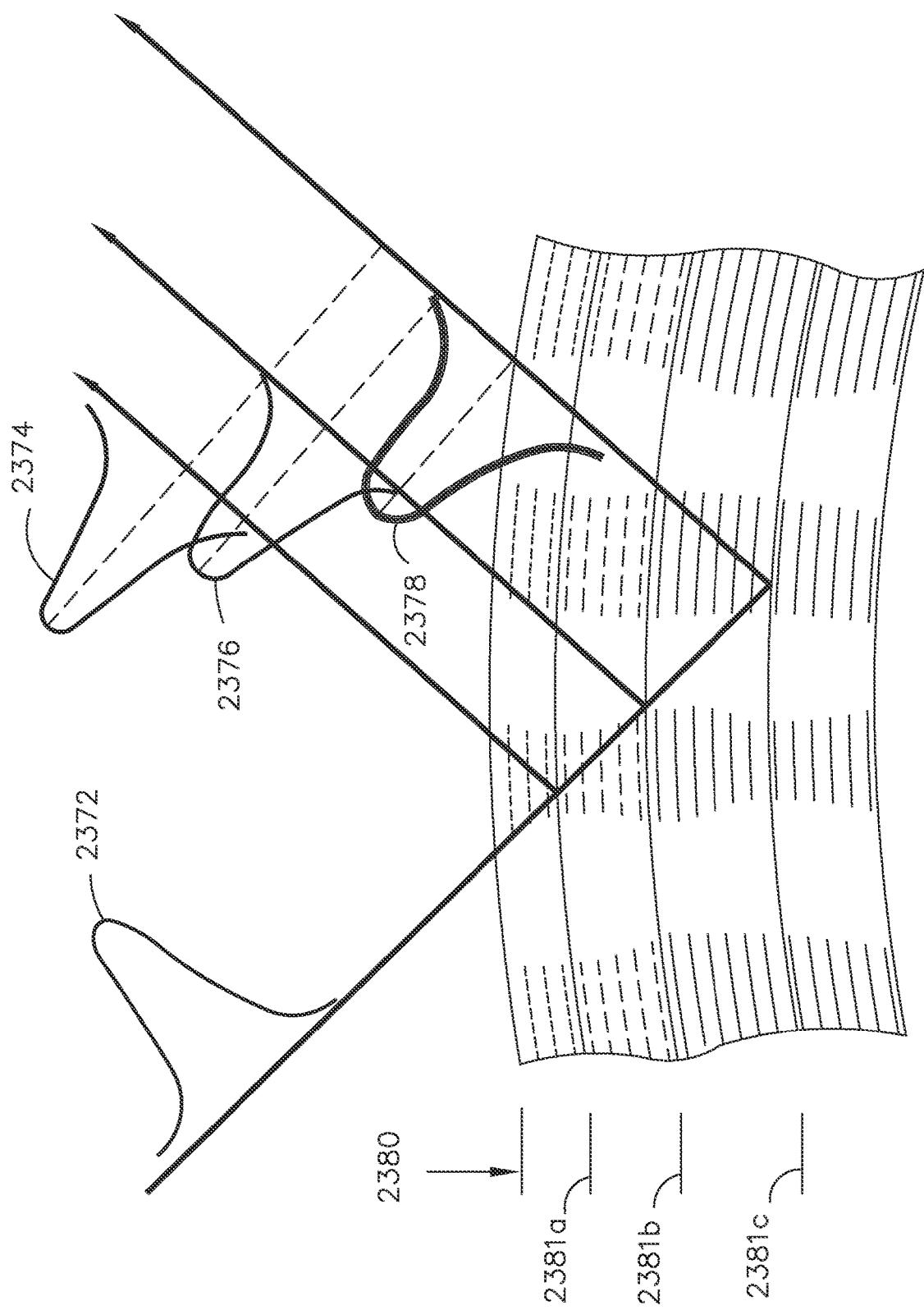


FIG. 167

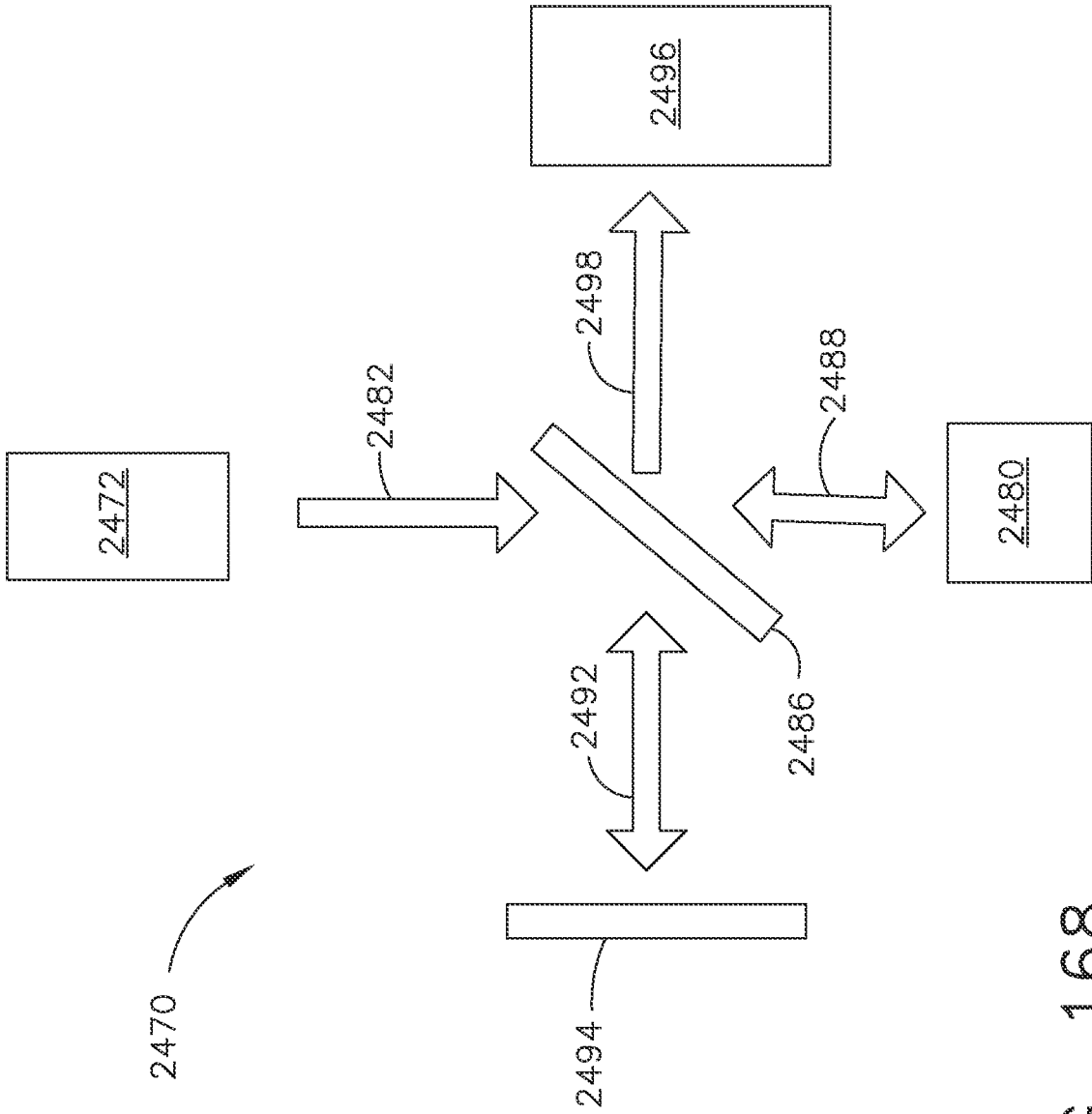


FIG. 168

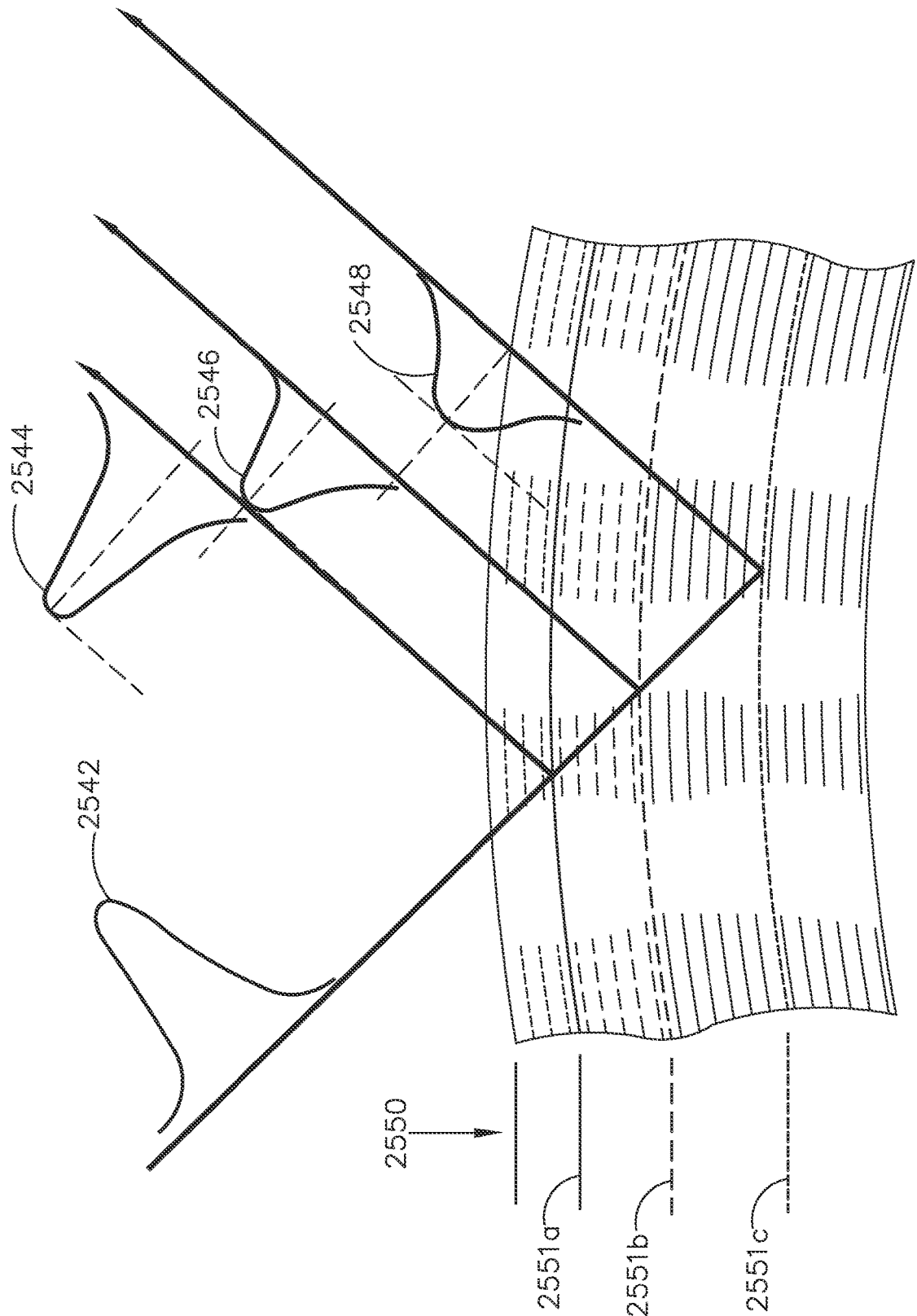


FIG. 169

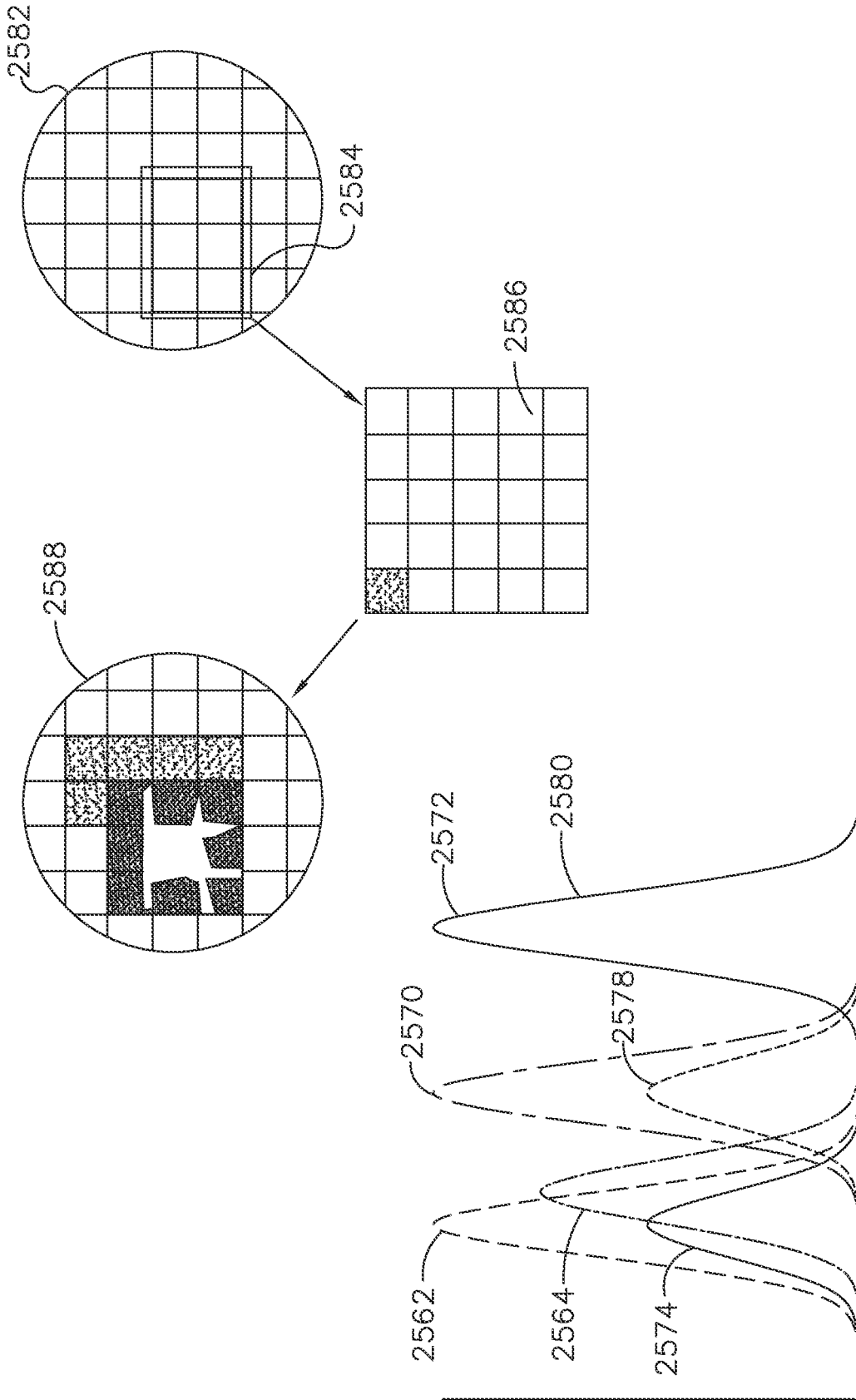


FIG. 170



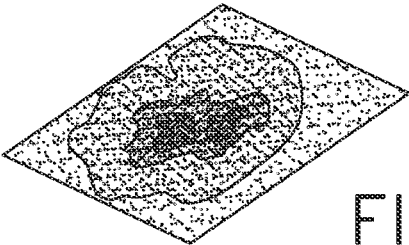


FIG. 171A

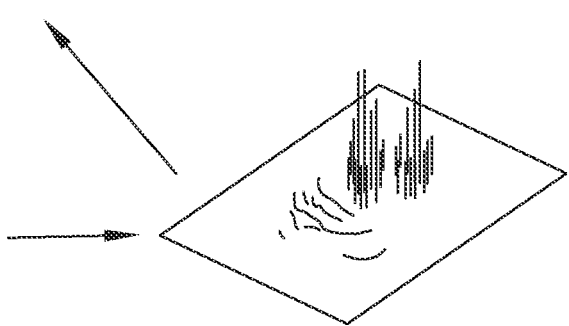


FIG. 171B

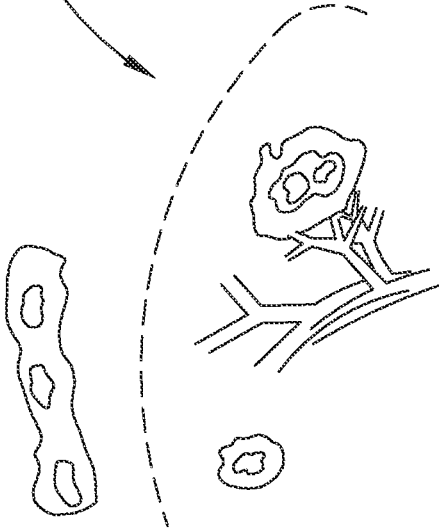


FIG. 171C

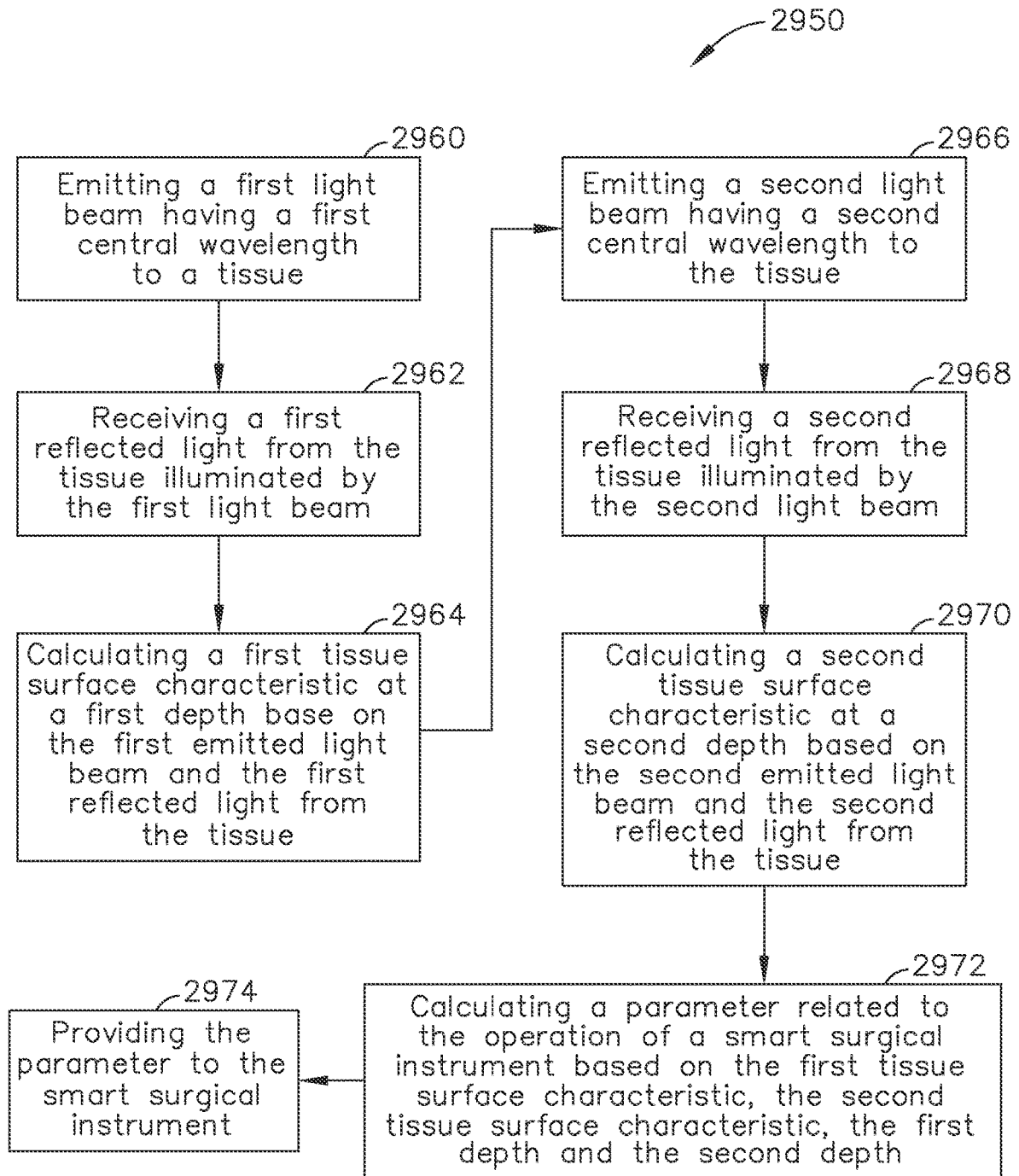


FIG. 172

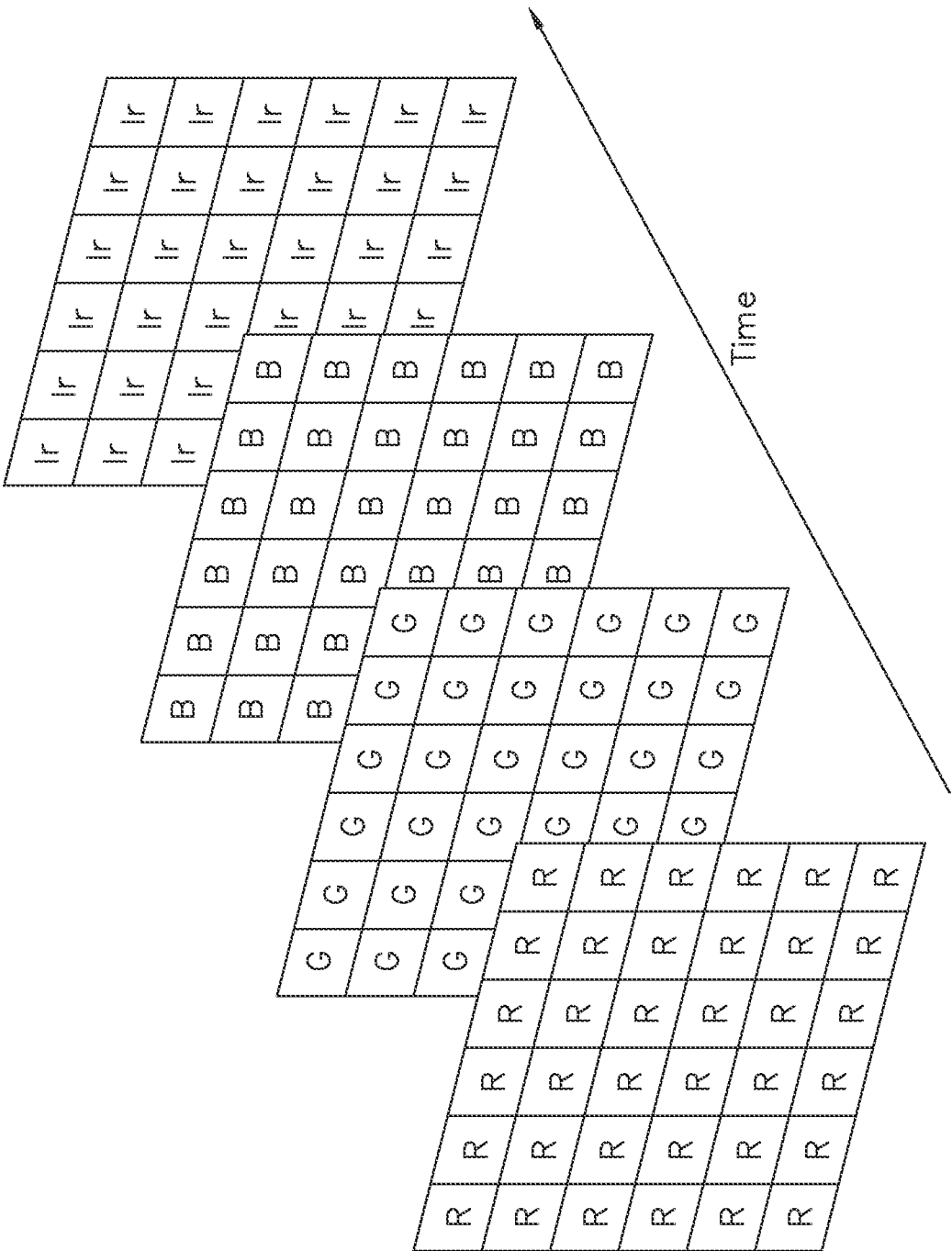


FIG. 173A

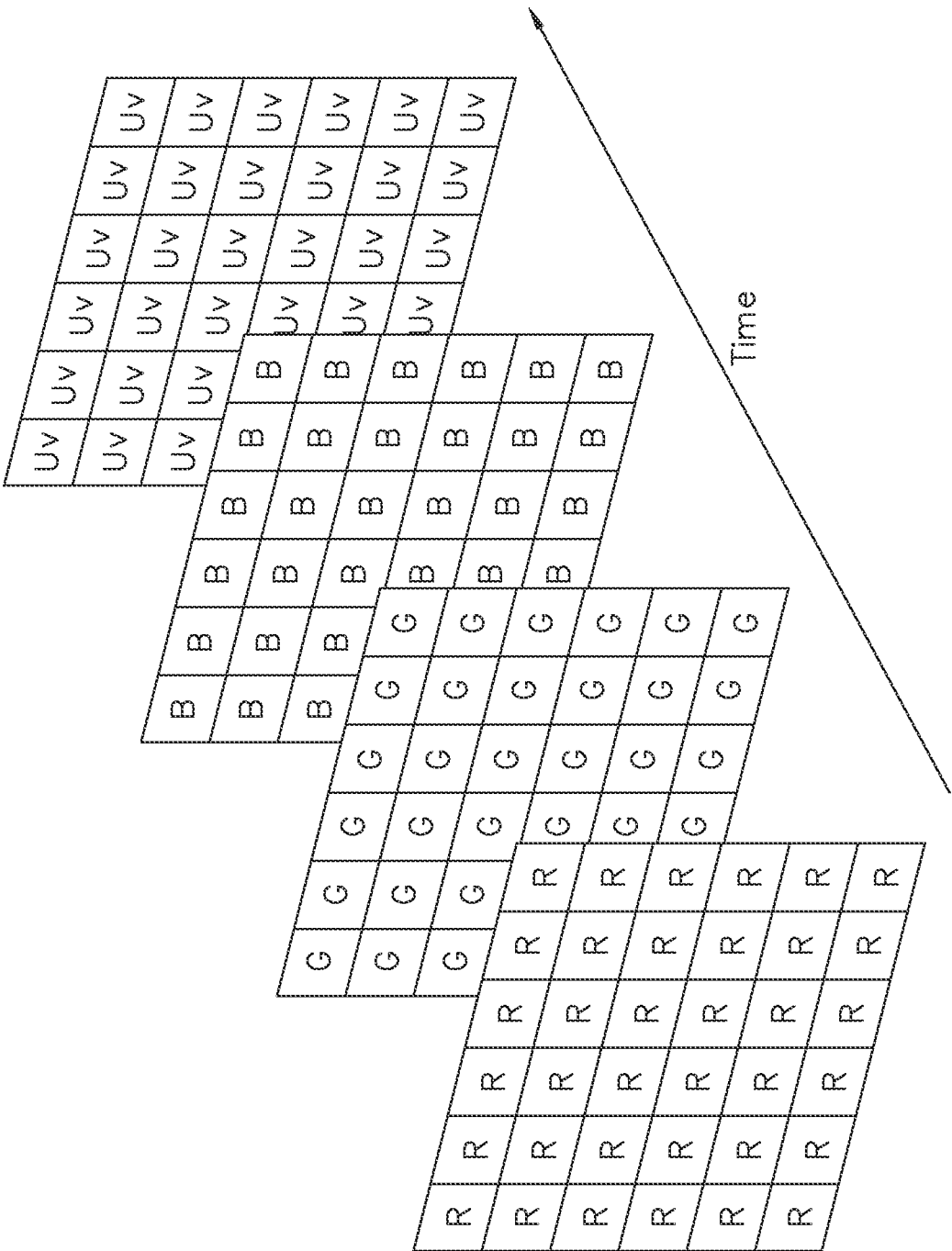


FIG. 173B

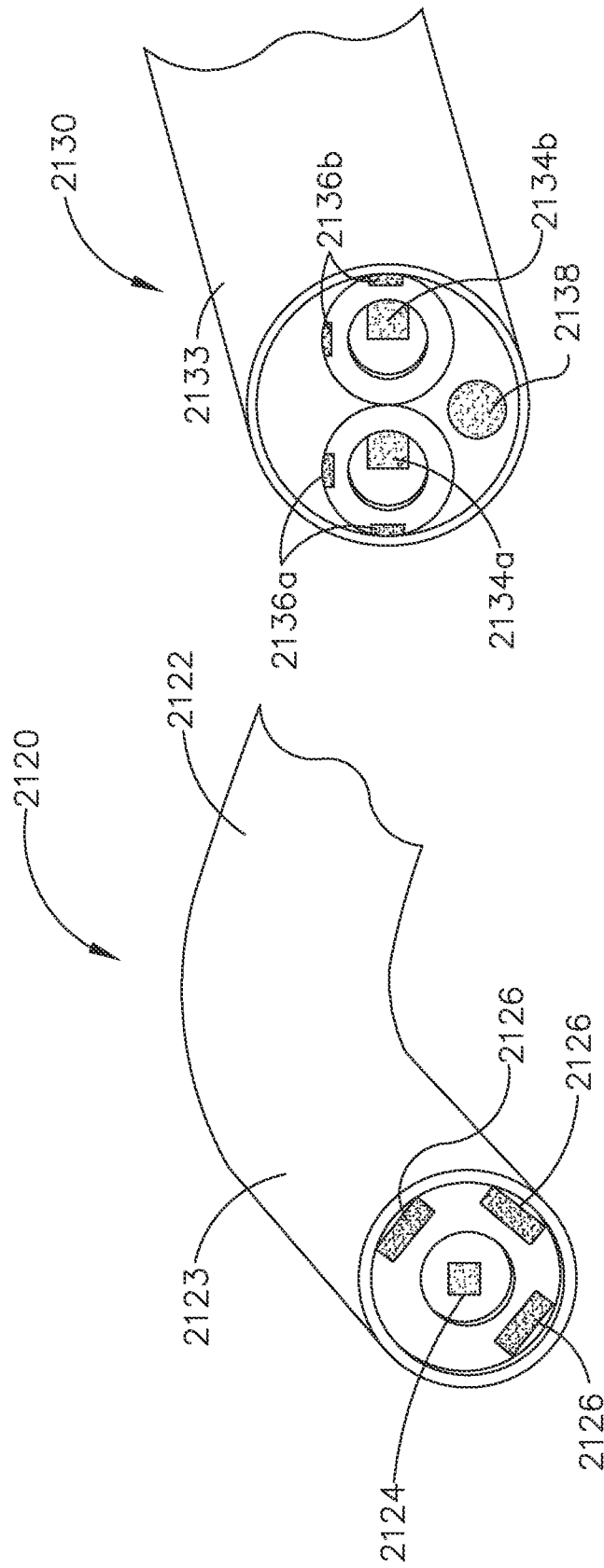


FIG. 174B

FIG. 174A

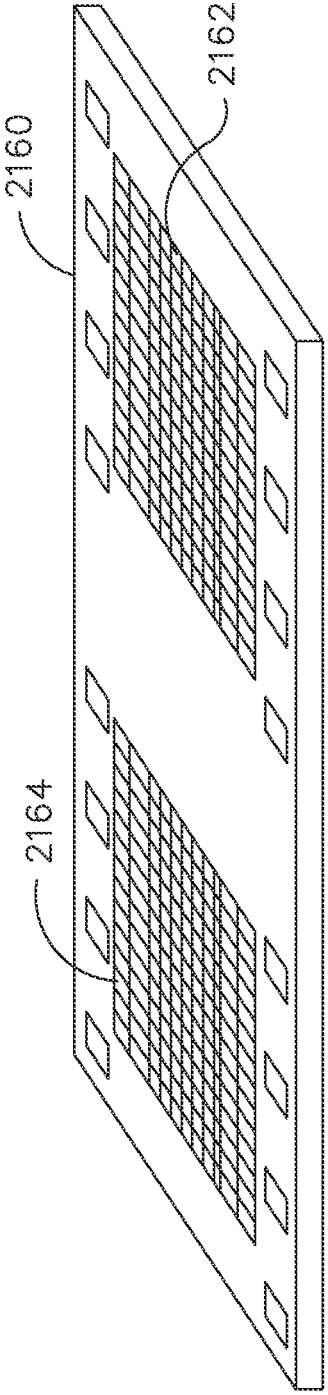


FIG. 174C

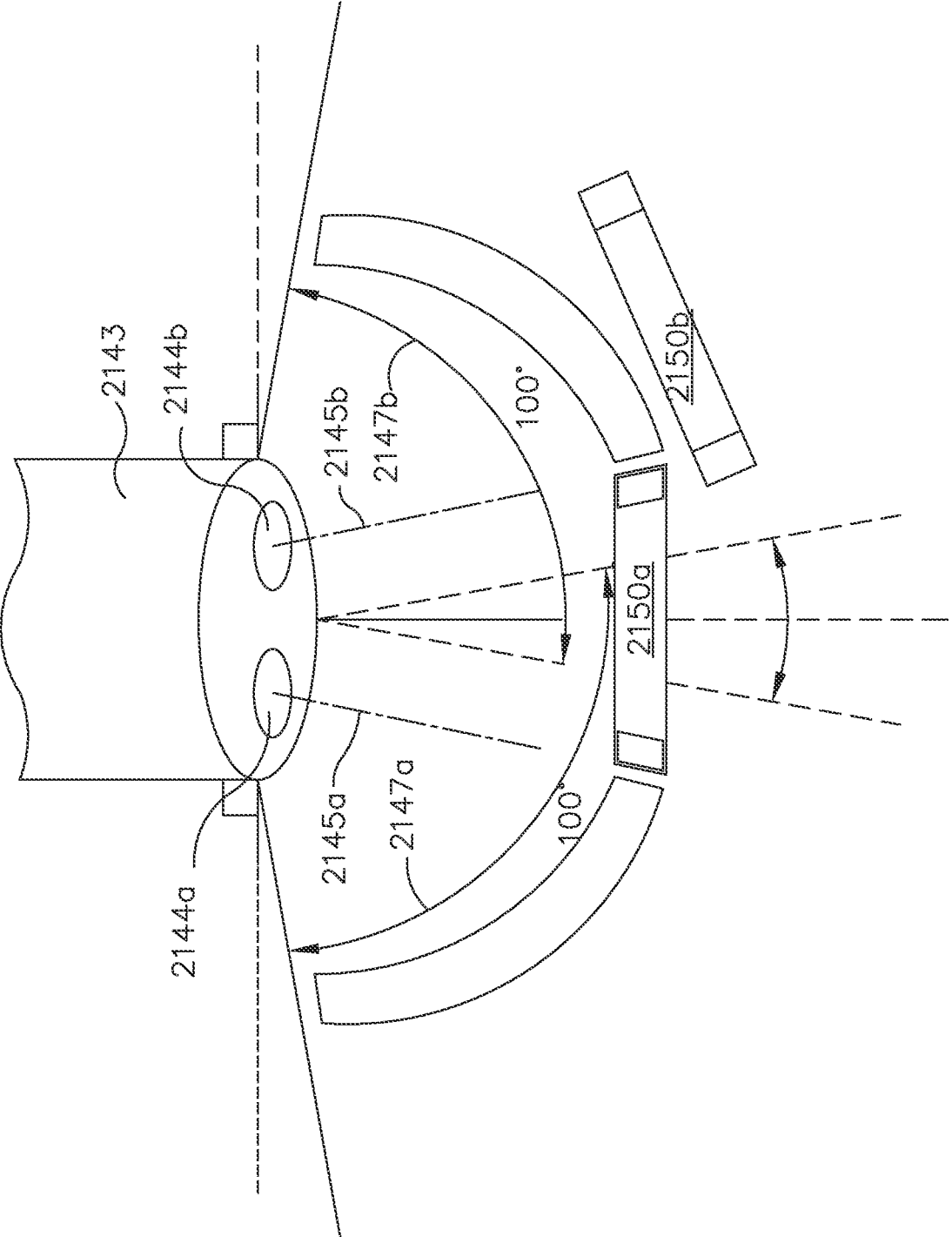


FIG. 175

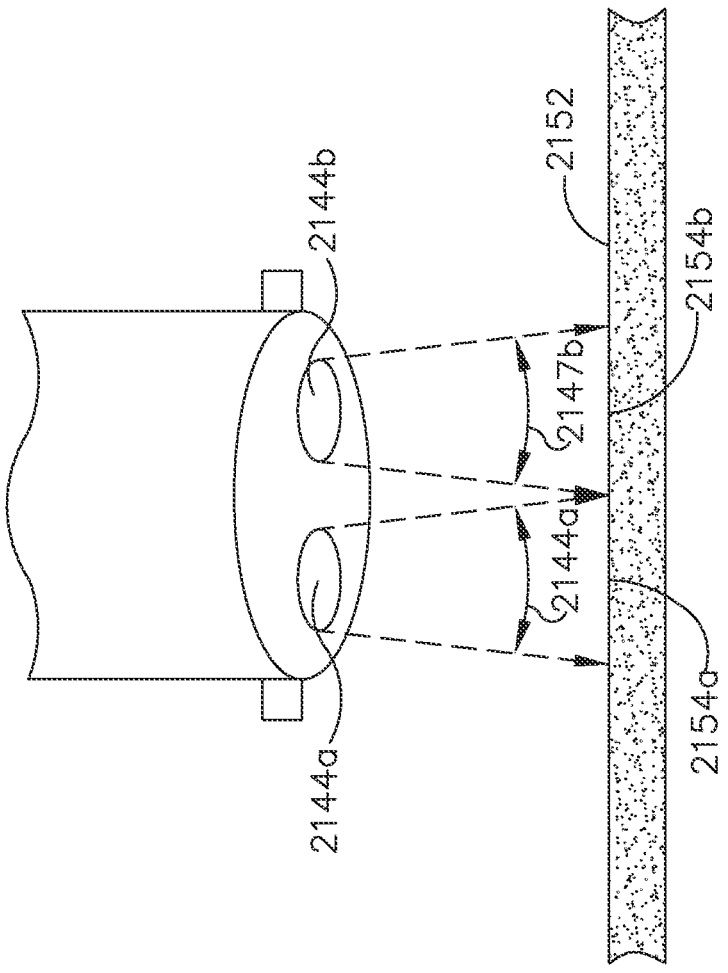


FIG. 176A



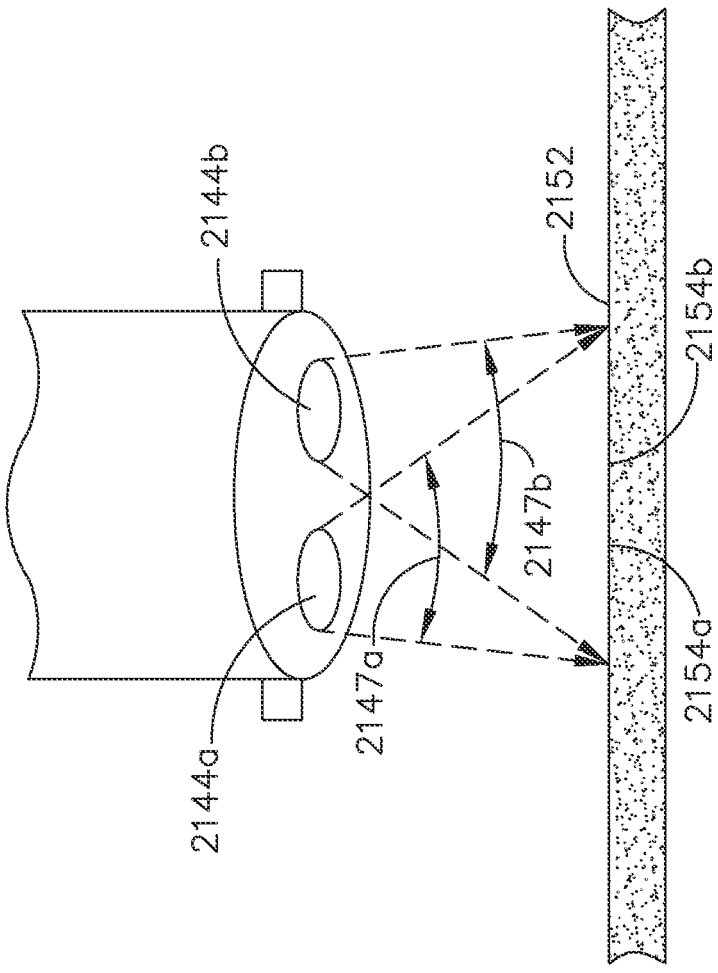


FIG. 176B

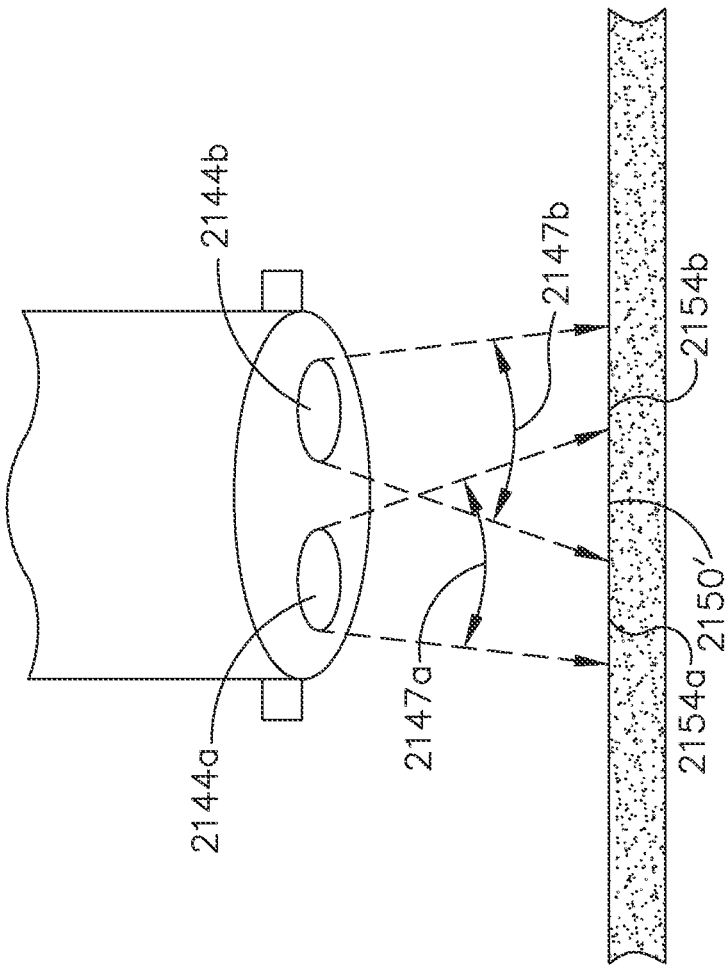


FIG. 176C

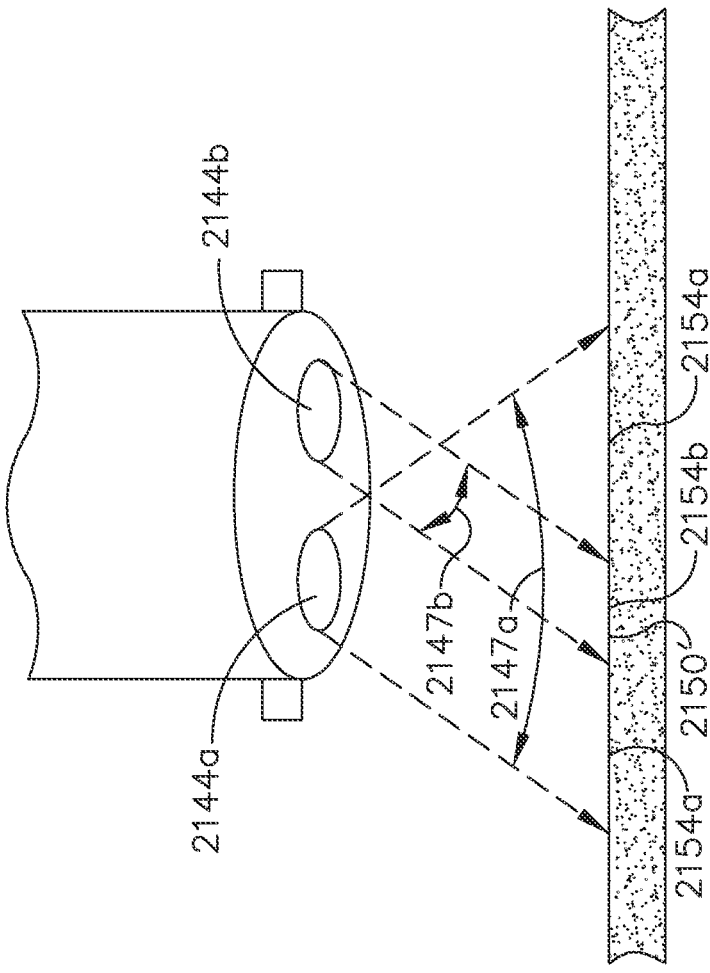


FIG. 176D

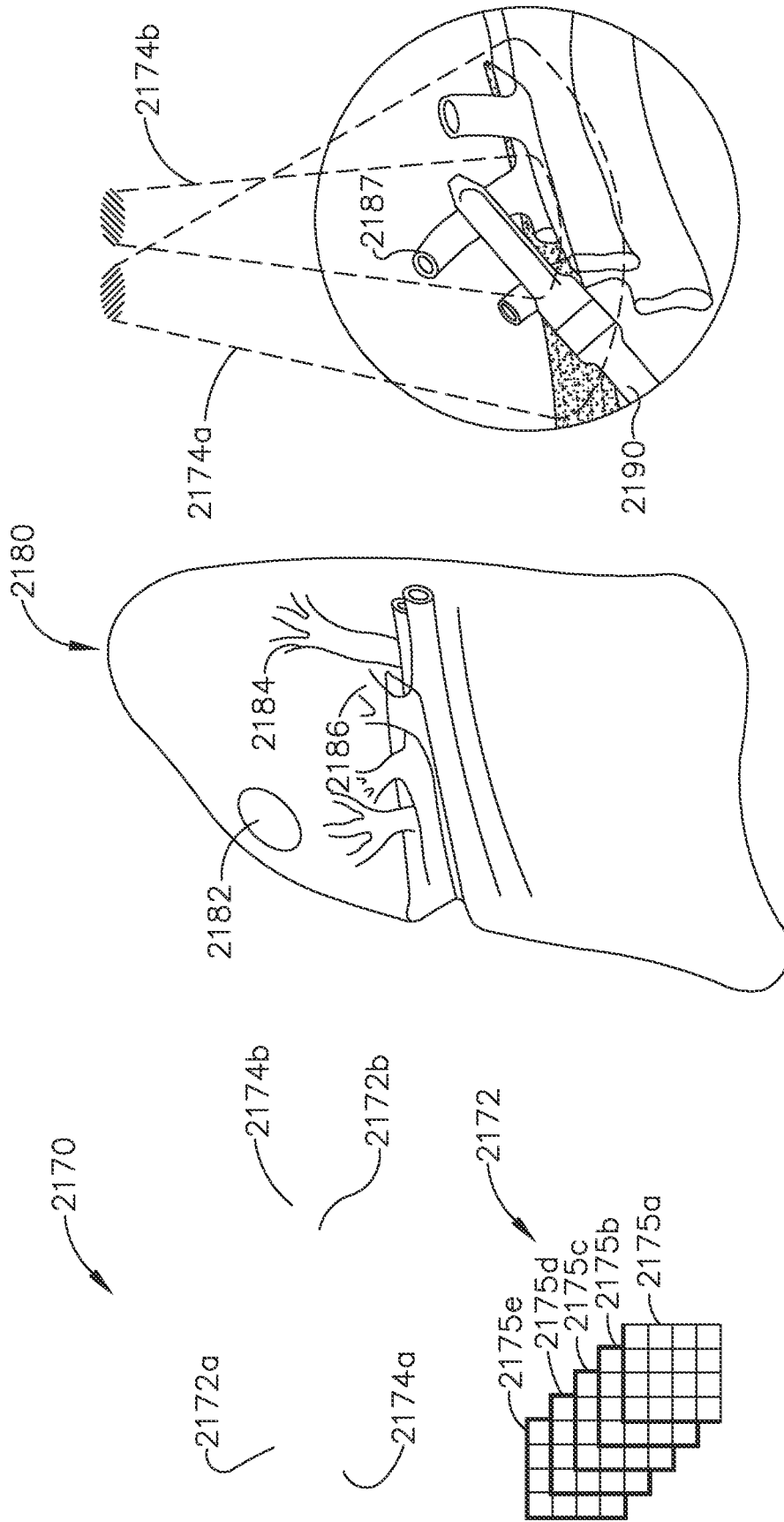


FIG. 177A FIG. 177B FIG. 177C

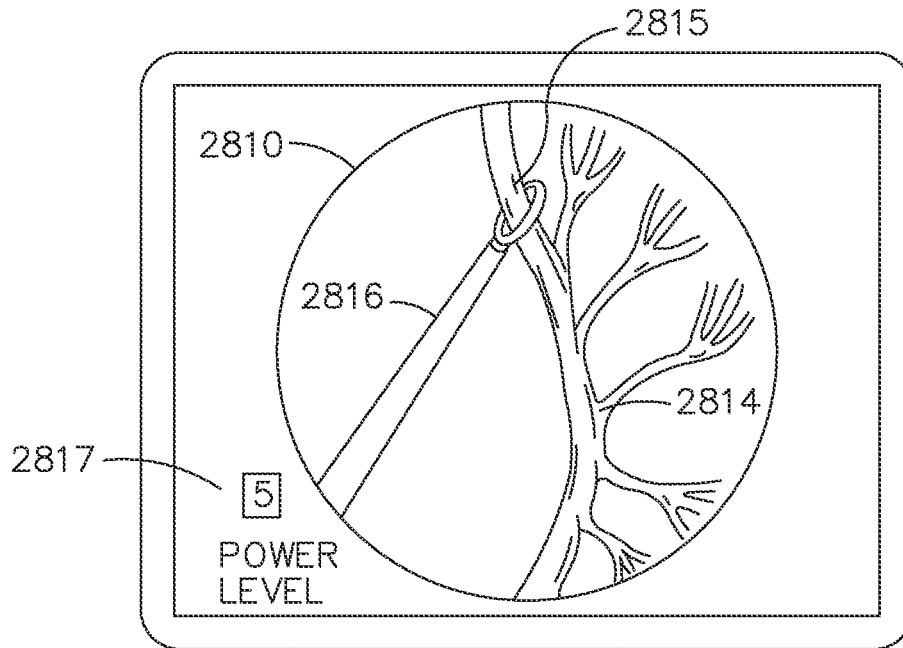


FIG. 178B

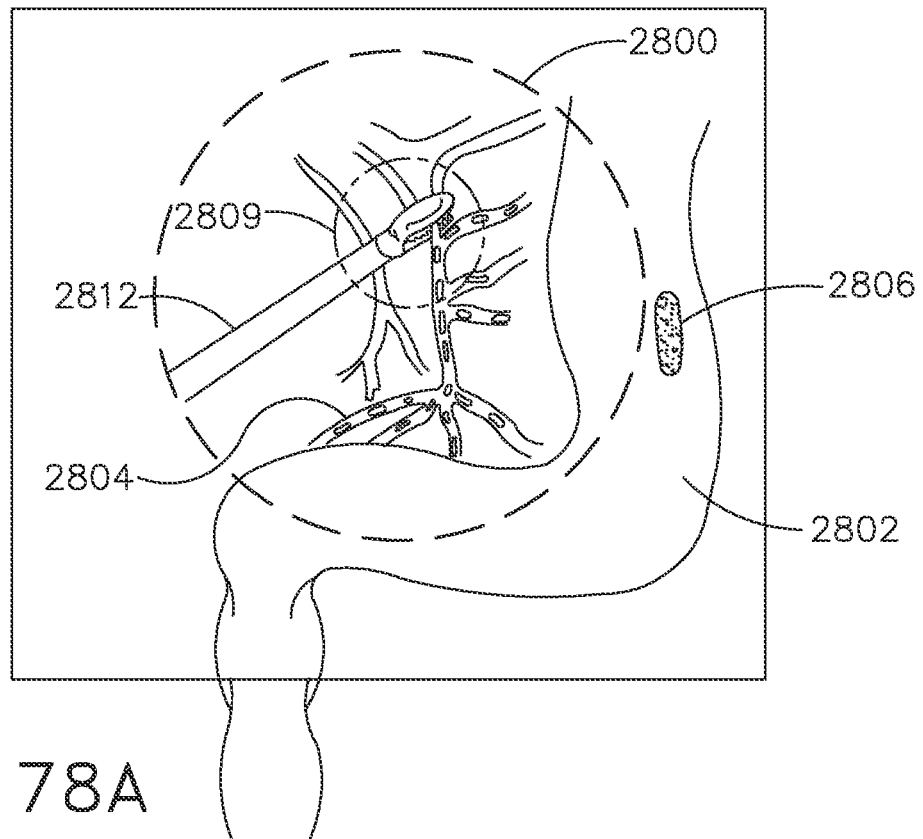


FIG. 178A

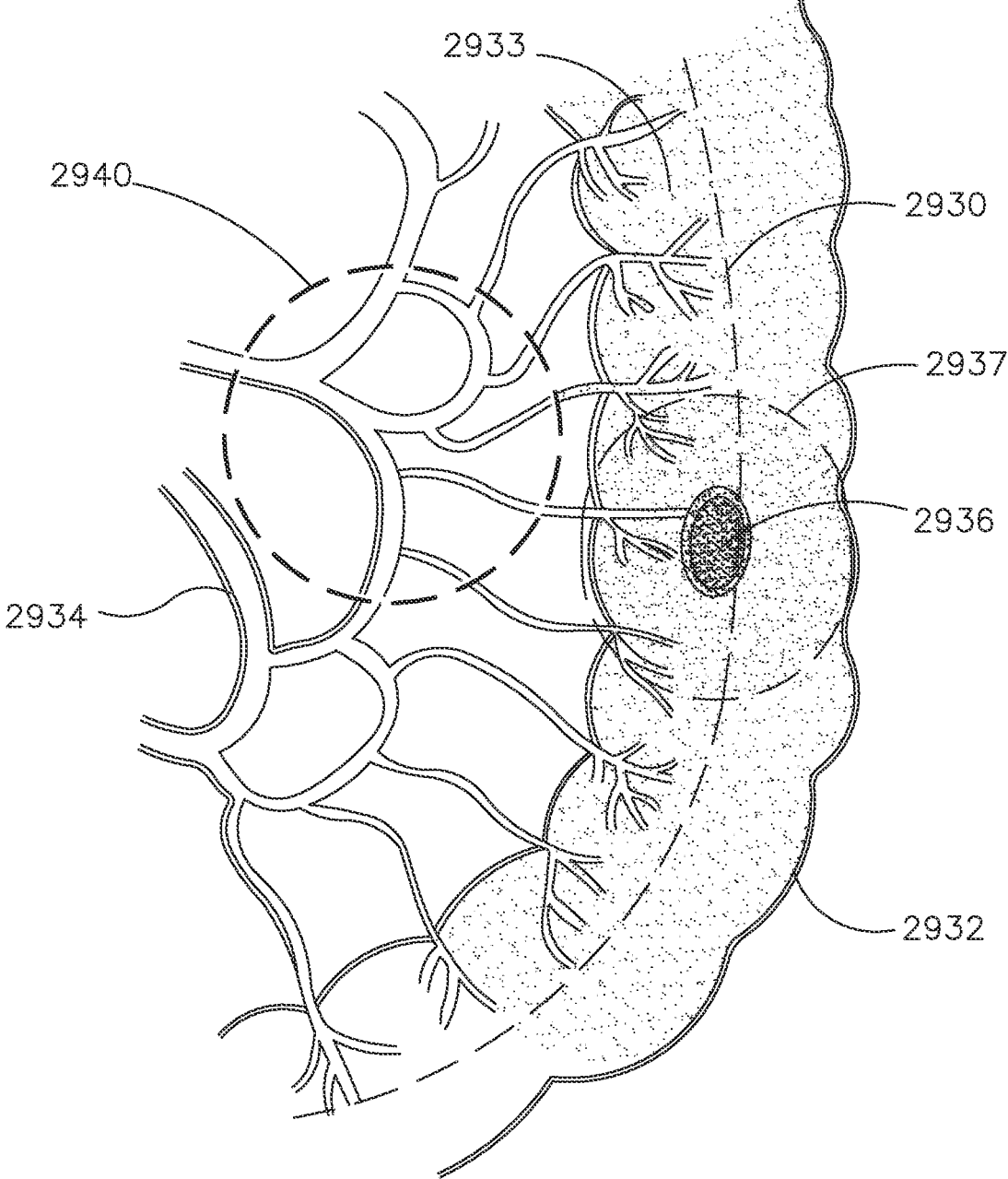


FIG. 179A

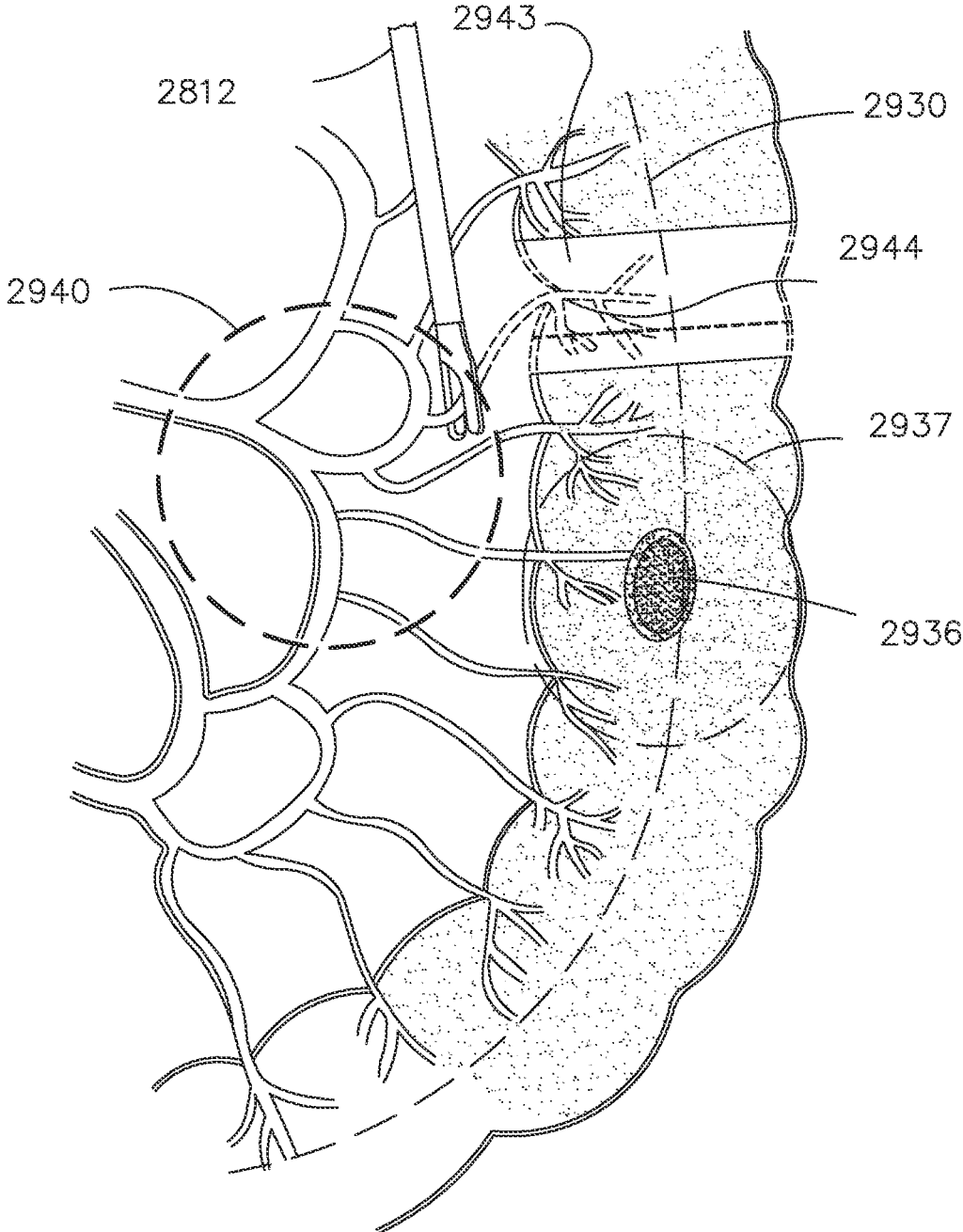


FIG. 179B

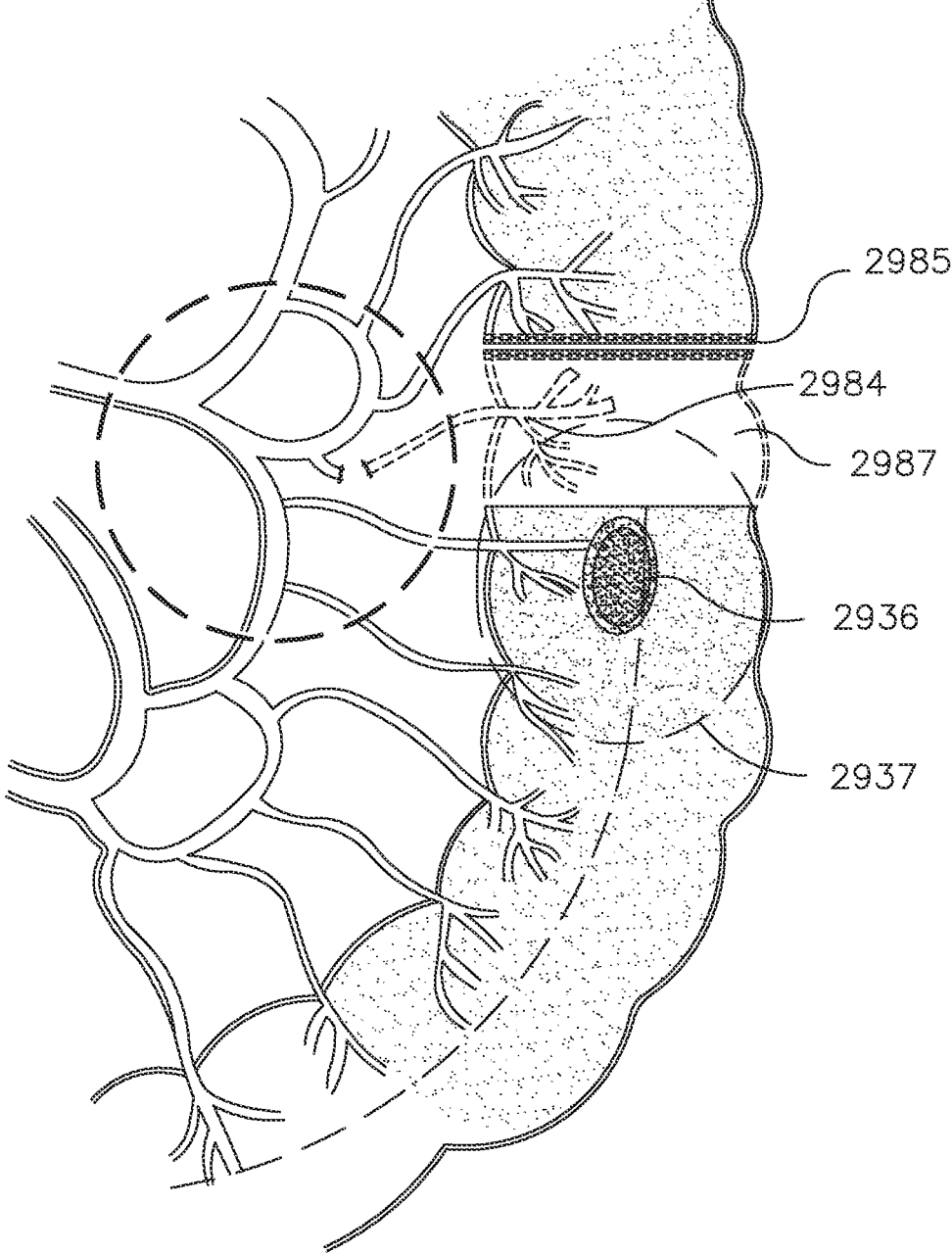


FIG. 179C



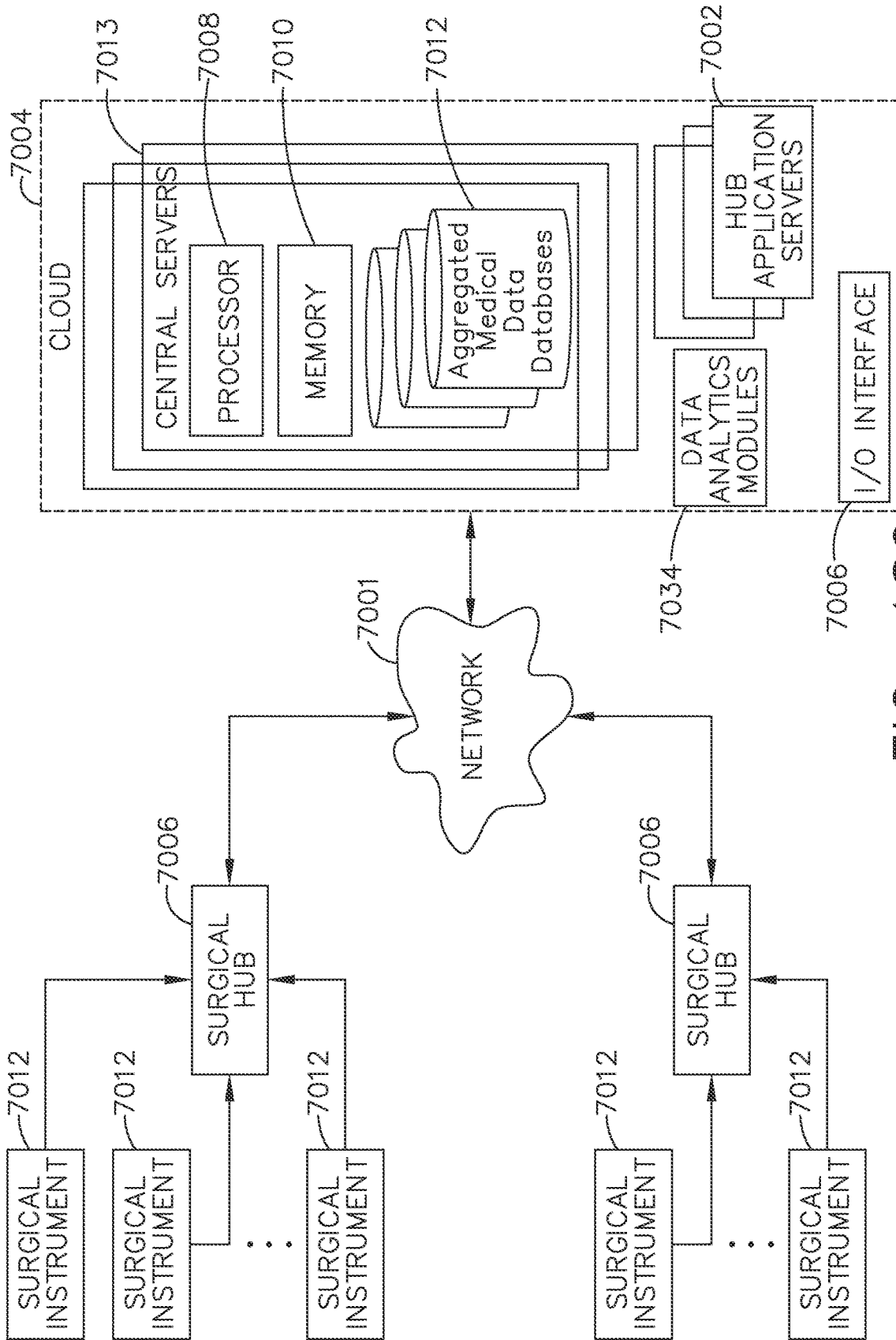


FIG. 180

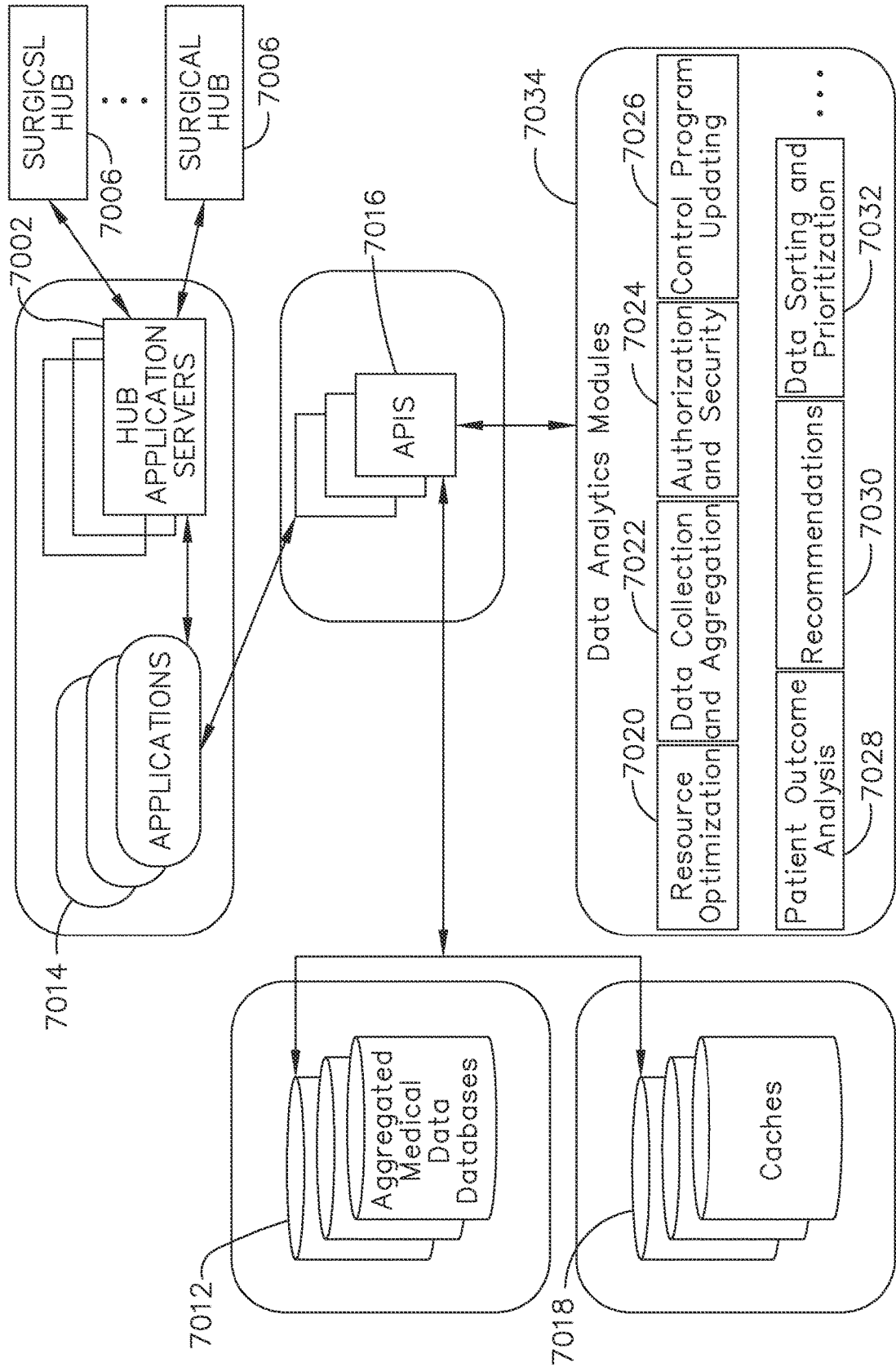


FIG. 181

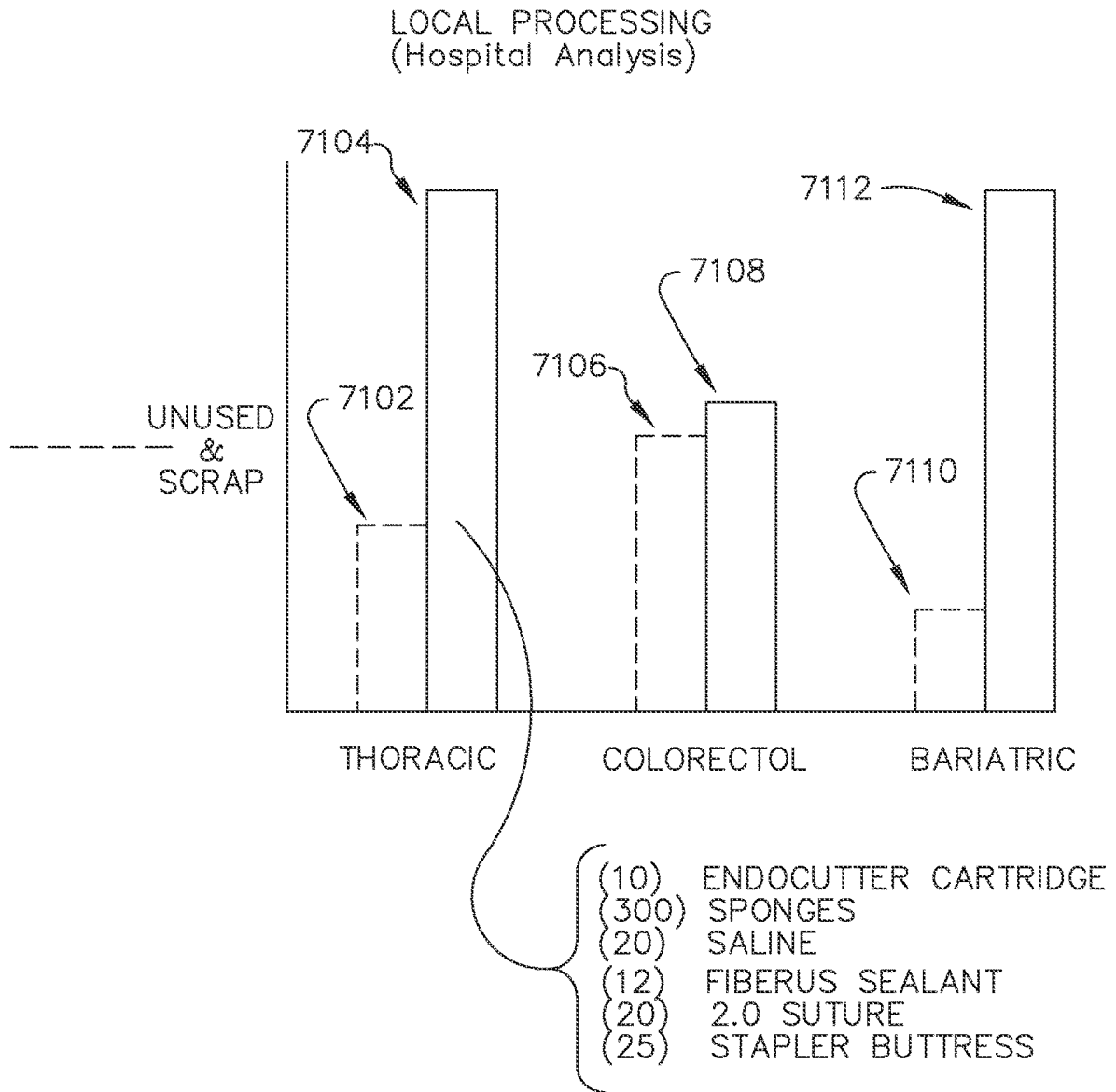


FIG. 182

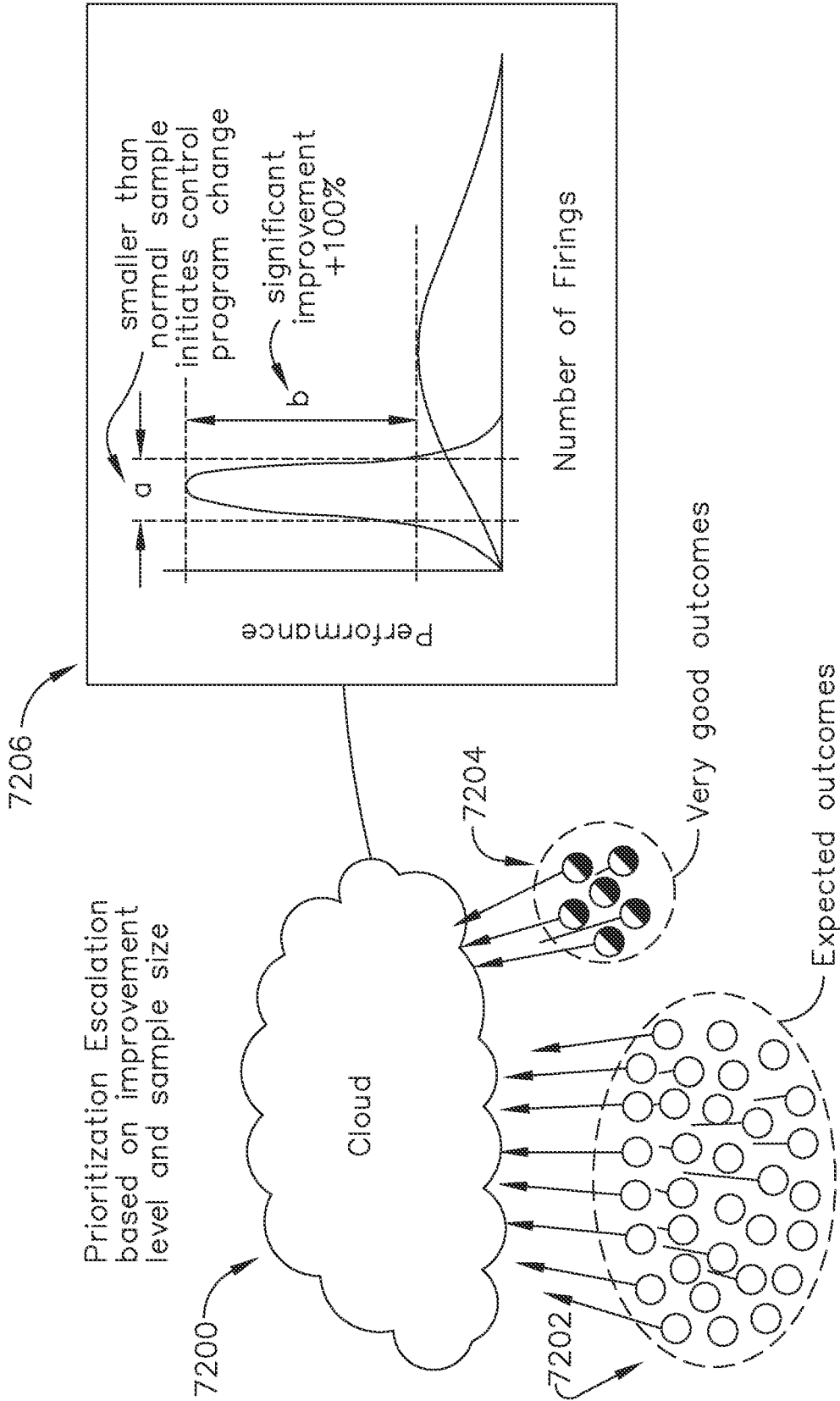


FIG. 183

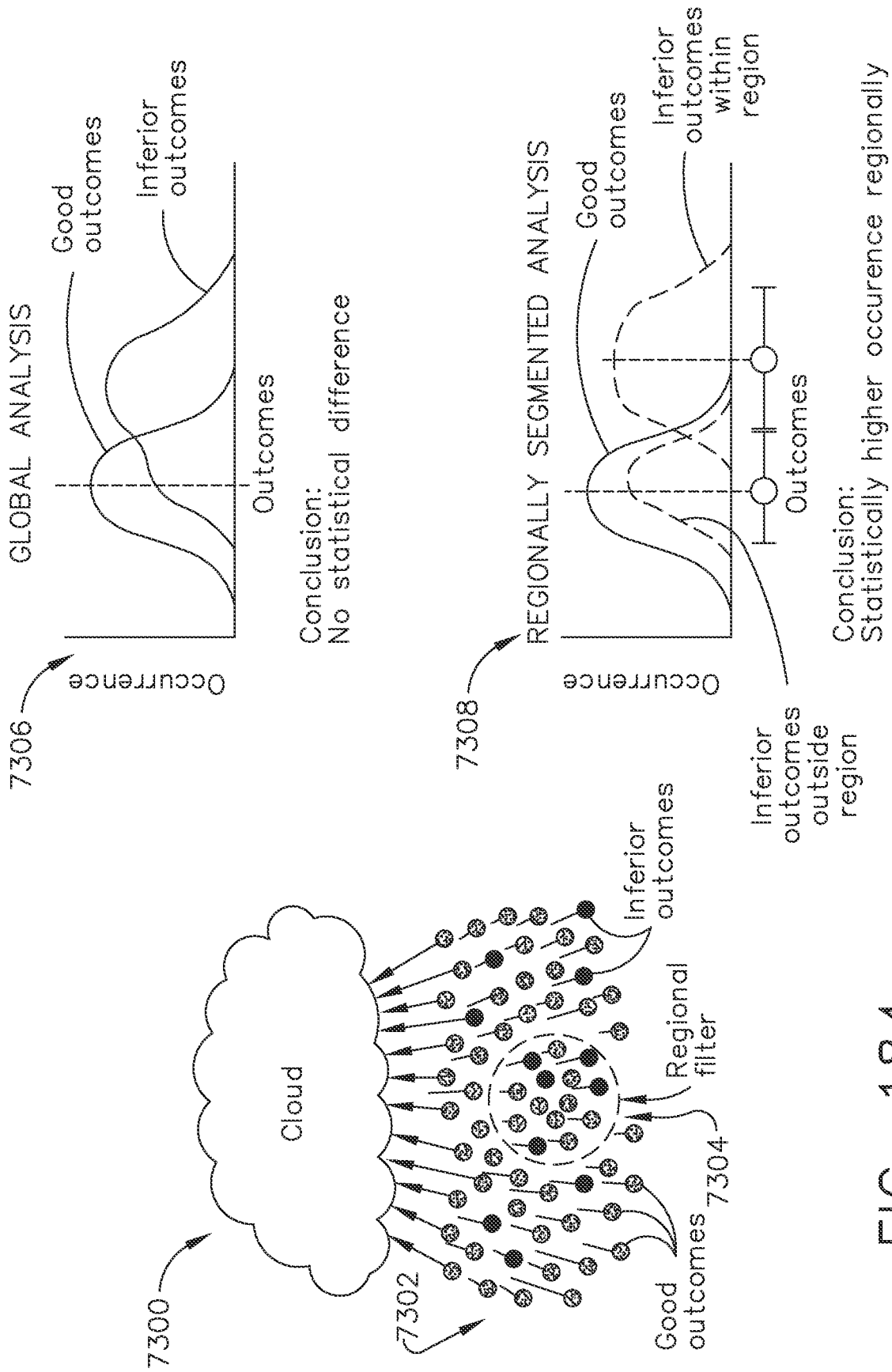


FIG. 184

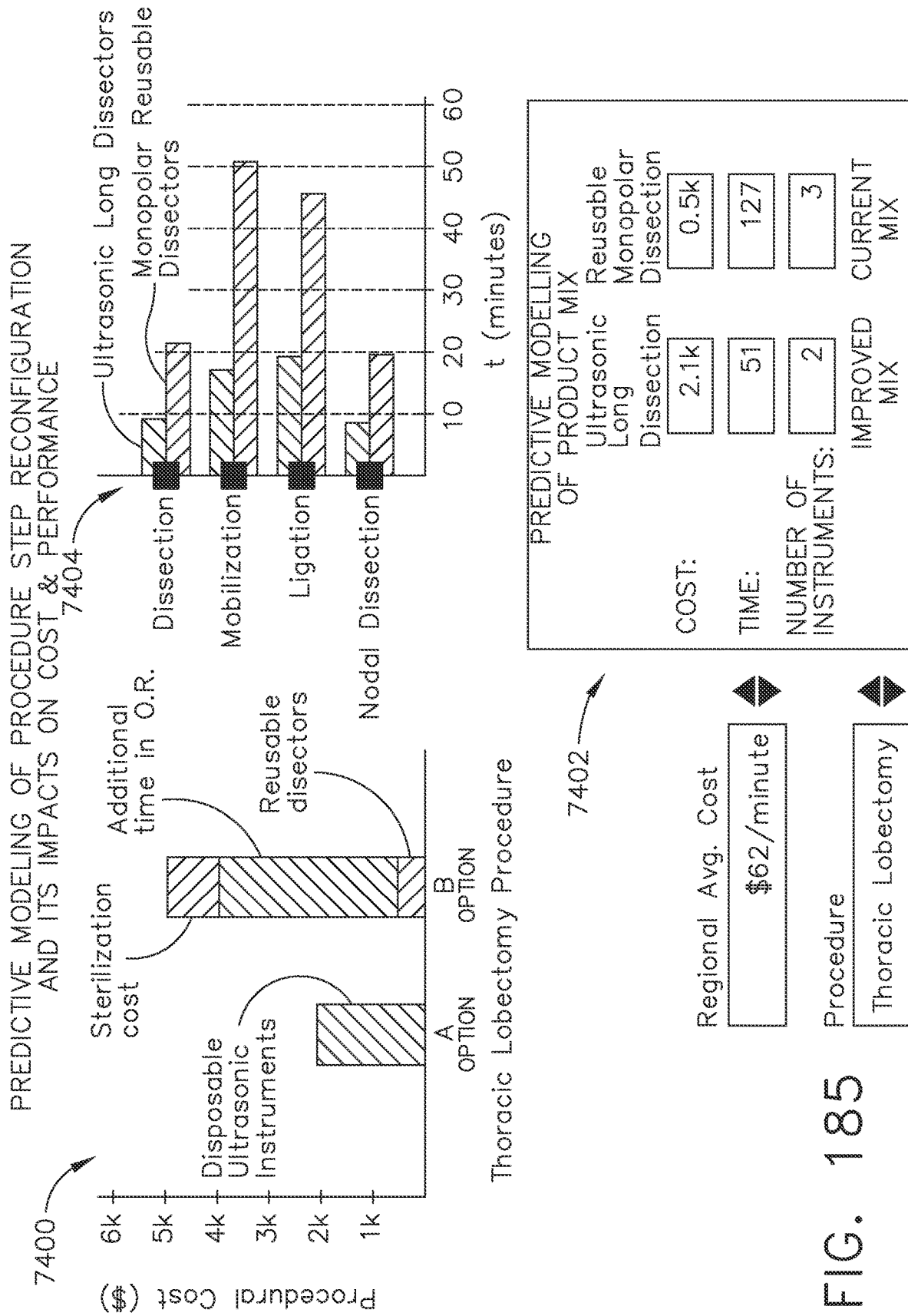


FIG. 185

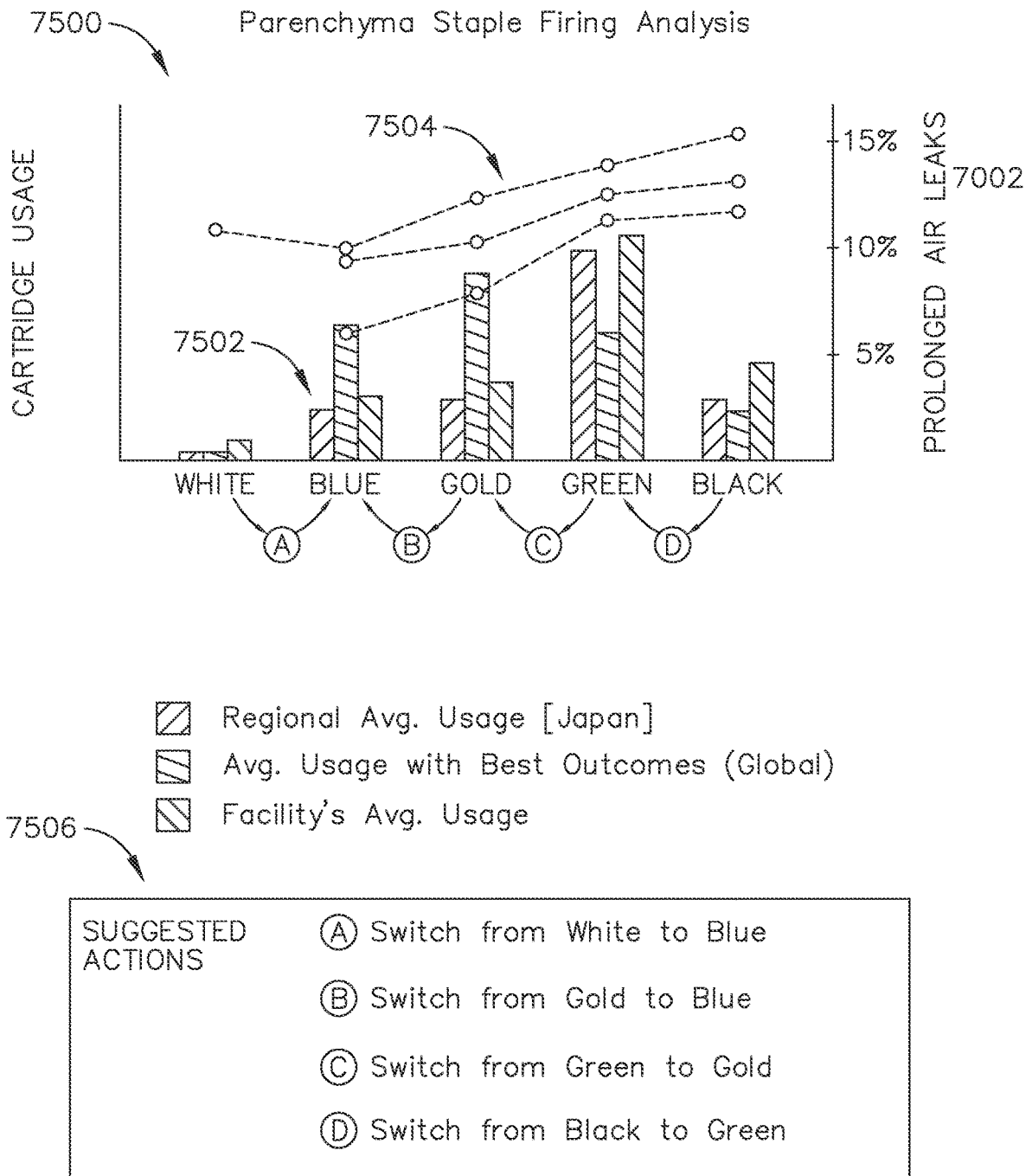


FIG. 186

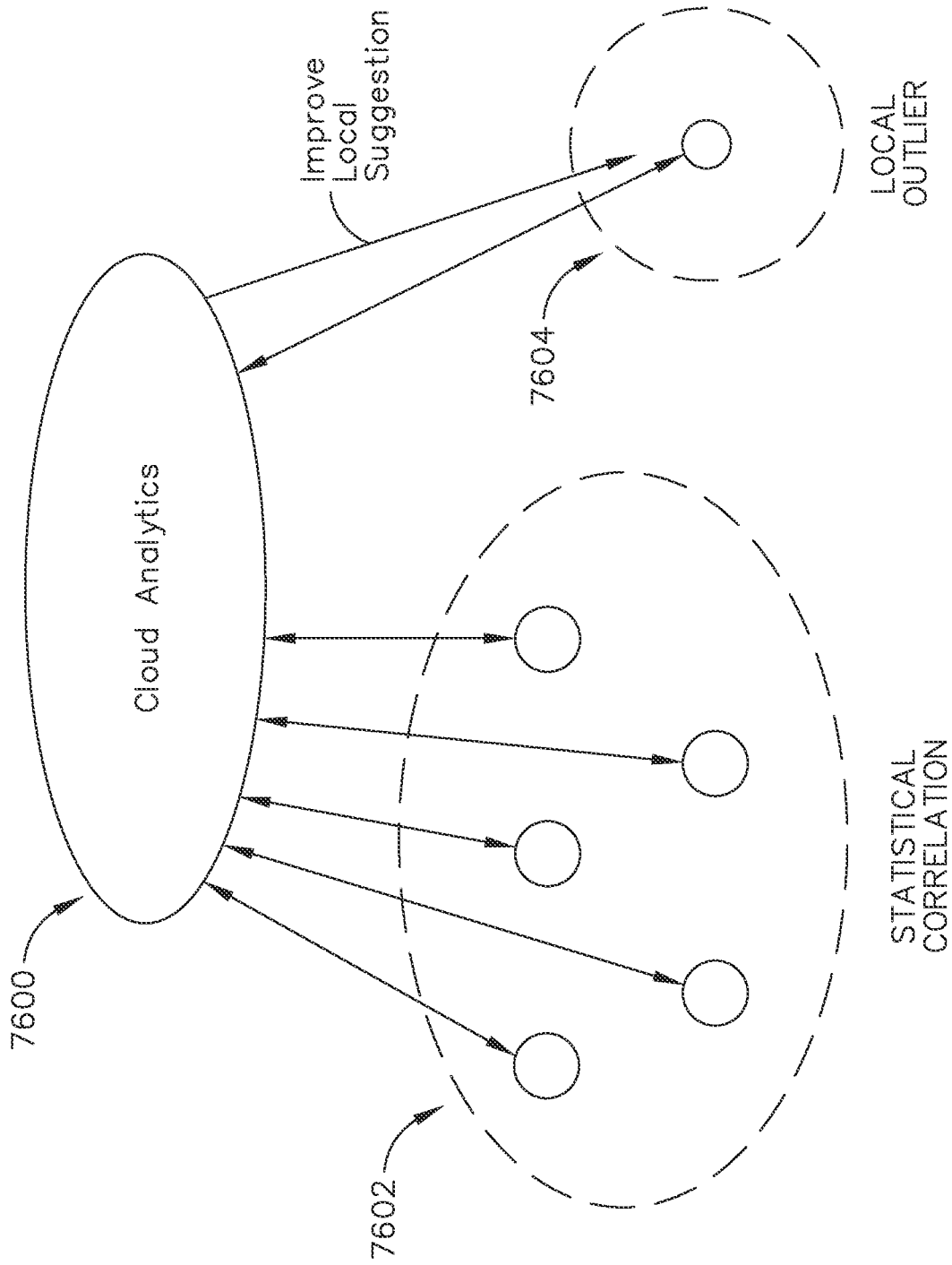
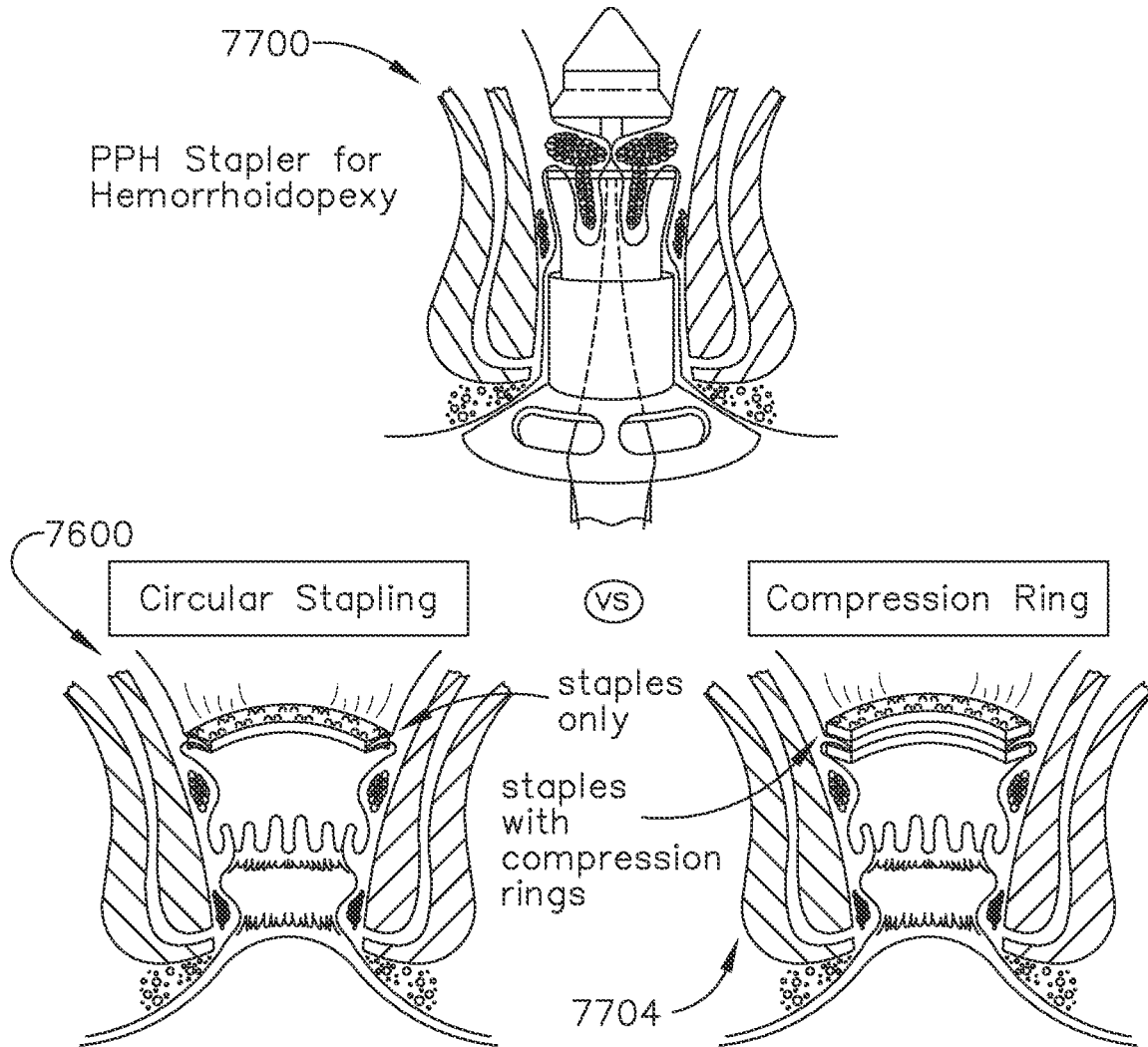


FIG. 187



Different Product Mix to Competitor  
with EQUIVALENT Outcomes



7706	Instrument Cost	\$130	<	\$142	Instrument Cost
	Time	38 min	<	45 min	Time
	Efficacy		<		Efficacy
	Current Mix	=		Suggested Mfg. equivalent of current mix	

FIG. 188

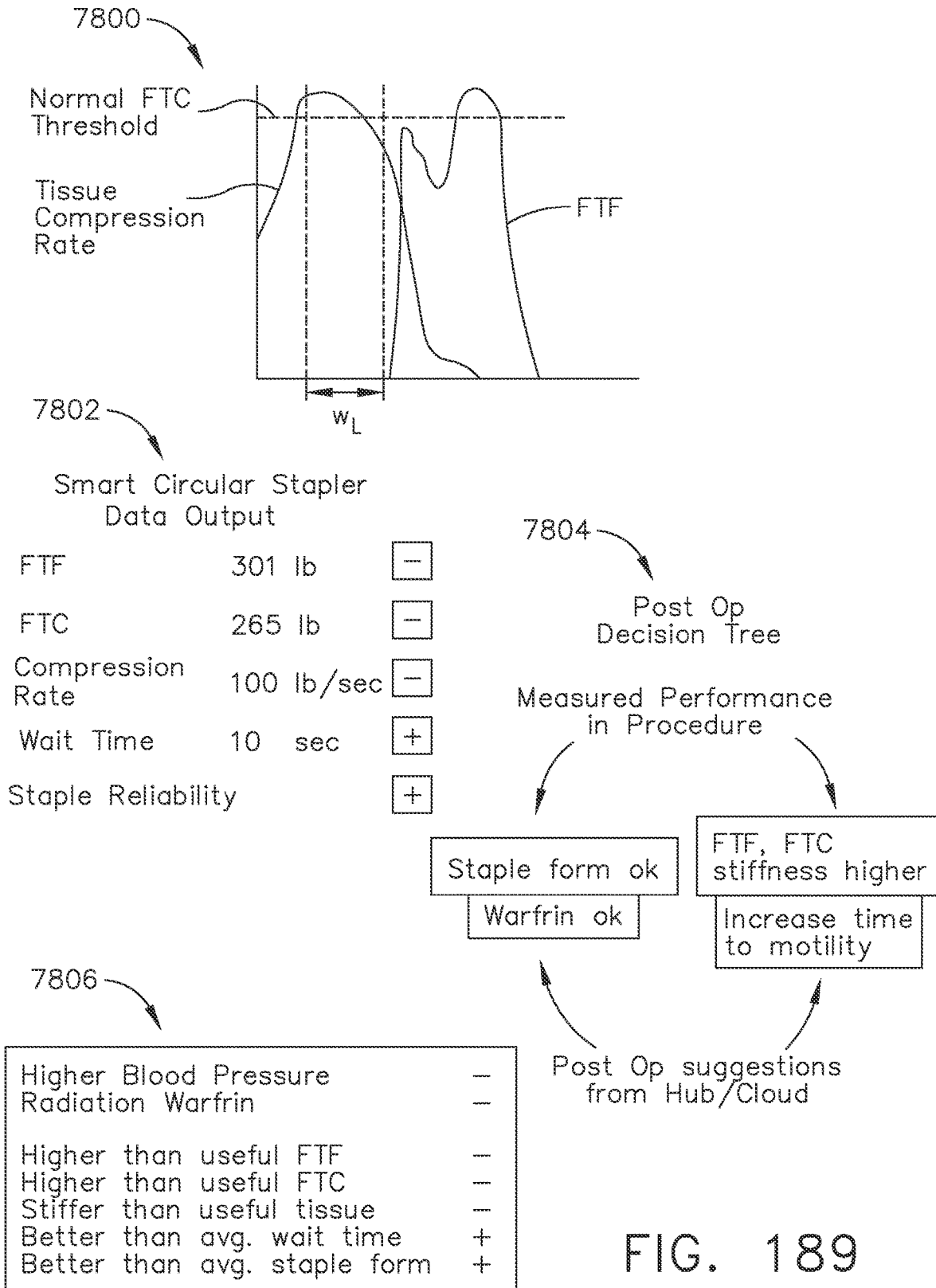


FIG. 189

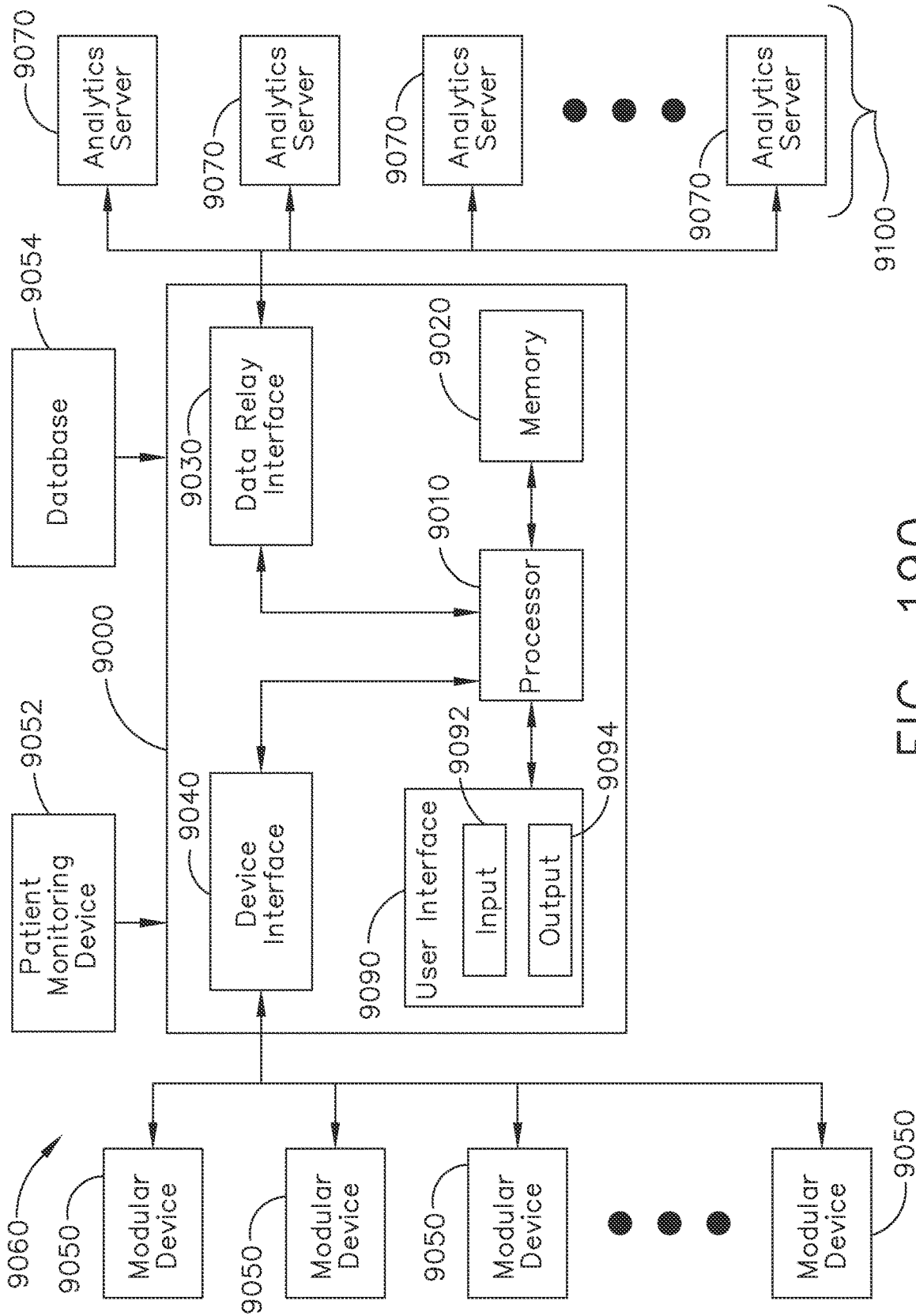


FIG. 190

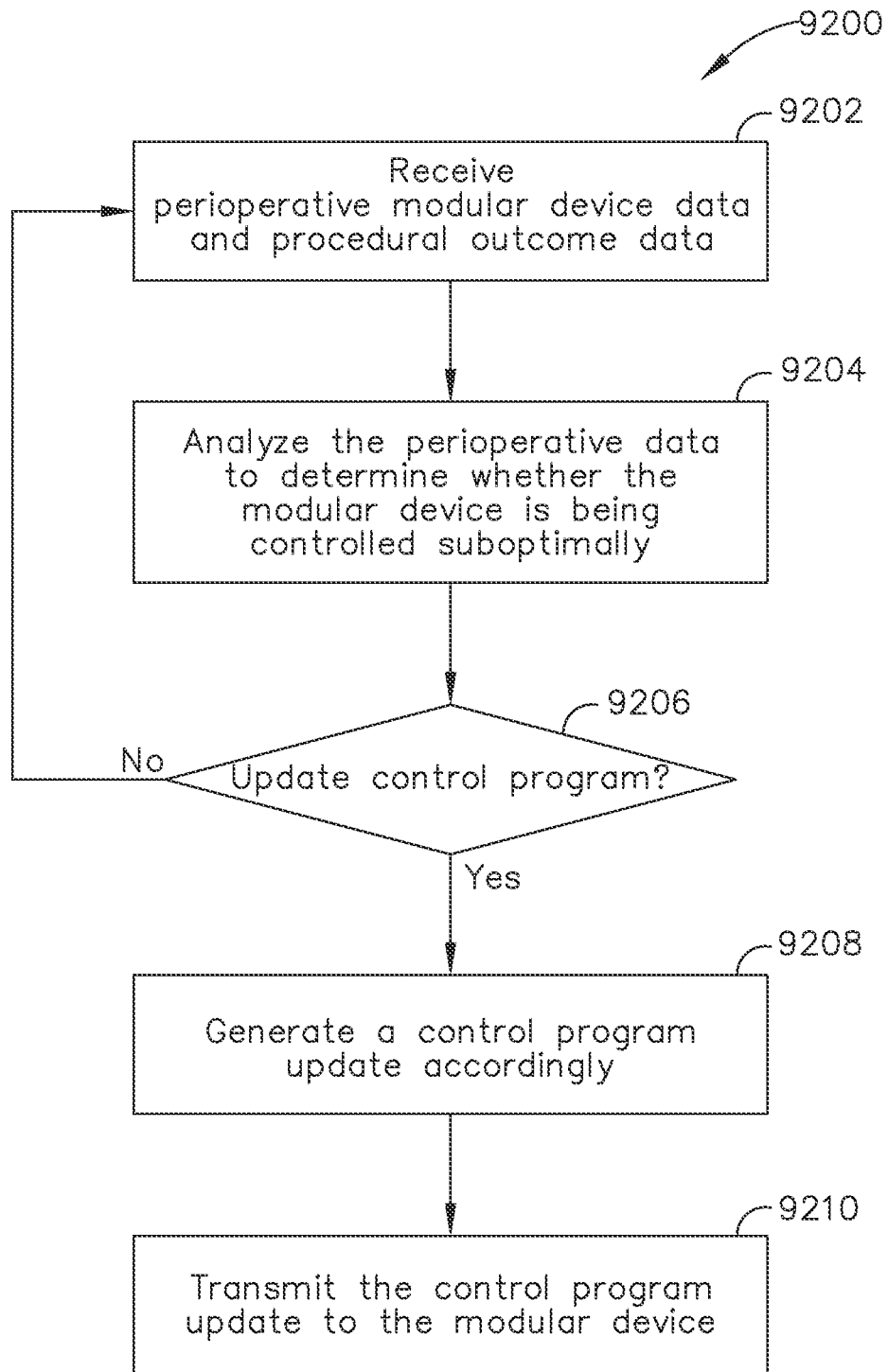


FIG. 191

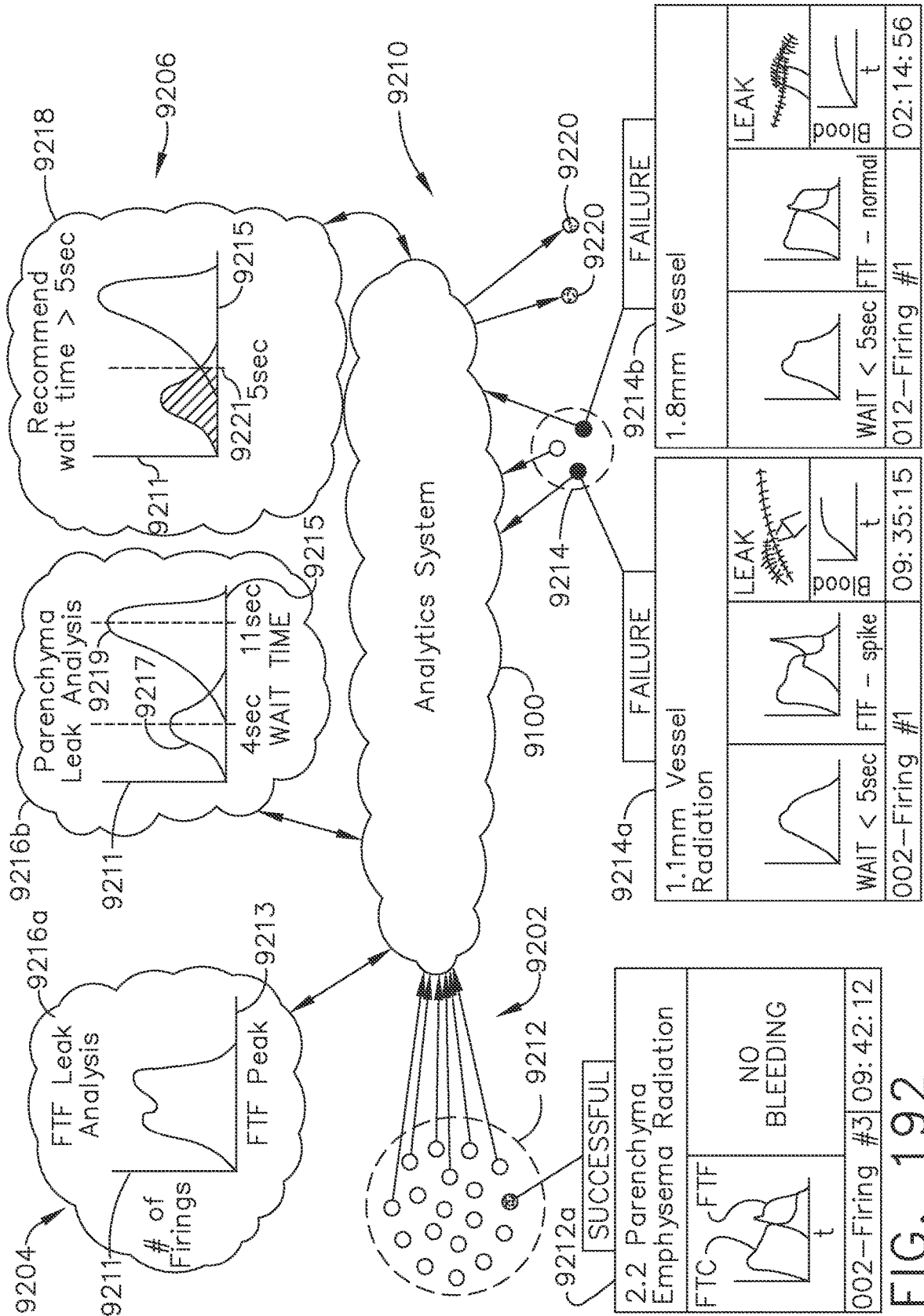


FIG. 192

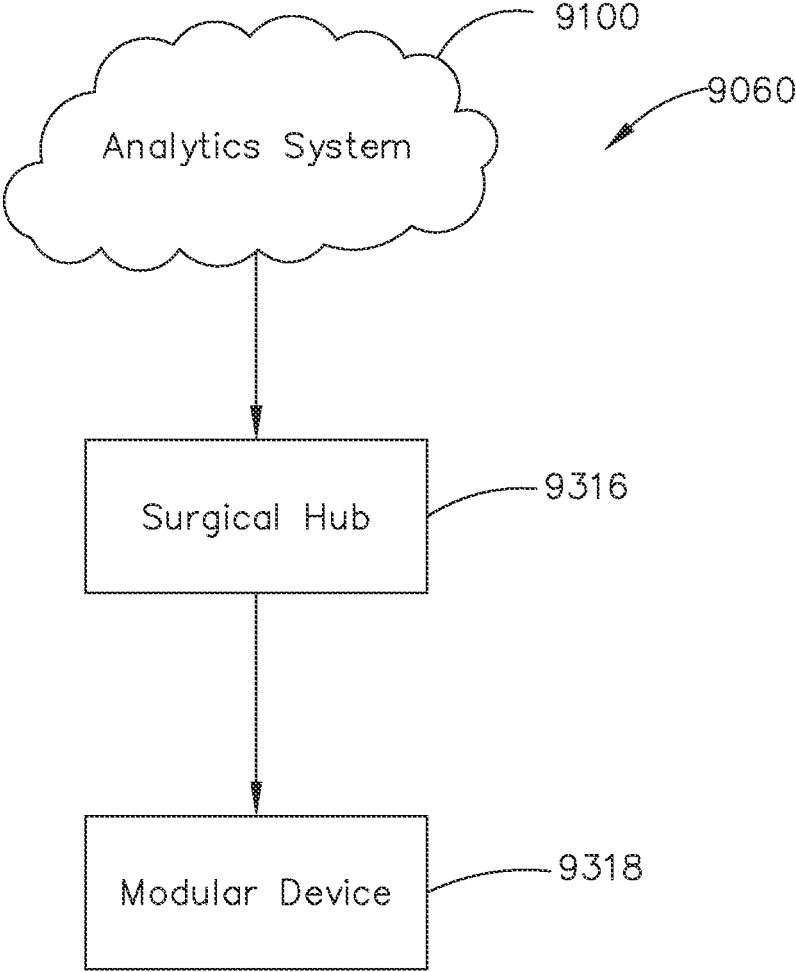


FIG. 193

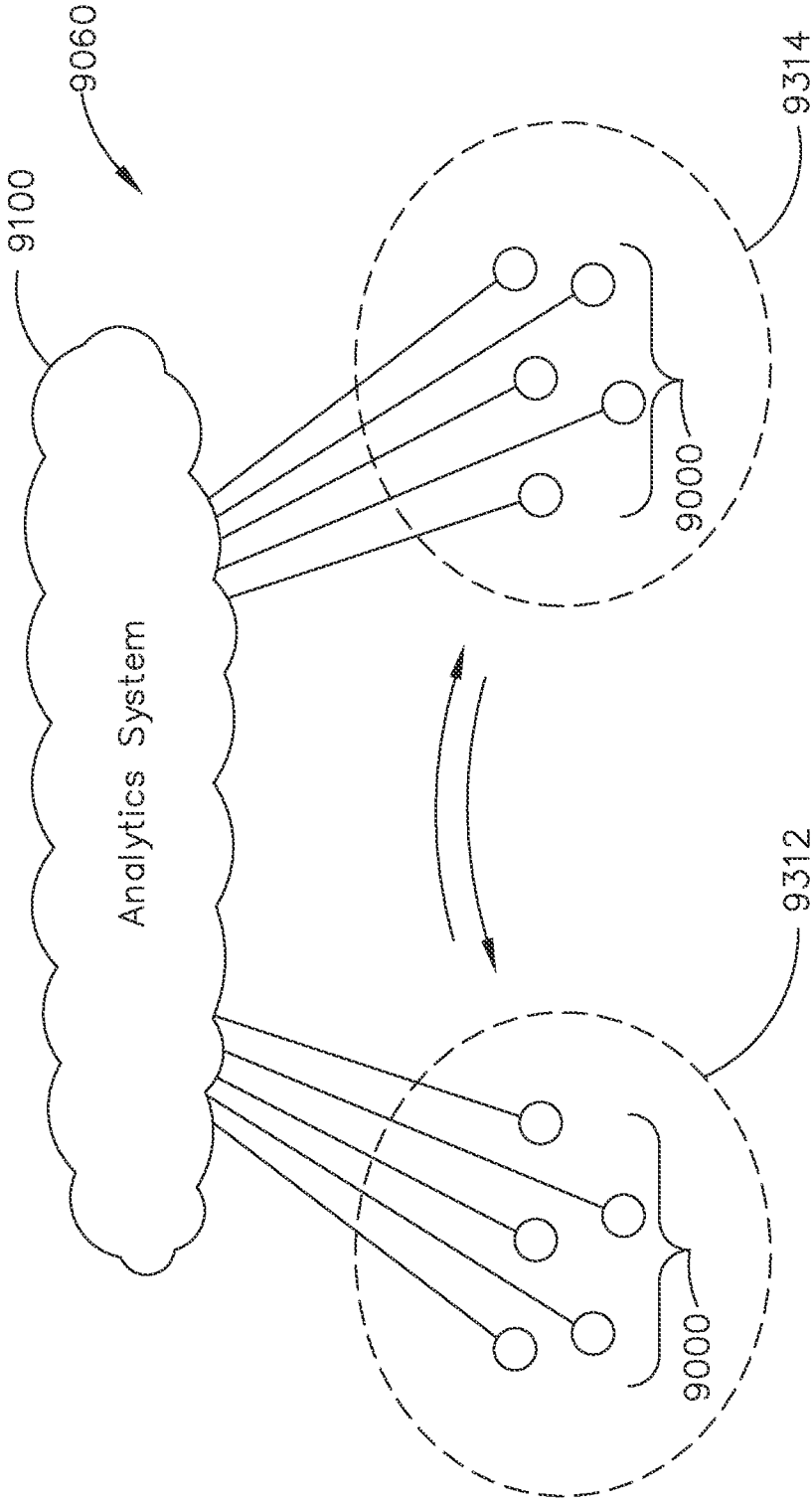


FIG. 194

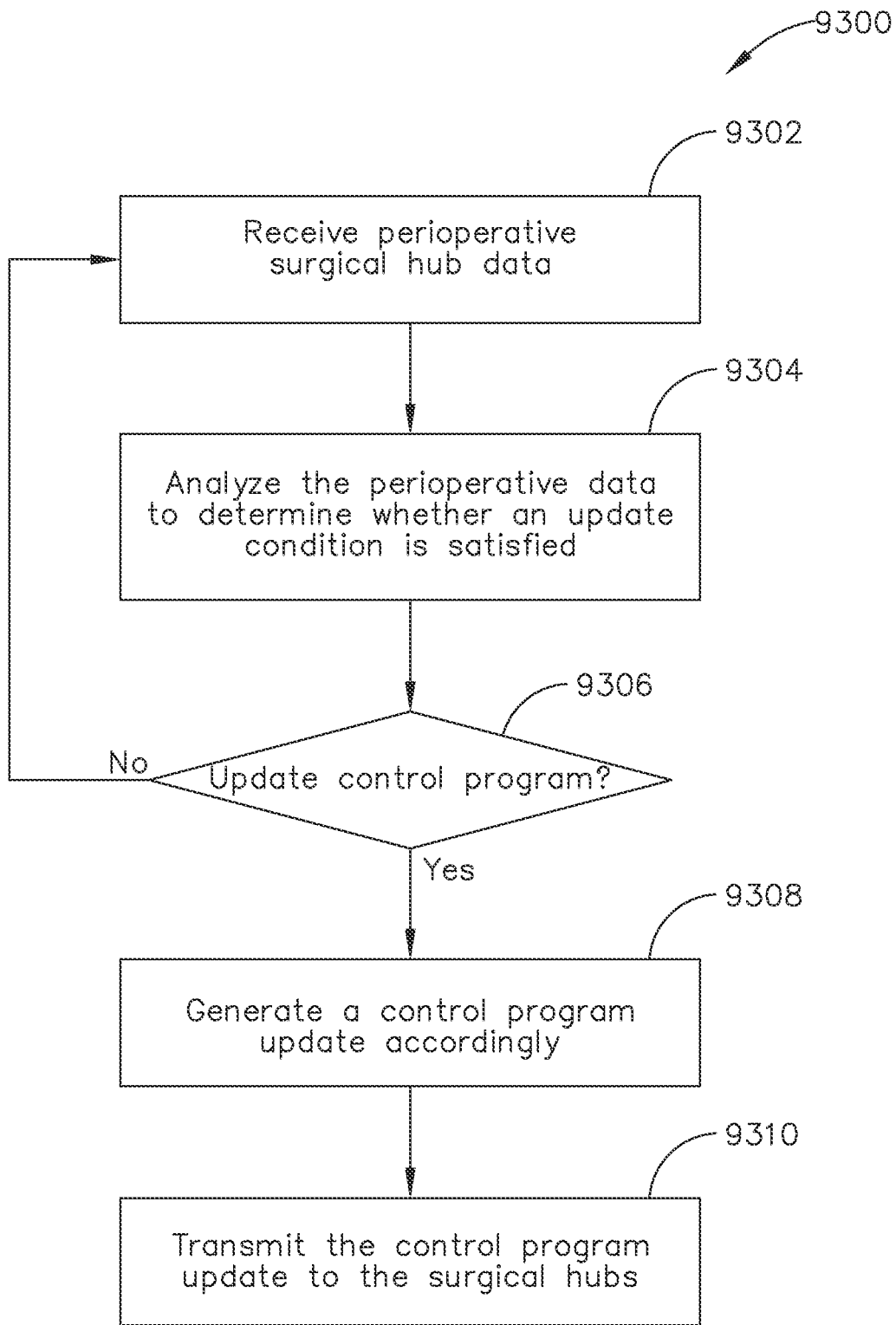


FIG. 195



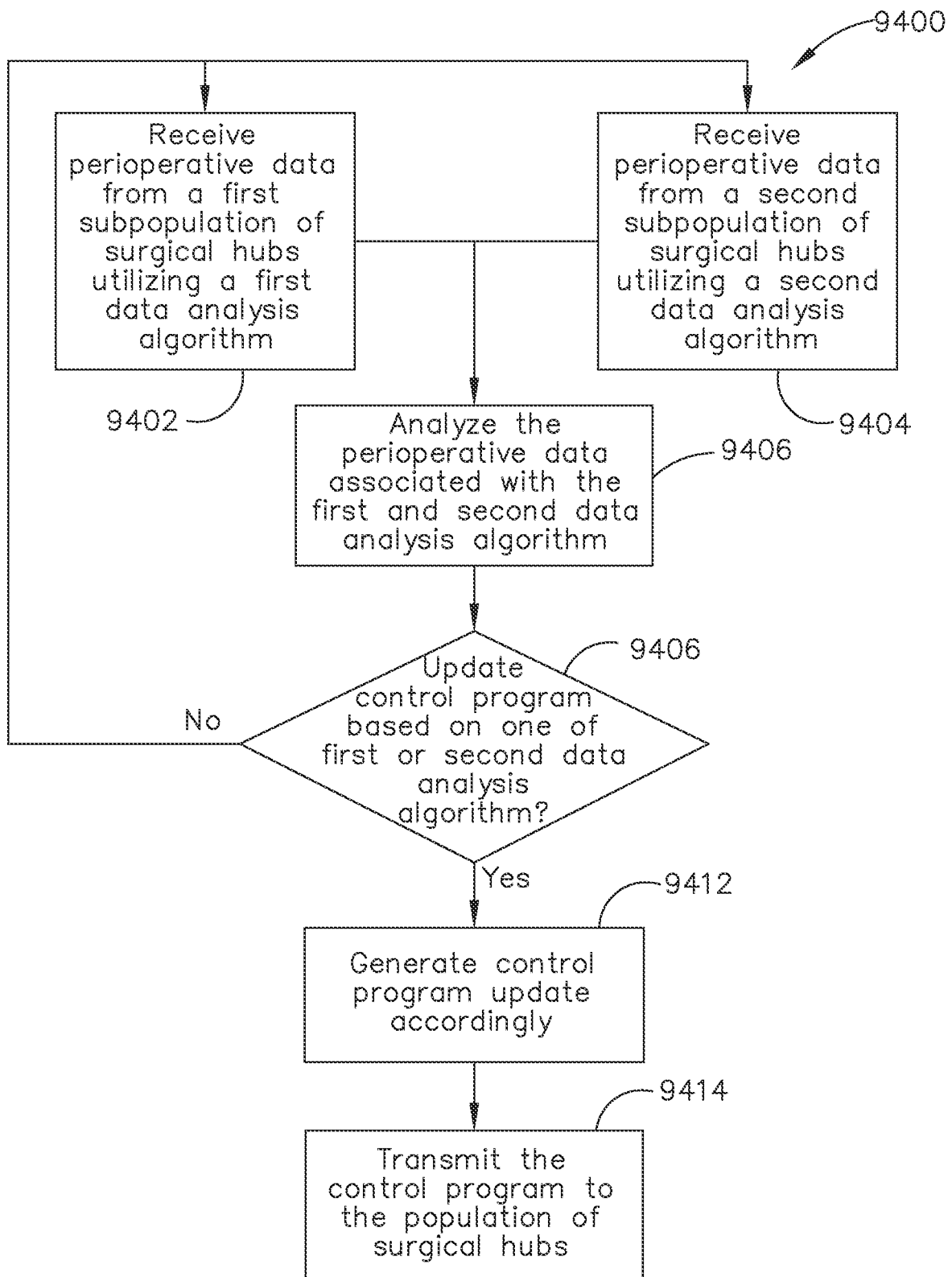
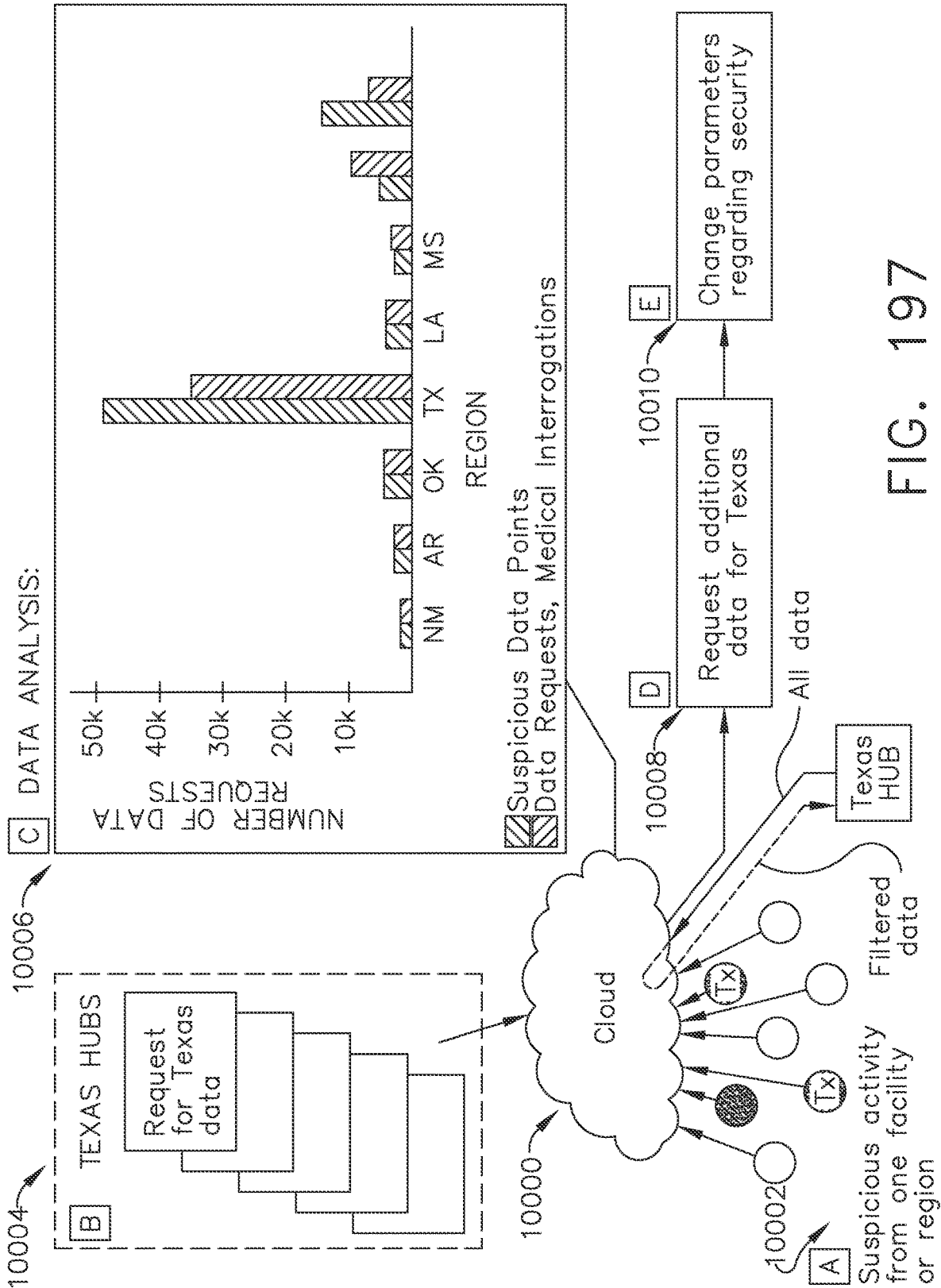
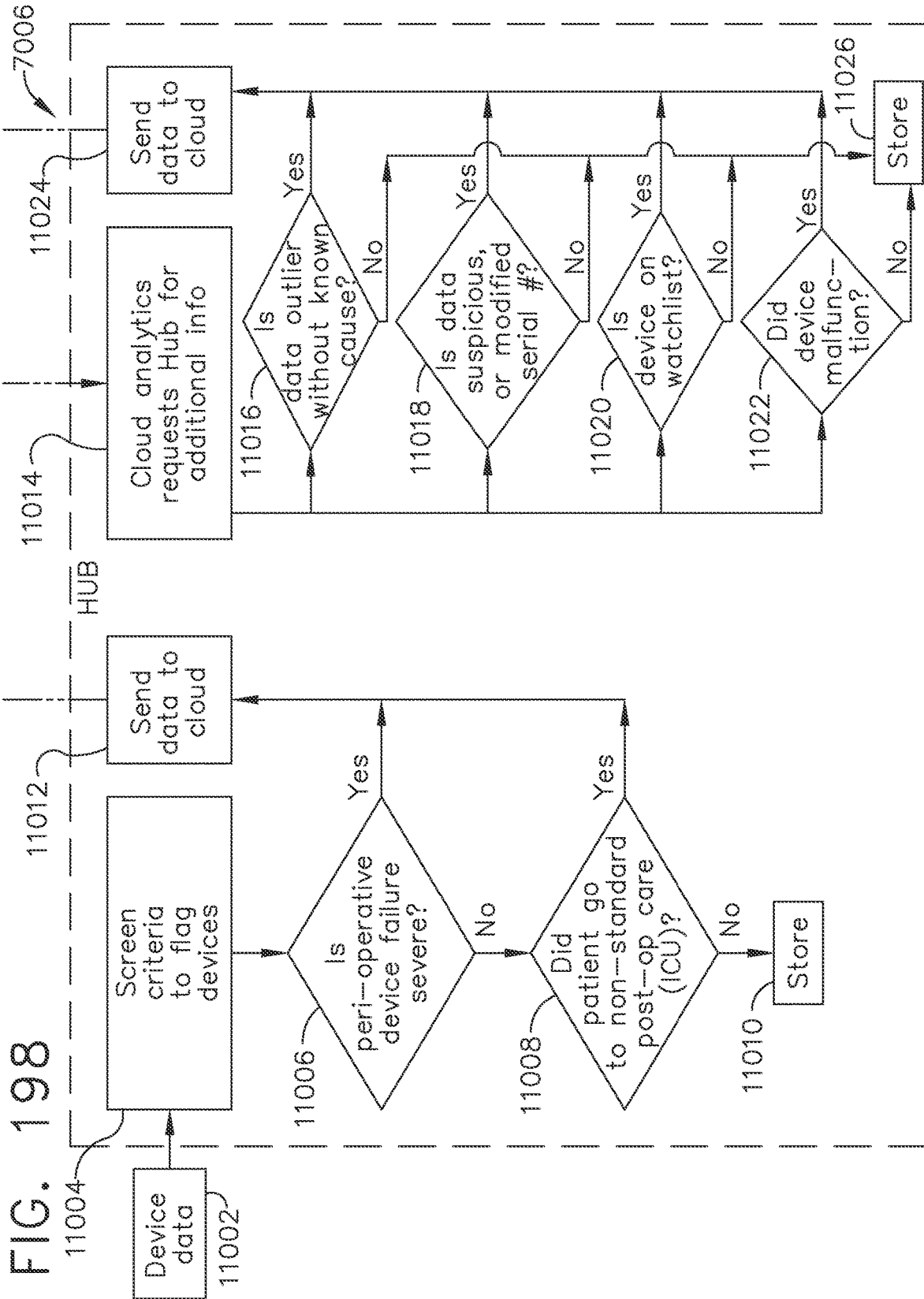
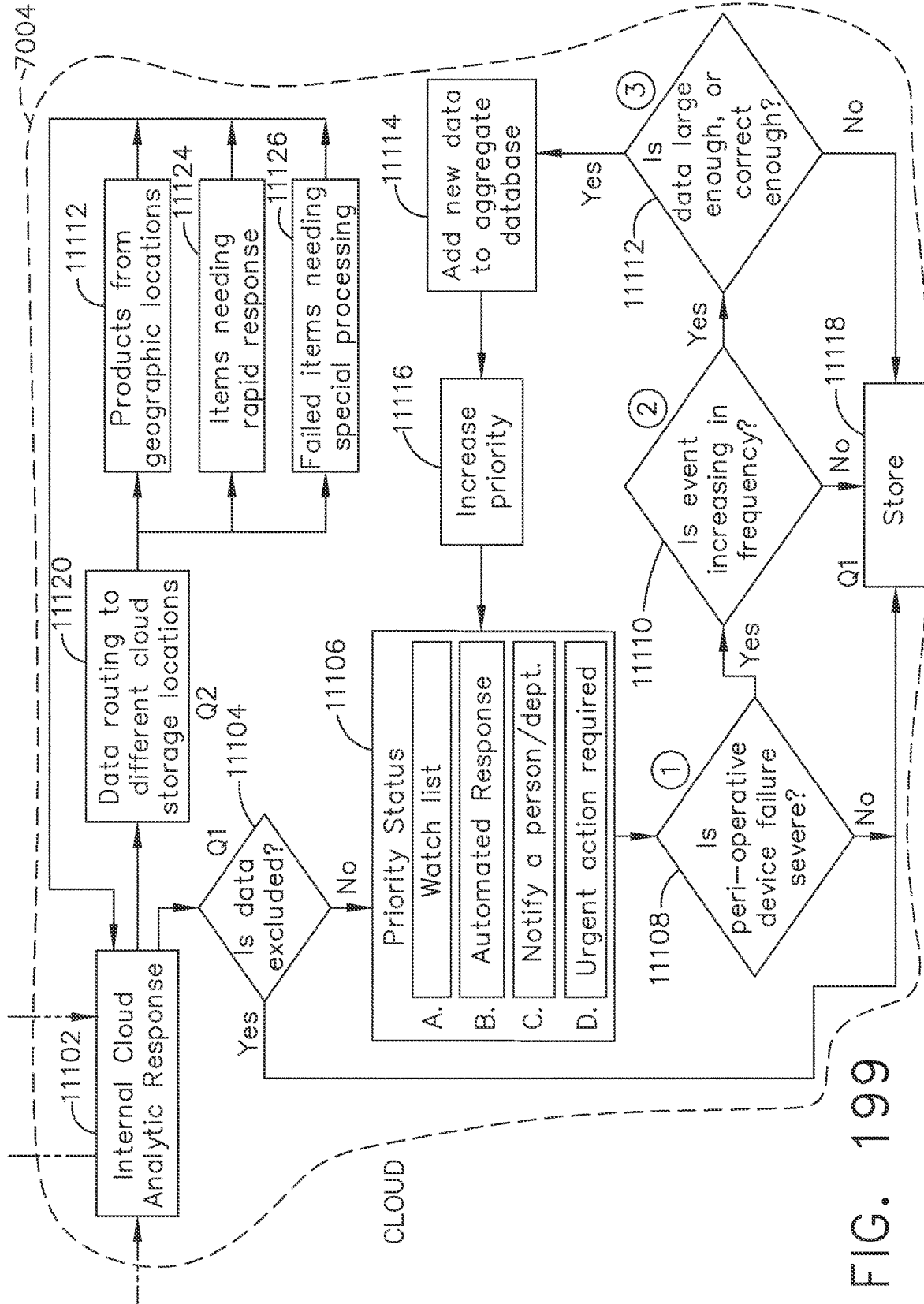


FIG. 196







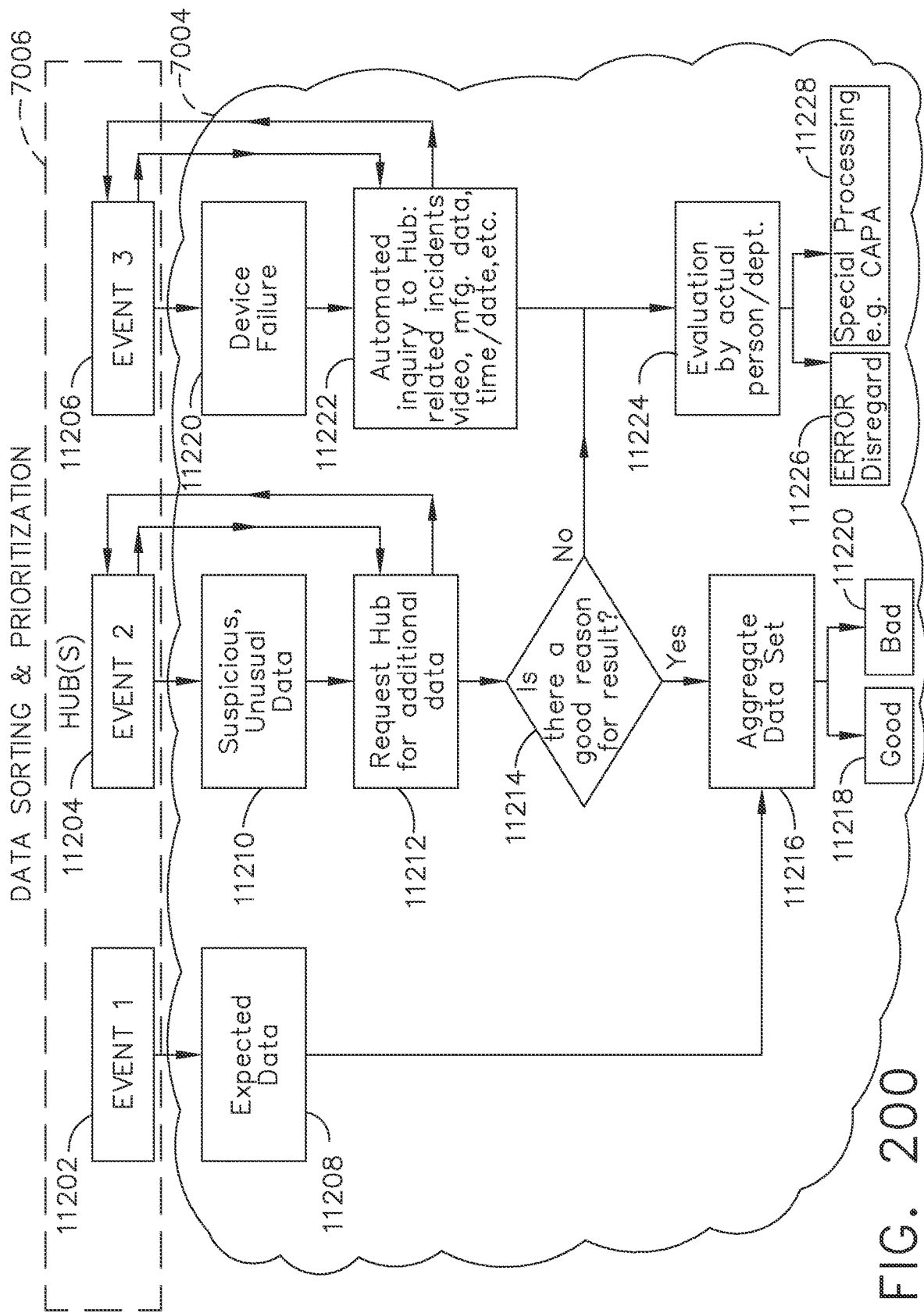


FIG. 200

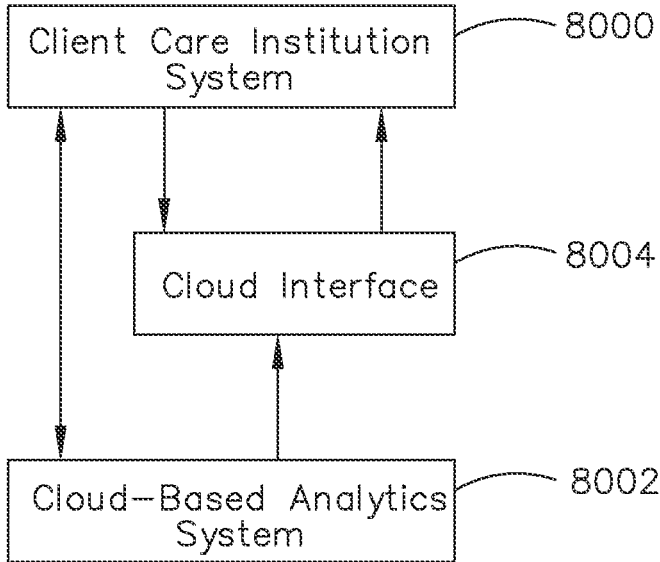


FIG. 201

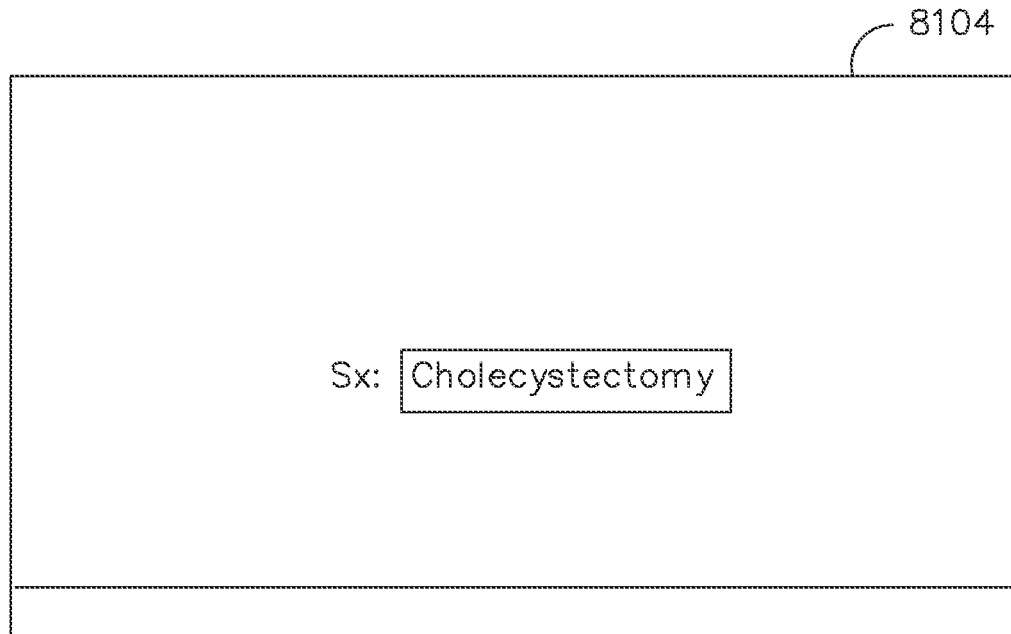


FIG. 202

8106

8104

Item:	Status:	Usage:	Remaining:
Handle A	In use (OR #2)	N/A	N/A
Handle B	Available	235/400	165
Handle C	Inadequate Usage	399/400	1
Handle D	Unauthorized ID	N/A	N/A
Handle E	Expired ID	N/A	N/A

8108

8110

8114

8112

8116

FIG. 203

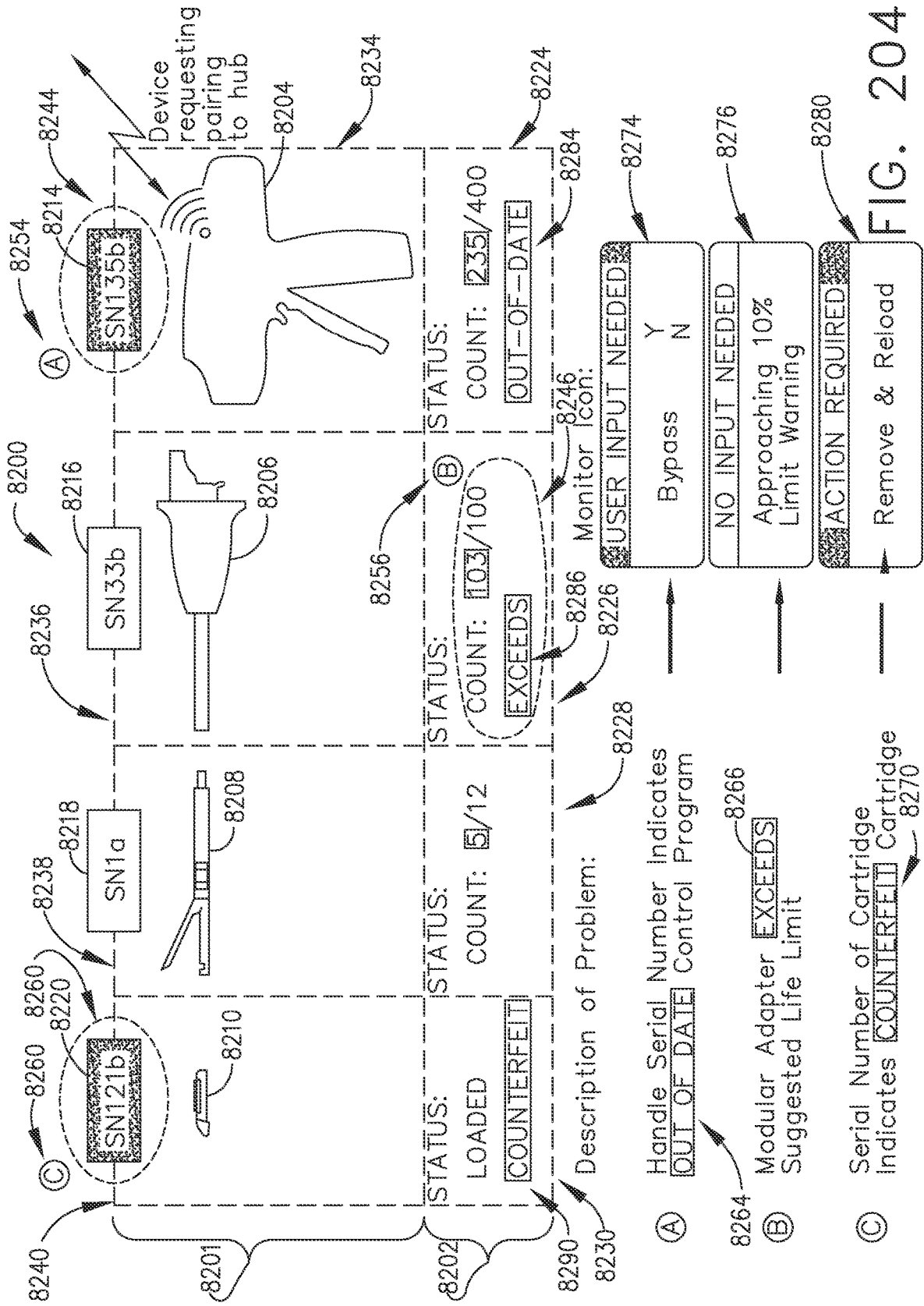


FIG. 204



**METHOD OF HUB COMMUNICATION,  
PROCESSING, DISPLAY, AND CLOUD  
ANALYTICS****CROSS-REFERENCE TO RELATED  
APPLICATIONS**

**[0001]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/773,778, titled METHOD FOR ADAPTIVE CONTROL SCHEMES FOR SURGICAL NETWORK CONTROL AND INTERACTION, filed Nov. 30, 2018, to U.S. Provisional Patent Application No. 62/773,728, titled METHOD FOR SITUATIONAL AWARENESS FOR SURGICAL NETWORK OR SURGICAL NETWORK CONNECTED DEVICE CAPABLE OF ADJUSTING FUNCTION BASED ON A SENSED SITUATION OR USAGE, filed Nov. 30, 2018, to U.S. Provisional Patent Application No. 62/773,741, titled METHOD FOR FACILITY DATA COLLECTION AND INTERPRETATION, filed Nov. 30, 2018, and to U.S. Provisional Patent Application No. 62/773,742, titled METHOD FOR CIRCULAR STAPLER CONTROL ALGORITHM ADJUSTMENT BASED ON SITUATIONAL AWARENESS, filed Nov. 30, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0002]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/750,529, titled METHOD FOR OPERATING A POWERED ARTICULATING MULTI-CLIP APPLIER, filed Oct. 25, 2018, to U.S. Provisional Patent Application No. 62/750,539, titled SURGICAL CLIP APPLIER, filed Oct. 25, 2018, and to U.S. Provisional Patent Application No. 62/750,555, titled SURGICAL CLIP APPLIER, filed Oct. 25, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0003]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/729,183, titled CONTROL FOR A SURGICAL NETWORK OR SURGICAL NETWORK CONNECTED DEVICE THAT ADJUSTS ITS FUNCTION BASED ON A SENSED SITUATION OR USAGE, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,177, titled AUTOMATED DATA SCALING, ALIGNMENT, AND ORGANIZING BASED ON PREDEFINED PARAMETERS WITHIN A SURGICAL NETWORK BEFORE TRANSMISSION, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,176, titled INDIRECT COMMAND AND CONTROL OF A FIRST OPERATING ROOM SYSTEM THROUGH THE USE OF

A SECOND OPERATING ROOM SYSTEM WITHIN A STERILE FIELD WHERE THE SECOND OPERATING ROOM SYSTEM HAS PRIMARY AND SECONDARY OPERATING MODES, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,185, titled POWERED STAPLING DEVICE THAT IS CAPABLE OF ADJUSTING FORCE, ADVANCEMENT SPEED, AND OVERALL STROKE OF CUTTING MEMBER OF THE DEVICE BASED ON SENSED PARAMETER OF FIRING OR CLAMPING, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,184, titled POWERED SURGICAL TOOL WITH A PREDEFINED ADJUSTABLE CONTROL ALGORITHM FOR CONTROLLING AT LEAST ONE END EFFECTOR PARAMETER AND A MEANS FOR LIMITING THE ADJUSTMENT, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,182, titled SENSING THE PATIENT POSITION AND CONTACT UTILIZING THE MONO-POLAR RETURN PAD ELECTRODE TO PROVIDE SITUATIONAL AWARENESS TO THE HUB, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,191, titled SURGICAL NETWORK RECOMMENDATIONS FROM REAL TIME ANALYSIS OF PROCEDURE VARIABLES AGAINST A BASELINE HIGHLIGHTING DIFFERENCES FROM THE OPTIMAL SOLUTION, filed Sep. 10, 2018, to U.S. Provisional Patent Application No. 62/729,195, titled ULTRASONIC ENERGY DEVICE WHICH VARIES PRESSURE APPLIED BY CLAMP ARM TO PROVIDE THRESHOLD CONTROL PRESSURE AT A CUT PROGRESSION LOCATION, filed Sep. 10, 2018, and to U.S. Provisional Patent Application No. 62/729,186, titled WIRELESS PAIRING OF A SURGICAL DEVICE WITH ANOTHER DEVICE WITHIN A STERILE SURGICAL FIELD BASED ON THE USAGE AND SITUATIONAL AWARENESS OF DEVICES, filed Sep. 10, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0004]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/721,995, titled CONTROLLING AN ULTRASONIC SURGICAL INSTRUMENT ACCORDING TO TISSUE LOCATION, filed Aug. 23, 2018, to U.S. Provisional Patent Application No. 62/721,998, titled SITUATIONAL AWARENESS OF ELECTRO-SURGICAL SYSTEMS, filed Aug. 23, 2018, to U.S. Provisional Patent Application No. 62/721,999, titled INTERRUPTION OF ENERGY DUE TO INADVERTENT CAPACITIVE COUPLING, filed Aug. 23, 2018, to U.S. Provisional Patent Application No. 62/721,994, titled BIPOLAR COMBINATION DEVICE THAT AUTOMATICALLY ADJUSTS PRESSURE BASED ON ENERGY MODALITY, filed Aug. 23, 2018, and to U.S. Provisional Patent Application No. 62/721,996, titled RADIO FREQUENCY ENERGY DEVICE FOR DELIVERING COMBINED ELECTRICAL SIGNALS, filed Aug. 23, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0005]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent

application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/692,747, titled SMART ACTIVATION OF AN ENERGY DEVICE BY ANOTHER DEVICE, filed on Jun. 30, 2018, to U.S. Provisional Patent Application No. 62/692,748, titled SMART ENERGY ARCHITECTURE, filed on Jun. 30, 2018, and to U.S. Provisional Patent Application No. 62/692,768, titled SMART ENERGY DEVICES, filed on Jun. 30, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0006]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/691,228, titled METHOD OF USING REINFORCED FLEX CIRCUITS WITH MULTIPLE SENSORS WITH ELECTROSURGICAL DEVICES, filed Jun. 28, 2018, to U.S. Provisional Patent Application No. 62/691,227, titled CONTROLLING A SURGICAL INSTRUMENT ACCORDING TO SENSED CLOSURE PARAMETERS, filed Jun. 28, 2018, to U.S. Provisional Patent Application No. 62/691,230, titled SURGICAL INSTRUMENT HAVING A FLEXIBLE ELECTRODE, filed Jun. 28, 2018, to U.S. Provisional Patent Application No. 62/691,219, titled SURGICAL EVACUATION SENSING AND MOTOR CONTROL, filed Jun. 28, 2018, to U.S. Provisional Patent Application No. 62/691,257, titled COMMUNICATION OF SMOKE EVACUATION SYSTEM PARAMETERS TO HUB OR CLOUD IN SMOKE EVACUATION MODULE FOR INTERACTIVE SURGICAL PLATFORM, filed Jun. 28, 2018, to U.S. Provisional Patent Application No. 62/691,262, titled SURGICAL EVACUATION SYSTEM WITH A COMMUNICATION CIRCUIT FOR COMMUNICATION BETWEEN A FILTER AND A SMOKE EVACUATION DEVICE, filed Jun. 28, 2018, and to U.S. Provisional Patent Application No. 62/691,251, titled DUAL IN-SERIES LARGE AND SMALL DROPLET FILTERS, filed Jun. 28, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0007]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/665,129, titled SURGICAL SUTURING SYSTEMS, filed May 1, 2018, to U.S. Provisional Patent Application No. 62/665,139, titled SURGICAL INSTRUMENTS COMPRISING CONTROL SYSTEMS, filed May 1, 2018, to U.S. Provisional Patent Application No. 62/665,177, titled SURGICAL INSTRUMENTS COMPRISING HANDLE ARRANGEMENTS, filed May 1, 2018, to U.S. Provisional Patent Application No. 62/665,128, titled MODULAR SURGICAL INSTRUMENTS, filed May 1, 2018, to U.S. Provisional Patent

Application No. 62/665,192, titled SURGICAL DISSECTORS, filed May 1, 2018, and to U.S. Provisional Patent Application No. 62/665,134, titled SURGICAL CLIP APPLIER, filed May 1, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0008]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/659,900, titled METHOD OF HUB COMMUNICATION, filed on Apr. 19, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0009]** The present application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/650,898, filed on Mar. 30, 2018, titled CAPACITIVE COUPLED RETURN PATH PAD WITH SEPARABLE ARRAY ELEMENTS, to U.S. Provisional Patent Application No. 62/650,887, titled SURGICAL SYSTEMS WITH OPTIMIZED SENSING CAPABILITIES, filed Mar. 30, 2018, to U.S. Provisional Patent Application No. 62/650,882, titled SMOKE EVACUATION MODULE FOR INTERACTIVE SURGICAL PLATFORM, filed Mar. 30, 2018, and to U.S. Provisional Patent Application No. 62/650,877, titled SURGICAL SMOKE EVACUATION SENSING AND CONTROLS, filed Mar. 30, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

**[0010]** This application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims the benefit of priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/649,302, titled INTERACTIVE SURGICAL SYSTEMS WITH ENCRYPTED COMMUNICATION CAPABILITIES, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,294, titled DATA STRIPPING METHOD TO INTERROGATE PATIENT RECORDS AND CREATE ANONYMIZED RECORD, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,300, titled SURGICAL HUB SITUATIONAL AWARENESS, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,309, titled SURGICAL HUB SPATIAL AWARENESS TO DETERMINE DEVICES IN OPERATING THEATER, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,310, titled COMPUTER IMPLEMENTED INTERACTIVE SURGICAL SYSTEMS, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,291, titled USE OF LASER LIGHT AND RED-GREEN-BLUE COLORATION TO DETERMINE PROPERTIES OF BACK SCATTERED LIGHT, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,296, titled ADAPTIVE CONTROL PROGRAM UPDATES FOR

SURGICAL DEVICES, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,333, titled CLOUD-BASED MEDICAL ANALYTICS FOR CUSTOMIZATION AND RECOMMENDATIONS TO A USER, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,327, titled CLOUD-BASED MEDICAL ANALYTICS FOR SECURITY AND AUTHENTICATION TRENDS AND REACTIVE MEASURES, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,315, titled DATA HANDLING AND PRIORITIZATION IN A CLOUD ANALYTICS NETWORK, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,313, titled CLOUD INTERFACE FOR COUPLED SURGICAL DEVICES, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,320, titled DRIVE ARRANGEMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS, filed Mar. 28, 2018, to U.S. Provisional Patent Application No. 62/649,307, titled AUTOMATIC TOOL ADJUSTMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS, filed Mar. 28, 2018, and to U.S. Provisional Patent Application No. 62/649,323, titled SENSING ARRANGEMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS, filed Mar. 28, 2018, the disclosure of each of which is herein incorporated by reference in its entirety.

[0011] This application is a continuation application claiming priority under 35 U.S.C. § 120 to U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, filed Dec. 4, 2018, now U.S. Patent Application Publication No. 2019/0206562, which also claims the benefit of priority under 35 U.S.C. § 119(e) to U.S. Provisional Patent Application No. 62/611,341, titled INTERACTIVE SURGICAL PLATFORM, filed Dec. 28, 2017, to U.S. Provisional Patent Application No. 62/611,340, titled CLOUD-BASED MEDICAL ANALYTICS, filed Dec. 28, 2017, and to U.S. Provisional Patent Application No. 62/611,339, titled ROBOT ASSISTED SURGICAL PLATFORM, filed Dec. 28, 2017, the disclosure of each of which is herein incorporated by reference in its entirety.

#### BACKGROUND

[0012] The present disclosure relates to various surgical systems. Surgical procedures are typically performed in surgical operating theaters or rooms in a healthcare facility such as, for example, a hospital. A sterile field is typically created around the patient. The sterile field may include the scrubbed team members, who are properly attired, and all furniture and fixtures in the area. Various surgical devices and systems are utilized in performance of a surgical procedure.

[0013] Furthermore, in the Digital and Information Age, medical systems and facilities are often slower to implement systems or procedures utilizing newer and improved technologies due to patient safety and a general desire for maintaining traditional practices. However, often times medical systems and facilities may lack communication and shared knowledge with other neighboring or similarly situated facilities as a result. To improve patient practices, it would be desirable to find ways to help interconnect medical systems and facilities better.

#### SUMMARY

[0014] In one aspect the present disclosure provides a method of displaying an operational parameter of a surgical

system. The method comprising: receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system; analyzing, by the cloud computing system, the first and the second usage data to correlate the first and the second usage data with surgical outcome data; determining, by the cloud computing system, based on the correlation, a recommended medical resource usage configuration; and displaying, on respective displays on the first and the second subset of surgical hubs, indications of the recommended medical resource usage configuration.

[0015] In another aspect the present disclosure provides a method of improving a surgical system, the method comprising: receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system, wherein the first subset of surgical hubs correspond to a first medical facility; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system, wherein the second subset of surgical hubs correspond to a second medical facility; analyzing, by the cloud computing system, the first and the second usage data to correlate the first and the second usage data with surgical outcome data; determining, by the cloud computing system, based on the correlation, that the first usage data is correlated with a first number of positive surgical outcomes and the second usage data is correlated with a second number of positive surgical outcomes, wherein the first number of positive surgical outcomes is greater than the second number of positive surgical outcomes; transmitting, by the cloud computing system, the first usage data to the second subset of surgical hubs; and determining, by the second subset of surgical hubs, an improved medical resource usage configuration based on the first usage data.

[0016] In another aspect the present disclosure provides a method of controlling security of a surgical system, the method comprising: receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system; analyzing, by the cloud computing system, the first and the second usage data to compare the first and the second usage data with security data; determining, by the cloud computing system, based on the comparison, that the first usage data is indicative of a security irregularity and the second usage data is indicative of an acceptable security status; determining, by the cloud computing system, a change in a security parameter based on the indicated security irregularity; transmitting, by the cloud computing system, the change in the security parameter to the second subset of surgical hubs; and changing, by the second subset of surgical hubs, the security parameter.

#### FIGURES

[0017] The features of various aspects are set forth with particularity in the appended claims. The various aspects, however, both as to organization and methods of operation, together with further objects and advantages thereof, may best be understood by reference to the following description, taken in conjunction with the accompanying drawings as follows.

[0018] FIG. 1 is a block diagram of a computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0019] FIG. 2 is a surgical system being used to perform a surgical procedure in an operating room, in accordance with at least one aspect of the present disclosure.

[0020] FIG. 3 is a surgical hub paired with a visualization system, a robotic system, and an intelligent instrument, in accordance with at least one aspect of the present disclosure.

[0021] FIG. 4 is a partial perspective view of a surgical hub enclosure, and of a combo generator module slidably receivable in a drawer of the surgical hub enclosure, in accordance with at least one aspect of the present disclosure.

[0022] FIG. 5 is a perspective view of a combo generator module with bipolar, ultrasonic, and monopolar contacts and a smoke evacuation component, in accordance with at least one aspect of the present disclosure.

[0023] FIG. 6 illustrates individual power bus attachments for a plurality of lateral docking ports of a lateral modular housing configured to receive a plurality of modules, in accordance with at least one aspect of the present disclosure.

[0024] FIG. 7 illustrates a vertical modular housing configured to receive a plurality of modules, in accordance with at least one aspect of the present disclosure.

[0025] FIG. 8 illustrates a surgical data network comprising a modular communication hub configured to connect modular devices located in one or more operating theaters of a healthcare facility, or any room in a healthcare facility specially equipped for surgical operations, to the cloud, in accordance with at least one aspect of the present disclosure.

[0026] FIG. 9 illustrates a computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0027] FIG. 10 illustrates a surgical hub comprising a plurality of modules coupled to the modular control tower, in accordance with at least one aspect of the present disclosure.

[0028] FIG. 11 illustrates one aspect of a Universal Serial Bus (USB) network hub device, in accordance with at least one aspect of the present disclosure.

[0029] FIG. 12 illustrates a logic diagram of a control system of a surgical instrument or tool, in accordance with at least one aspect of the present disclosure.

[0030] FIG. 13 illustrates a control circuit configured to control aspects of the surgical instrument or tool, in accordance with at least one aspect of the present disclosure.

[0031] FIG. 14 illustrates a combinational logic circuit configured to control aspects of the surgical instrument or tool, in accordance with at least one aspect of the present disclosure.

[0032] FIG. 15 illustrates a sequential logic circuit configured to control aspects of the surgical instrument or tool, in accordance with at least one aspect of the present disclosure.

[0033] FIG. 16 illustrates a surgical instrument or tool comprising a plurality of motors which can be activated to perform various functions, in accordance with at least one aspect of the present disclosure.

[0034] FIG. 17 is a schematic diagram of a robotic surgical instrument configured to operate a surgical tool described herein, in accordance with at least one aspect of the present disclosure.

[0035] FIG. 18 illustrates a block diagram of a surgical instrument programmed to control the distal translation of a displacement member, in accordance with at least one aspect of the present disclosure.

[0036] FIG. 19 is a schematic diagram of a surgical instrument configured to control various functions, in accordance with at least one aspect of the present disclosure.

[0037] FIG. 20 is a simplified block diagram of a generator configured to provide inductorless tuning, among other benefits, in accordance with at least one aspect of the present disclosure.

[0038] FIG. 21 illustrates an example of a generator, which is one form of the generator of FIG. 20, in accordance with at least one aspect of the present disclosure.

[0039] FIG. 22 illustrates a combination generator, in accordance with at least one aspect of the present disclosure.

[0040] FIG. 23 illustrates a method of capturing data from a combination generator and communicating the captured generator data to a cloud-based system, in accordance with at least one aspect of the present disclosure.

[0041] FIG. 24 illustrates a data packet of combination generator data, in accordance with at least one aspect of the present disclosure.

[0042] FIG. 25 illustrates an encryption algorithm, in accordance with at least one aspect of the present disclosure.

[0043] FIG. 26 illustrates another encryption algorithm, in accordance with at least one aspect of the present disclosure.

[0044] FIG. 27 illustrates yet another encryption algorithm, in accordance with at least one aspect of the present disclosure.

[0045] FIG. 28 illustrates a high-level representation of a datagram, in accordance with at least one aspect of the present disclosure.

[0046] FIG. 29 illustrates a more detailed representation of the datagram of FIG. 28, in accordance with at least one aspect of the present disclosure.

[0047] FIG. 30 illustrates another representation of the datagram of FIG. 28, in accordance with at least one aspect of the present disclosure.

[0048] FIG. 31 illustrates a method of identifying surgical data associated with a failure event and communicating the identified surgical data to a cloud-based system on a prioritized basis, in accordance with at least one aspect of the present disclosure.

[0049] FIG. 32 illustrates yet another representation of the datagram of FIG. 28, in accordance with at least one aspect of the present disclosure.

[0050] FIG. 33 illustrates a partial artificial timeline of a surgical procedure performed in an operating room via a surgical system, in accordance with at least one aspect of the present disclosure.

[0051] FIG. 34 illustrates ultrasonic pinging of an operating room wall to determine a distance between a surgical hub and the operating room wall, in accordance with at least one aspect of the present disclosure.

[0052] FIG. 35 is a logic flow diagram of a process depicting a control program or a logic configuration for surgical hub pairing with surgical devices of a surgical system that are located within the bounds of an operating room, in accordance with at least one aspect of the present disclosure.

[0053] FIG. 36 is a logic flow diagram of a process depicting a control program or a logic configuration for selectively forming and severing connections between

devices of a surgical system, in accordance with at least one aspect of the present disclosure.

[0054] FIG. 37 is a logic flow diagram of a process depicting a control program or a logic configuration for selectively reevaluating the bounds of an operating room after detecting a new device, in accordance with at least one aspect of the present disclosure.

[0055] FIG. 38 is a logic flow diagram of a process depicting a control program or a logic configuration for selectively reevaluating the bounds of an operating room after disconnection of a paired device, in accordance with at least one aspect of the present disclosure.

[0056] FIG. 39 is a logic flow diagram of a process depicting a control program or a logic configuration for reevaluating the bounds of an operating room by a surgical hub after detecting a change in the position of the surgical hub, in accordance with at least one aspect of the present disclosure.

[0057] FIG. 40 is a logic flow diagram of a process depicting a control program or a logic configuration for selectively forming connections between devices of a surgical system, in accordance with at least one aspect of the present disclosure.

[0058] FIG. 41 is a logic flow diagram of a process depicting a control program or a logic configuration for selectively forming and severing connections between devices of a surgical system, in accordance with at least one aspect of the present disclosure.

[0059] FIG. 42 illustrates a surgical hub pairing a first device and a second device of a surgical system in an operating room, in accordance with at least one aspect of the present disclosure.

[0060] FIG. 43 illustrates a surgical hub unpairing a first device and a second device of a surgical system in an operating room, and pairing the first device with a third device in the operating room, in accordance with at least one aspect of the present disclosure.

[0061] FIG. 44 is a logic flow diagram of a process depicting a control program or a logic configuration for forming an severing connections between devices of a surgical system in an operating room during a surgical procedure based on progression of the steps of the surgical procedure, in accordance with at least one aspect of the present disclosure.

[0062] FIG. 45 is a logic flow diagram of a process depicting a control program or a logic configuration for overlaying information derived from one or more still frames of a livestream of a remote surgical site onto the livestream, in accordance with at least one aspect of the present disclosure.

[0063] FIG. 46 is a logic flow diagram of a process depicting a control program or a logic configuration for differentiating among surgical steps of a surgical procedure, in accordance with at least one aspect of the present disclosure.

[0064] FIG. 47 is a logic flow diagram of a process 3230 depicting a control program or a logic configuration for differentiating among surgical steps of a surgical procedure, in accordance with at least one aspect of the present disclosure.

[0065] FIG. 48 is a logic flow diagram of a process 3240 depicting a control program or a logic configuration for identifying a staple cartridge from information derived from

one or more still frames of staples deployed from the staple cartridge into tissue, in accordance with at least one aspect of the present disclosure.

[0066] FIG. 49 is a partial view of a surgical system in an operating room, the surgical system including a surgical hub that has an imaging module in communication with an imaging device at a remote surgical site, in accordance with at least one aspect of the present disclosure.

[0067] FIG. 50 illustrates a partial view of stapled tissue that received a first staple firing and a second staple firing arranged end-to-end, in accordance with at least one aspect of the present disclosure.

[0068] FIG. 51 illustrates three rows of staples deployed on one side of a tissue stapled and cut by a surgical stapler, in accordance with at least one aspect of the present disclosure.

[0069] FIG. 52 illustrates a non-anodized staple and an anodized staple, in accordance with at least one aspect of the present disclosure.

[0070] FIG. 53 is a logic flow diagram of a process depicting a control program or a logic configuration for coordinating a control arrangement between surgical hubs, in accordance with at least one aspect of the present disclosure.

[0071] FIG. 54 illustrates an interaction between two surgical hubs in an operating room, in accordance with at least one aspect of the present disclosure.

[0072] FIG. 55 is a logic flow diagram of a process depicting a control program or a logic configuration for coordinating a control arrangement between surgical hubs, in accordance with at least one aspect of the present disclosure.

[0073] FIG. 56 illustrates an interaction between two surgical hubs in different operating rooms ("OR1" and "OR3"), in accordance with at least one aspect of the present disclosure.

[0074] FIG. 57 illustrates a secondary display in an operating room ("OR3") showing a surgical site in a colorectal procedure, in accordance with at least one aspect of the present disclosure.

[0075] FIG. 58 illustrates a personal interface or tablet in OR1 displaying the surgical site of OR3, in accordance with at least one aspect of the present disclosure.

[0076] FIG. 59 illustrates an expanded view of the surgical site of OR3 displayed on a primary display of OR1, in accordance with at least one aspect of the present disclosure.

[0077] FIG. 60 illustrates a personal interface or tablet displaying a layout of OR1 that shows available displays, in accordance with at least one aspect of the present disclosure.

[0078] FIG. 61 illustrates a recommendation of a transection location of a surgical site of OR3 made by a surgical operator in OR1 via a personal interface or tablet in OR1, in accordance with at least one aspect of the present disclosure.

[0079] FIG. 62 is a diagram illustrating a technique for interacting with a patient Electronic Medical Record (EMR) database, in accordance with at least one aspect of the present disclosure.

[0080] FIG. 63 illustrates a process of anonymizing a surgical procedure by substituting an artificial time measure for a real time clock for all information stored internally within the instrument, robot, surgical hub, and/or hospital computer equipment, in accordance with at least one aspect of the present disclosure.

[0081] FIG. 64 illustrates ultrasonic pinging of an operating room wall to determine a distance between a surgical hub and the operating room wall, in accordance with at least one aspect of the present disclosure.

[0082] FIG. 65 illustrates a diagram depicting the process of importing patient data stored in an Electronic Medical Record (EMR) database, stripping the patient data, and identifying smart device implications, in accordance with at least one aspect of the present disclosure.

[0083] FIG. 66 illustrates the application of cloud based analytics to redacted and stripped patient data and independent data pairs, in accordance with at least one aspect of the present disclosure.

[0084] FIG. 67 is a logic flow diagram of a process depicting a control program or a logic configuration for associating patient data sets from first and second sources of data, in accordance with at least one aspect of the present disclosure.

[0085] FIG. 68 is a logic flow diagram of a process depicting a control program or a logic configuration for stripping data to extract relevant portions of the data to configure and operate the surgical hub and modules (e.g., instruments) coupled to the surgical hub, in accordance with at least one aspect of the present disclosure.

[0086] FIG. 69 illustrates a self-describing data packet comprising self-describing data, in accordance with at least one aspect of the present disclosure.

[0087] FIG. 70 is a logic flow diagram of a process depicting a control program or a logic configuration for using data packets comprising self-describing data, in accordance with at least one aspect of the present disclosure.

[0088] FIG. 71 is a logic flow diagram of a process depicting a control program or a logic configuration for using data packets comprising self-describing data, in accordance with at least one aspect of the present disclosure.

[0089] FIG. 72 is a diagram of a tumor embedded in the right superior posterior lobe of the right lung, in accordance with at least one aspect of the present disclosure.

[0090] FIG. 73 is a diagram of a lung tumor resection surgical procedure including four separate firings of a surgical stapler to seal and cut bronchial vessels exposed in the fissure leading to and from the upper and lower lobes of the right lung shown in FIG. 72, in accordance with at least one aspect of the present disclosure.

[0091] FIG. 74 is a graphical illustration of a force-to-close (FTC) versus time curve and a force-to-fire (FTF) versus time curve characterizing the first firing of device 002 as shown in FIG. 72, in accordance with at least one aspect of the present disclosure.

[0092] FIG. 75 is a diagram of a staple line visualization laser Doppler to evaluate the integrity of staple line seals by monitoring bleeding of a vessel after a firing of a surgical stapler, in accordance with at least one aspect of the present disclosure.

[0093] FIG. 76 illustrates a paired data set grouped by surgery, in accordance with at least one aspect of the present disclosure.

[0094] FIG. 77 is a diagram of the right lung.

[0095] FIG. 78 is a diagram of the bronchial tree including the trachea and bronchi of the lung.

[0096] FIG. 79 is a logic flow diagram of a process depicting a control program or a logic configuration for storing paired anonymous data sets grouped by surgery, in accordance with at least one aspect of the present disclosure.

[0097] FIG. 80 is a logic flow diagram of a process depicting a control program or a logic configuration for determining rate, frequency, and type of data to transfer to a remote cloud-based analytics network, in accordance with at least one aspect of the present disclosure.

[0098] FIG. 81 illustrates a diagram of a situationally aware surgical system, in accordance with at least one aspect of the present disclosure.

[0099] FIG. 82A illustrates a logic flow diagram of a process for controlling a modular device according to contextual information derived from received data, in accordance with at least one aspect of the present disclosure.

[0100] FIG. 82B illustrates a logic flow diagram of a process for controlling a second modular device according to contextual information derived from perioperative data received from a first modular device, in accordance with at least one aspect of the present disclosure.

[0101] FIG. 82C illustrates a logic flow diagram of a process for controlling a second modular device according to contextual information derived from perioperative data received from a first modular device and the second modular device, in accordance with at least one aspect of the present disclosure.

[0102] FIG. 82D illustrates a logic flow diagram of a process for controlling a third modular device according to contextual information derived from perioperative data received from a first modular device and a second modular device, in accordance with at least one aspect of the present disclosure.

[0103] FIG. 83A illustrates a diagram of a surgical hub communicably coupled to a particular set of modular devices and an Electronic Medical Record (EMR) database, in accordance with at least one aspect of the present disclosure.

[0104] FIG. 83B illustrates a diagram of a smoke evacuator including pressure sensors, in accordance with at least one aspect of the present disclosure.

[0105] FIG. 84A illustrates a logic flow diagram of a process for determining a procedure type according to smoke evacuator perioperative data, in accordance with at least one aspect of the present disclosure.

[0106] FIG. 84B illustrates a logic flow diagram of a process for determining a procedure type according to smoke evacuator, insufflator, and medical imaging device perioperative data, in accordance with at least one aspect of the present disclosure.

[0107] FIG. 84C illustrates a logic flow diagram of a process for determining a procedure type according to medical imaging device perioperative data, in accordance with at least one aspect of the present disclosure.

[0108] FIG. 84D illustrates a logic flow diagram of a process for determining a procedural step according to insufflator perioperative data, in accordance with at least one aspect of the present disclosure.

[0109] FIG. 84E illustrates a logic flow diagram of a process for determining a procedural step according to energy generator perioperative data, in accordance with at least one aspect of the present disclosure.

[0110] FIG. 84F illustrates a logic flow diagram of a process for determining a procedural step according to energy generator perioperative data, in accordance with at least one aspect of the present disclosure.

[0111] FIG. 84G illustrates a logic flow diagram of a process for determining a procedural step according to

stapler perioperative data, in accordance with at least one aspect of the present disclosure.

[0112] FIG. 84H illustrates a logic flow diagram of a process for determining a patient status according to ventilator, pulse oximeter, blood pressure monitor, and/or EKG monitor perioperative data, in accordance with at least one aspect of the present disclosure.

[0113] FIG. 84I illustrates a logic flow diagram of a process for determining a patient status according to pulse oximeter, blood pressure monitor, and/or EKG monitor perioperative data, in accordance with at least one aspect of the present disclosure.

[0114] FIG. 84J illustrates a logic flow diagram of a process for determining a patient status according to ventilator perioperative data, in accordance with at least one aspect of the present disclosure.

[0115] FIG. 85A illustrates a scanner coupled to a surgical hub for scanning a patient wristband, in accordance with at least one aspect of the present disclosure.

[0116] FIG. 85B illustrates a scanner coupled to a surgical hub for scanning a list of surgical items, in accordance with at least one aspect of the present disclosure.

[0117] FIG. 86 illustrates a timeline of an illustrative surgical procedure and the inferences that the surgical hub can make from the data detected at each step in the surgical procedure, in accordance with at least one aspect of the present disclosure.

[0118] FIG. 87A illustrates a flow diagram depicting the process of importing patient data stored in an EMR database and deriving inferences therefrom, in accordance with at least one aspect of the present disclosure.

[0119] FIG. 87B illustrates a flow diagram depicting the process of determining control adjustments corresponding to the derived inferences from FIG. 87A, in accordance with at least one aspect of the present disclosure.

[0120] FIG. 88 illustrates a block diagram of a computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0121] FIG. 89 illustrates a logic flow diagram of tracking data associated with an operating theater event, in accordance with at least one aspect of the present disclosure.

[0122] FIG. 90 illustrates a diagram depicting how the data tracked by the surgical hub can be parsed to provide increasingly detailed metrics, in accordance with at least one aspect of the present disclosure.

[0123] FIG. 91 illustrates a bar graph depicting the number of patients operated on relative to the days of a week for different operating rooms, in accordance with at least one aspect of the present disclosure.

[0124] FIG. 92 illustrates a bar graph depicting the total downtime between procedures relative to the days of a week for a particular operating room, in accordance with at least one aspect of the present disclosure.

[0125] FIG. 93 illustrates a bar graph depicting the total downtime per day of the week depicted in FIG. 92 broken down according to each individual downtime instance, in accordance with at least one aspect of the present disclosure.

[0126] FIG. 94 illustrates a bar graph depicting the average procedure length relative to the days of a week for a particular operating room, in accordance with at least one aspect of the present disclosure.

[0127] FIG. 95 illustrates a bar graph depicting procedure length relative to procedure type, in accordance with at least one aspect of the present disclosure.

[0128] FIG. 96 illustrates a bar graph depicting the average completion time for particular procedural steps for different types of thoracic procedures, in accordance with at least one aspect of the present disclosure.

[0129] FIG. 97 illustrates a bar graph depicting procedure time relative to procedure types, in accordance with at least one aspect of the present disclosure.

[0130] FIG. 98 illustrates a bar graph depicting operating room downtime relative to the time of day, in accordance with at least one aspect of the present disclosure.

[0131] FIG. 99 illustrates a bar graph depicting operating room downtime relative to the day of the week, in accordance with at least one aspect of the present disclosure.

[0132] FIG. 100 illustrates a pair of pie charts depicting the percentage of time that the operating theater is utilized, in accordance with at least one aspect of the present disclosure.

[0133] FIG. 101 illustrates a bar graph depicting consumed and unused surgical items relative to procedure type, in accordance with at least one aspect of the present disclosure.

[0134] FIG. 102 illustrates a logic flow diagram of a process for storing data from the modular devices and patient information database for comparison, in accordance with at least one aspect of the present disclosure.

[0135] FIG. 103 illustrates a diagram of a distributed computing system, in accordance with at least one aspect of the present disclosure.

[0136] FIG. 104 illustrates a logic flow diagram of a process for shifting distributed computing resources, in accordance with at least one aspect of the present disclosure.

[0137] FIG. 105 illustrates a diagram of an imaging system and a surgical instrument bearing a calibration scale, in accordance with at least one aspect of the present disclosure.

[0138] FIG. 106 illustrates a diagram of a surgical instrument centered on a linear staple transection line using the benefit of centering tools and techniques described in connection with FIGS. 107-119, in accordance with at least one aspect of the present disclosure.

[0139] FIGS. 107-109 illustrate a process of aligning an anvil trocar of a circular stapler to a staple overlap portion of a linear staple line created by a double-stapling technique, in accordance with at least one aspect of the present disclosure, where:

[0140] FIG. 107 illustrates an anvil trocar of a circular stapler that is not aligned with a staple overlap portion of a linear staple line created by a double-stapling technique;

[0141] FIG. 108 illustrates an anvil trocar of a circular stapler that is aligned with the center of the staple overlap portion of the linear staple line created by a double-stapling technique; and

[0142] FIG. 109 illustrates a centering tool displayed on a surgical hub display showing a staple overlap portion of a linear staple line created by a double-stapling technique to be cut out by a circular stapler, where the anvil trocar is not aligned with the staple overlap portion of the double staple line as shown in FIG. 107.

[0143] FIGS. 110 and 111 illustrate a before image and an after image of a centering tool, in accordance with at least one aspect of the present disclosure, where:

[0144] FIG. 110 illustrates an image of a projected cut path of an anvil trocar and circular knife before alignment with the target alignment ring circumscribing the image of the

linear staple line over the image of the staple overlap portion presented on a surgical hub display; and

[0145] FIG. 111 illustrates an image of a projected cut path of an anvil trocar and circular knife after alignment with the target alignment ring circumscribing the image of the linear staple line over the image of the staple overlap portion presented on a surgical hub display.

[0146] FIGS. 112-114 illustrate a process of aligning an anvil trocar of a circular stapler to a center of a linear staple line, in accordance with at least one aspect of the present disclosure, where:

[0147] FIG. 112 illustrates the anvil trocar out of alignment with the center of the linear staple line;

[0148] FIG. 113 illustrates the anvil trocar in alignment with the center of the linear staple line; and

[0149] FIG. 114 illustrates a centering tool displayed on a surgical hub display of a linear staple line, where the anvil trocar is not aligned with the staple overlap portion of the double staple line as shown in FIG. 112.

[0150] FIG. 115 is an image of a standard reticle field view of a linear staple line transection of a surgical as viewed through a laparoscope displayed on the surgical hub display, in accordance with at least one aspect of the present disclosure.

[0151] FIG. 116 is an image of a laser-assisted reticle field of view of the surgical site shown in FIG. 115 before the anvil trocar and circular knife of the circular stapler are aligned to the center of the linear staple line, in accordance with at least one aspect of the present disclosure.

[0152] FIG. 117 is an image of a laser-assisted reticle field of view of the surgical site shown in FIG. 116 after the anvil trocar and circular knife of the circular stapler are aligned to the center of the linear staple line, in accordance with at least one aspect of the present disclosure.

[0153] FIG. 118 illustrates a non-contact inductive sensor implementation of a non-contact sensor to determine an anvil trocar location relative to the center of a staple line transection, in accordance with at least one aspect of the present disclosure.

[0154] FIGS. 119A and 119B illustrate one aspect of a non-contact capacitive sensor implementation of the non-contact sensor to determine an anvil trocar location relative to the center of a staple line transection, in accordance with at least one aspect of the present disclosure, where:

[0155] FIG. 119A shows the non-contact capacitive sensor without a nearby metal target; and

[0156] FIG. 119B shows the non-contact capacitive sensor near a metal target.

[0157] FIG. 120 is a logic flow diagram of a process depicting a control program or a logic configuration for aligning a surgical instrument, in accordance with at least one aspect of the present disclosure.

[0158] FIG. 121 illustrates a primary display of the surgical hub comprising a global and local display, in accordance with at least one aspect of the present disclosure.

[0159] FIG. 122 illustrates a primary display of the surgical hub, in accordance with at least one aspect of the present disclosure.

[0160] FIG. 123 illustrates a clamp stabilization sequence over a five second period, in accordance with at least one aspect of the present disclosure.

[0161] FIG. 124 illustrates a diagram of four separate wide angle view images of a surgical site at four separate times during the procedure, in accordance with at least one aspect of the present disclosure.

[0162] FIG. 125 is a graph of tissue creep clamp stabilization curves for two tissue types, in accordance with at least one aspect of the present disclosure.

[0163] FIG. 126 is a graph of time dependent proportionate fill of a clamp force stabilization curve, in accordance with at least one aspect of the present disclosure.

[0164] FIG. 127 is a graph of the role of tissue creep in the clamp force stabilization curve, in accordance with at least one aspect of the present disclosure.

[0165] FIGS. 128A and 128B illustrate two graphs for determining when the clamped tissue has reached creep stability, in accordance with at least one aspect of the present disclosure, where:

[0166] FIG. 128A illustrates a curve that represents a vector tangent angle  $d\theta$  as a function of time; and

[0167] FIG. 128B illustrates a curve that represents change in force-to-close ( $\Delta FTC$ ) as a function of time.

[0168] FIG. 129 illustrates an example of an augmented video image of a pre-operative video image augmented with data identifying displayed elements, in accordance with at least one aspect of the present disclosure.

[0169] FIG. 130 is a logic flow diagram of a process depicting a control program or a logic configuration to display images, in accordance with at least one aspect of the present disclosure.

[0170] FIG. 131 illustrates a communication system comprising an intermediate signal combiner positioned in the communication path between an imaging module and a surgical hub display, in accordance with at least one aspect of the present disclosure.

[0171] FIG. 132 illustrates an independent interactive headset worn by a surgeon to communicate data to the surgical hub, according to one aspect of the present disclosure.

[0172] FIG. 133 illustrates a method for controlling the usage of a device, in accordance with at least one aspect of the present disclosure, in accordance with at least one aspect of the present disclosure.

[0173] FIG. 134 illustrates a surgical system that includes a handle having a controller and a motor, an adapter releasably coupled to the handle, and a loading unit releasably coupled to the adapter, in accordance with at least one aspect of the present disclosure.

[0174] FIG. 135 illustrates a verbal Automated Endoscopic System for Optimal Positioning (AESOP) camera positioning system, in accordance with at least one aspect of the present disclosure.

[0175] FIG. 136 illustrates a multi-functional surgical control system and switching interface for virtual operating room integration, in accordance with at least one aspect of the present disclosure.

[0176] FIG. 137 illustrates a diagram of a beam source and combined beam detector system utilized as a device control mechanism in an operating theater, in accordance with at least one aspect of the present disclosure.

[0177] FIGS. 138A-E illustrate various types of sterile field control and data input consoles, in accordance with at least one aspect of the present disclosure, where:

[0178] FIG. 138A illustrates a single zone sterile field control and data input console;



[0179] FIG. 138B illustrates a multi zone sterile field control and data input console;

[0180] FIG. 138C illustrates a tethered sterile field control and data input console;

[0181] FIG. 138D illustrates a battery operated sterile field control and data input console; and

[0182] FIG. 138E illustrates a battery operated sterile field control and data input console.

[0183] FIGS. 139A-139B illustrate a sterile field console in use in a sterile field during a surgical procedure, in accordance with at least one aspect of the present disclosure, where:

[0184] FIG. 139A shows the sterile field console positioned in the sterile field near two surgeons engaged in an operation; and

[0185] FIG. 139B shows one of the surgeons tapping the touchscreen of the sterile field console.

[0186] FIG. 140 illustrates a process for accepting consult feeds from another operating room, in accordance with at least one aspect of the present disclosure.

[0187] FIG. 141 illustrates a standard technique for estimating vessel path and depth and device trajectory, in accordance with at least one aspect of the present disclosure.

[0188] FIGS. 142A-142D illustrate multiple real time views of images of a virtual anatomical detail for dissection, in accordance with at least one aspect of the present disclosure, where:

[0189] FIG. 142A is a perspective view of the virtual anatomical detail;

[0190] FIG. 142B is a side view of the virtual anatomical detail;

[0191] FIG. 142C is a perspective view of the virtual anatomical detail; and

[0192] FIG. 142D is a side view of the virtual anatomical detail.

[0193] FIGS. 143A-143B illustrate a touchscreen display that may be used within the sterile field, in accordance with at least one aspect of the present disclosure, where:

[0194] FIG. 143A illustrates an image of a surgical site displayed on a touchscreen display in portrait mode;

[0195] FIG. 143B shows the touchscreen display rotated in landscape mode and the surgeon uses his index finger to scroll the image in the direction of the arrows;

[0196] FIG. 143C shows the surgeon using his index finger and thumb to pinch open the image in the direction of the arrows to zoom in;

[0197] FIG. 143D shows the surgeon using his index finger and thumb to pinch close the image in the direction of the arrows to zoom out; and

[0198] FIG. 143E shows the touchscreen display rotated in two directions indicated by arrows to enable the surgeon to view the image in different orientations.

[0199] FIG. 144 illustrates a surgical site employing a smart retractor comprising a direct interface control to a surgical hub, in accordance with at least one aspect of the present disclosure.

[0200] FIG. 145 illustrates a surgical site with a smart flexible sticker display attached to the body of a patient, in accordance with at least one aspect of the present disclosure.

[0201] FIG. 146 is a logic flow diagram of a process depicting a control program or a logic configuration to communicate from inside a sterile field to a device located outside the sterile field, in accordance with at least one aspect of the present disclosure.

[0202] FIG. 147 illustrates a system for performing surgery, in accordance with at least one aspect of the present disclosure.

[0203] FIG. 148 illustrates a second layer of information overlaying a first layer of information, in accordance with at least one aspect of the present disclosure.

[0204] FIG. 149 depicts a perspective view of a surgeon using a surgical instrument that includes a handle assembly housing and a wireless circuit board during a surgical procedure, with the surgeon wearing a set of safety glasses, in accordance with at least one aspect of the present disclosure.

[0205] FIG. 150 is a schematic diagram of a feedback control system for controlling a surgical instrument, in accordance with at least one aspect of the present disclosure.

[0206] FIG. 151 illustrates a feedback controller that includes an on-screen display module and a heads up display (HUD) module, in accordance with at least one aspect of the present disclosure.

[0207] FIG. 152A illustrates a visualization system that may be incorporated into a surgical system, in accordance with at least one aspect of the present disclosure.

[0208] FIG. 152B illustrates a top plan view of a hand unit of the visualization system of FIG. 152A, in accordance with at least one aspect of the present disclosure.

[0209] FIG. 152C illustrates a side plan view of the hand unit depicted in FIG. 152A along with an imaging sensor disposed therein, in accordance with at least one aspect of the present disclosure.

[0210] FIG. 152D illustrates a plurality of an imaging sensors depicted in FIG. 152C, in accordance with at least one aspect of the present disclosure.

[0211] FIG. 153A illustrates a plurality of laser emitters that may be incorporated in the visualization system of FIG. 152A, in accordance with at least one aspect of the present disclosure.

[0212] FIG. 153B illustrates illumination of an image sensor having a Bayer pattern of color filters, in accordance with at least one aspect of the present disclosure.

[0213] FIG. 153C illustrates a graphical representation of the operation of a pixel array for a plurality of frames, in accordance with at least one aspect of the present disclosure.

[0214] FIG. 153D illustrates a schematic of an example of an operation sequence of chrominance and luminance frames, in accordance with at least one aspect of the present disclosure.

[0215] FIG. 153E illustrates an example of sensor and emitter patterns, in accordance with at least one aspect of the present disclosure.

[0216] FIG. 153F illustrates a graphical representation of the operation of a pixel array, in accordance with at least one aspect of the present disclosure.

[0217] FIG. 154 illustrates a schematic of one example of instrumentation for NIR spectroscopy, according to one aspect of the present disclosure.

[0218] FIG. 155 illustrates schematically one example of instrumentation for determining NIRS based on Fourier transform infrared imaging, in accordance with at least one aspect of the present disclosure.

[0219] FIGS. 156A-C illustrate a change in wavelength of light scattered from moving blood cells, in accordance with at least one aspect of the present disclosure.

[0220] FIG. 157 illustrates an aspect of instrumentation that may be used to detect a Doppler shift in laser light

scattered from portions of a tissue, in accordance with at least one aspect of the present disclosure.

[0221] FIG. 158 illustrates schematically some optical effects on light impinging on a tissue having subsurface structures, in accordance with at least one aspect of the present disclosure.

[0222] FIG. 159 illustrates an example of the effects on a Doppler analysis of light impinging on a tissue sample having subsurface structures, in accordance with at least one aspect of the present disclosure.

[0223] FIGS. 160A-C illustrate schematically the detection of moving blood cells at a tissue depth based on a laser Doppler analysis at a variety of laser wavelengths, in accordance with at least one aspect of the present disclosure.

[0224] FIG. 160D illustrates the effect of illuminating a CMOS imaging sensor with a plurality of light wavelengths over time, in accordance with at least one aspect of the present disclosure.

[0225] FIG. 161 illustrates an example of a use of Doppler imaging to detect the present of subsurface blood vessels, in accordance with at least one aspect of the present disclosure.

[0226] FIG. 162 illustrates a method to identify a subsurface blood vessel based on a Doppler shift of blue light due to blood cells flowing therethrough, in accordance with at least one aspect of the present disclosure.

[0227] FIG. 163 illustrates schematically localization of a deep subsurface blood vessel, in accordance with at least one aspect of the present disclosure.

[0228] FIG. 164 illustrates schematically localization of a shallow subsurface blood vessel, in accordance with at least one aspect of the present disclosure.

[0229] FIG. 165 illustrates a composite image comprising a surface image and an image of a subsurface blood vessel, in accordance with at least one aspect of the present disclosure.

[0230] FIG. 166 is a flow chart of a method for determining a depth of a surface feature in a piece of tissue, in accordance with at least one aspect of the present disclosure.

[0231] FIG. 167 illustrates the effect of the location and characteristics of non-vascular structures on light impinging on a tissue sample, in accordance with at least one aspect of the present disclosure.

[0232] FIG. 168 schematically depicts one example of components used in a full field OCT device, in accordance with at least one aspect of the present disclosure.

[0233] FIG. 169 illustrates schematically the effect of tissue anomalies on light reflected from a tissue sample, in accordance with at least one aspect of the present disclosure.

[0234] FIG. 170 illustrates an image display derived from a combination of tissue visualization modalities, in accordance with at least one aspect of the present disclosure.

[0235] FIGS. 171A-C illustrate several aspects of displays that may be provided to a surgeon for a visual identification of a combination of surface and sub-surface structures of a tissue in a surgical site, in accordance with at least one aspect of the present disclosure.

[0236] FIG. 172 is a flow chart of a method for providing information related to a characteristic of a tissue to a smart surgical instrument, in accordance with at least one aspect of the present disclosure.

[0237] FIGS. 173A and 173B illustrate a multi-pixel light sensor receiving by light reflected by a tissue illuminated by sequential exposure to red, green, blue, and infrared light,

and red, green, blue, and ultraviolet laser light sources, respectively, in accordance with at least one aspect of the present disclosure.

[0238] FIGS. 174A and 174B illustrate the distal end of an elongated camera probe having a single light sensor and two light sensors, respectively, in accordance with at least one aspect of the present disclosure.

[0239] FIG. 174C illustrates a perspective view of an example of a monolithic sensor having a plurality of pixel arrays, in accordance with at least one aspect of the present disclosure.

[0240] FIG. 175 illustrates one example of a pair of fields of view available to two image sensors of an elongated camera probe, in accordance with at least one aspect of the present disclosure.

[0241] FIGS. 176A-D illustrate additional examples of a pair of fields of view available to two image sensors of an elongated camera probe, in accordance with at least one aspect of the present disclosure.

[0242] FIGS. 177A-C illustrate an example of the use of an imaging system incorporating the features disclosed in FIG. 176D, in accordance with at least one aspect of the present disclosure.

[0243] FIGS. 178A and 178B depict another example of a dual imaging system, in accordance with at least one aspect of the present disclosure.

[0244] FIGS. 179A-C illustrate examples of a sequence of surgical steps which may benefit from the use of multi-image analysis at the surgical site, in accordance with at least one aspect of the present disclosure.

[0245] FIG. 180 is a block diagram of the computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0246] FIG. 181 is a block diagram which illustrates the functional architecture of the computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0247] FIG. 182 is an example illustration of a tabulation of various resources correlated to particular types of surgical categories, in accordance with at least one aspect of the present disclosure.

[0248] FIG. 183 provides an example illustration of how data is analyzed by the cloud system to provide a comparison between multiple facilities to compare use of resources, in accordance with at least one aspect of the present disclosure.

[0249] FIG. 184 illustrates one example of how the cloud system may determine efficacy trends from an aggregated set of data across whole regions, in accordance with at least one aspect of the present disclosure.

[0250] FIG. 185 provides an example illustration of some types of analysis the cloud system may be configured to perform to provide the predicting modeling, in accordance with at least one aspect of the present disclosure.

[0251] FIG. 186 provides a graphical illustration of a type of example analysis the cloud system may perform to provide these recommendations, in accordance with at least one aspect of the present disclosure.

[0252] FIG. 187 provides an illustration of how the cloud system may conduct analysis to identify a statistical correlation to a local issue that is tied to how a device is used in the localized setting, in accordance with at least one aspect of the present disclosure.

[0253] FIG. 188 provides a graphical illustration of an example of how some devices may satisfy an equivalent use compared to an intended device, and that the cloud system may determine such equivalent use, in accordance with at least one aspect of the present disclosure.

[0254] FIG. 189 provides various examples of how some data may be used as variables in deciding how a post-operative decision tree may branch out, in accordance with at least one aspect of the present disclosure.

[0255] FIG. 190 illustrates a block diagram of a computer-implemented interactive surgical system that is configured to adaptively generate control program updates for modular devices, in accordance with at least one aspect of the present disclosure.

[0256] FIG. 191 illustrates a logic flow diagram of a process for updating the control program of a modular device, in accordance with at least one aspect of the present disclosure.

[0257] FIG. 192 illustrates a diagram of an illustrative analytics system updating a surgical instrument control program, in accordance with at least one aspect of the present disclosure.

[0258] FIG. 193 illustrates a diagram of an analytics system pushing an update to a modular device through a surgical hub, in accordance with at least one aspect of the present disclosure.

[0259] FIG. 194 illustrates a diagram of a computer-implemented interactive surgical system that is configured to adaptively generate control program updates for surgical hubs, in accordance with at least one aspect of the present disclosure.

[0260] FIG. 195 illustrates a logic flow diagram of a process for updating the control program of a surgical hub, in accordance with at least one aspect of the present disclosure.

[0261] FIG. 196 illustrates a logic flow diagram of a process for updating the data analysis algorithm of a control program of a surgical hub, in accordance with at least one aspect of the present disclosure.

[0262] FIG. 197 provides an illustration of example functionality by a cloud medical analytics system for providing improved security and authentication to multiple medical facilities that are interconnected, in accordance with at least one aspect of the present disclosure.

[0263] FIG. 198 is a flow diagram of the computer-implemented interactive surgical system programmed to use screening criteria to determine critical data and to push requests to a surgical hub to obtain additional data, in accordance with at least one aspect of the present disclosure.

[0264] FIG. 199 is a flow diagram of an aspect of responding to critical data by the computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0265] FIG. 200 is a flow diagram of an aspect of data sorting and prioritization by the computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure.

[0266] FIG. 201 illustrates an example system for implementing automated inventory control, in accordance with at least one aspect of the present disclosure.

[0267] FIG. 202 illustrates one example of an institution's cloud interface through which a proposed surgical procedure may be entered, in accordance with at least one aspect of the present disclosure.

[0268] FIG. 203 illustrates one example of an institution's cloud interface through which a cloud-based system provides knowledge regarding the availability and/or usability of inventory items associated with an entered surgical procedure based on system-defined constraints, in accordance with at least one aspect of the present disclosure.

[0269] FIG. 204 illustrates a surgical tool including modular components wherein the status of each modular component is evaluated based on system-defined constraints, in accordance with at least one aspect of the present disclosure.

#### DESCRIPTION

[0270] Applicant of the present application owns the following U.S. Patent Applications, filed on Dec. 4, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:

[0271] U.S. patent application Ser. No. 16/209,416, titled METHOD OF HUB COMMUNICATION, PROCESSING, DISPLAY, AND CLOUD ANALYTICS, now U.S. Patent Application Publication No. 2019/0206562;

[0272] U.S. patent application Ser. No. 16/209,395, titled METHOD OF HUB COMMUNICATION, now U.S. Patent Application Publication No. 2019/0201136;

[0273] U.S. patent application Ser. No. 16/209,403, titled METHOD OF CLOUD BASED DATA ANALYTICS FOR USE WITH THE HUB, now U.S. Patent Application Publication No. 2019/0206569;

[0274] U.S. patent application Ser. No. 16/209,407, titled METHOD OF ROBOTIC HUB COMMUNICATION, DETECTION, AND CONTROL, now U.S. Patent Application Publication No. 2019/0201137;

[0275] U.S. patent application Ser. No. 16/209,423, titled METHOD OF COMPRESSING TISSUE WITHIN A STAPLING DEVICE AND SIMULTANEOUSLY DISPLAYING THE LOCATION OF THE TISSUE WITHIN THE JAWS, now U.S. Patent Application Publication No. 2019/0200981;

[0276] U.S. patent application Ser. No. 16/209,427, titled METHOD OF USING REINFORCED FLEXIBLE CIRCUITS WITH MULTIPLE SENSORS TO OPTIMIZE PERFORMANCE OF RADIO FREQUENCY DEVICES, now U.S. Patent Application Publication No. 2019/0208641;

[0277] U.S. patent application Ser. No. 16/209,433, titled METHOD OF SENSING PARTICULATE FROM SMOKE EVACUATED FROM A PATIENT, ADJUSTING THE PUMP SPEED BASED ON THE SENSED INFORMATION, AND COMMUNICATING THE FUNCTIONAL PARAMETERS OF THE SYSTEM TO THE HUB, now U.S. Patent Application Publication No. 2019/0201594;

[0278] U.S. patent application Ser. No. 16/209,447, titled METHOD FOR SMOKE EVACUATION FOR SURGICAL HUB, now U.S. Patent Application Publication No. 2019/0201045;

[0279] U.S. patent application Ser. No. 16/209,453, titled METHOD FOR CONTROLLING SMART ENERGY DEVICES, now U.S. Patent Application Publication No. 2019/0201046;

[0280] U.S. patent application Ser. No. 16/209,458, titled METHOD FOR SMART ENERGY DEVICE INFRASTRUCTURE, now U.S. Patent Application Publication No. 2019/0201047;

- [0281] U.S. patent application Ser. No. 16/209,465, titled METHOD FOR ADAPTIVE CONTROL SCHEMES FOR SURGICAL NETWORK CONTROL AND INTERACTION, now U.S. Patent Application Publication No. 2019/0206563;
- [0282] U.S. patent application Ser. No. 16/209,478, titled METHOD FOR SITUATIONAL AWARENESS FOR SURGICAL NETWORK OR SURGICAL NETWORK CONNECTED DEVICE CAPABLE OF ADJUSTING FUNCTION BASED ON A SENSED SITUATION OR USAGE, now U.S. Patent Application Publication No. 2019/0104919;
- [0283] U.S. patent application Ser. No. 16/209,490, titled METHOD FOR FACILITY DATA COLLECTION AND INTERPRETATION, now U.S. Patent Application Publication No. 2019/0206564; and
- [0284] U.S. patent application Ser. No. 16/209,491, titled METHOD FOR CIRCULAR STAPLER CONTROL ALGORITHM ADJUSTMENT BASED ON SITUATIONAL AWARENESS, now U.S. Patent Application Publication No. 2019/0200998.
- [0285] Applicant of the present application owns the following U.S. Patent Applications, filed on Nov. 6, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0286] U.S. patent application Ser. No. 16/182,224, titled SURGICAL NETWORK, INSTRUMENT, AND CLOUD RESPONSES BASED ON VALIDATION OF RECEIVED DATASET AND AUTHENTICATION OF ITS SOURCE AND INTEGRITY;
- [0287] U.S. patent application Ser. No. 16/182,230, titled SURGICAL SYSTEM FOR PRESENTING INFORMATION INTERPRETED FROM EXTERNAL DATA;
- [0288] U.S. patent application Ser. No. 16/182,233, titled SURGICAL SYSTEMS WITH AUTONOMOUSLY ADJUSTABLE CONTROL PROGRAMS;
- [0289] U.S. patent application Ser. No. 16/182,239, titled ADJUSTMENT OF DEVICE CONTROL PROGRAMS BASED ON STRATIFIED CONTEXTUAL DATA IN ADDITION TO THE DATA;
- [0290] U.S. patent application Ser. No. 16/182,243, titled SURGICAL HUB AND MODULAR DEVICE RESPONSE ADJUSTMENT BASED ON SITUATIONAL AWARENESS;
- [0291] U.S. patent application Ser. No. 16/182,248, titled DETECTION AND ESCALATION OF SECURITY RESPONSES OF SURGICAL INSTRUMENTS TO INCREASING SEVERITY THREATS;
- [0292] U.S. patent application Ser. No. 16/182,251, titled INTERACTIVE SURGICAL SYSTEM;
- [0293] U.S. patent application Ser. No. 16/182,260, titled AUTOMATED DATA SCALING, ALIGNMENT, AND ORGANIZING BASED ON PREDEFINED PARAMETERS WITHIN SURGICAL NETWORKS;
- [0294] U.S. patent application Ser. No. 16/182,267, titled SENSING THE PATIENT POSITION AND CONTACT UTILIZING THE MONO-POLAR RETURN PAD ELECTRODE TO PROVIDE SITUATIONAL AWARENESS TO THE HUB;
- [0295] U.S. patent application Ser. No. 16/182,249, titled POWERED SURGICAL TOOL WITH PREDEFINED ADJUSTABLE CONTROL ALGORITHM FOR CONTROLLING END EFFECTOR PARAMETER;
- [0296] U.S. patent application Ser. No. 16/182,246, titled ADJUSTMENTS BASED ON AIRBORNE PARTICLE PROPERTIES;
- [0297] U.S. patent application Ser. No. 16/182,256, titled ADJUSTMENT OF A SURGICAL DEVICE FUNCTION BASED ON SITUATIONAL AWARENESS;
- [0298] U.S. patent application Ser. No. 16/182,242, titled REAL-TIME ANALYSIS OF COMPREHENSIVE COST OF ALL INSTRUMENTATION USED IN SURGERY UTILIZING DATA FLUIDITY TO TRACK INSTRUMENTS THROUGH STOCKING AND IN-HOUSE PROCESSES;
- [0299] U.S. patent application Ser. No. 16/182,255, titled USAGE AND TECHNIQUE ANALYSIS OF SURGEON/STAFF PERFORMANCE AGAINST A BASELINE TO OPTIMIZE DEVICE UTILIZATION AND PERFORMANCE FOR BOTH CURRENT AND FUTURE PROCEDURES;
- [0300] U.S. patent application Ser. No. 16/182,269, titled IMAGE CAPTURING OF THE AREAS OUTSIDE THE ABDOMEN TO IMPROVE PLACEMENT AND CONTROL OF A SURGICAL DEVICE IN USE;
- [0301] U.S. patent application Ser. No. 16/182,278, titled COMMUNICATION OF DATA WHERE A SURGICAL NETWORK IS USING CONTEXT OF THE DATA AND REQUIREMENTS OF A RECEIVING SYSTEM/USER TO INFLUENCE INCLUSION OR LINKAGE OF DATA AND METADATA TO ESTABLISH CONTINUITY;
- [0302] U.S. patent application Ser. No. 16/182,290, titled SURGICAL NETWORK RECOMMENDATIONS FROM REAL TIME ANALYSIS OF PROCEDURE VARIABLES AGAINST A BASELINE HIGHLIGHTING DIFFERENCES FROM THE OPTIMAL SOLUTION;
- [0303] U.S. patent application Ser. No. 16/182,232, titled CONTROL OF A SURGICAL SYSTEM THROUGH A SURGICAL BARRIER;
- [0304] U.S. patent application Ser. No. 16/182,227, titled SURGICAL NETWORK DETERMINATION OF PRIORITIZATION OF COMMUNICATION, INTERACTION, OR PROCESSING BASED ON SYSTEM OR DEVICE NEEDS;
- [0305] U.S. patent application Ser. No. 16/182,231, titled WIRELESS PAIRING OF A SURGICAL DEVICE WITH ANOTHER DEVICE WITHIN A STERILE SURGICAL FIELD BASED ON THE USAGE AND SITUATIONAL AWARENESS OF DEVICES;
- [0306] U.S. patent application Ser. No. 16/182,229, titled ADJUSTMENT OF STAPLE HEIGHT OF AT LEAST ONE ROW OF STAPLES BASED ON THE SENSED TISSUE THICKNESS OR FORCE IN CLOSING;
- [0307] U.S. patent application Ser. No. 16/182,234, titled STAPLING DEVICE WITH BOTH COMPULSORY AND DISCRETIONARY LOCKOUTS BASED ON SENSED PARAMETERS;
- [0308] U.S. patent application Ser. No. 16/182,240, titled POWERED STAPLING DEVICE CONFIG-

- URED TO ADJUST FORCE, ADVANCEMENT SPEED, AND OVERALL STROKE OF CUTTING MEMBER BASED ON SENSED PARAMETER OF FIRING OR CLAMPING;
- [0309] U.S. patent application Ser. No. 16/182,235, titled VARIATION OF RADIO FREQUENCY AND ULTRASONIC POWER LEVEL IN COOPERATION WITH VARYING CLAMP ARM PRESSURE TO ACHIEVE PREDEFINED HEAT FLUX OR POWER APPLIED TO TISSUE; and
- [0310] U.S. patent application Ser. No. 16/182,238, titled ULTRASONIC ENERGY DEVICE WHICH VARIES PRESSURE APPLIED BY CLAMP ARM TO PROVIDE THRESHOLD CONTROL PRESSURE AT A CUT PROGRESSION LOCATION.
- [0311] Applicant of the present application owns the following U.S. Patent Applications that were filed on Oct. 26, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0312] U.S. patent application Ser. No. 16/172,303, titled METHOD FOR OPERATING A POWERED ARTICULATING MULTI-CLIP APPLIER;
- [0313] U.S. patent application Ser. No. 16/172,130, titled CLIP APPLIER COMPRISING INTER-CHANGEABLE CLIP RELOADS;
- [0314] U.S. patent application Ser. No. 16/172,066, titled CLIP APPLIER COMPRISING A MOVABLE CLIP MAGAZINE;
- [0315] U.S. patent application Ser. No. 16/172,078, titled CLIP APPLIER COMPRISING A ROTATABLE CLIP MAGAZINE;
- [0316] U.S. patent application Ser. No. 16/172,087, titled CLIP APPLIER COMPRISING CLIP ADVANCING SYSTEMS;
- [0317] U.S. patent application Ser. No. 16/172,094, titled CLIP APPLIER COMPRISING A CLIP CRIMPING SYSTEM;
- [0318] U.S. patent application Ser. No. 16/172,128, titled CLIP APPLIER COMPRISING A RECIPROCATING CLIP ADVANCING MEMBER;
- [0319] U.S. patent application Ser. No. 16/172,168, titled CLIP APPLIER COMPRISING A MOTOR CONTROLLER;
- [0320] U.S. patent application Ser. No. 16/172,164, titled SURGICAL SYSTEM COMPRISING A SURGICAL TOOL AND A SURGICAL HUB;
- [0321] U.S. patent application Ser. No. 16/172,328, titled METHOD OF HUB COMMUNICATION WITH SURGICAL INSTRUMENT SYSTEMS;
- [0322] U.S. patent application Ser. No. 16/172,280, titled METHOD FOR PRODUCING A SURGICAL INSTRUMENT COMPRISING A SMART ELECTRICAL SYSTEM;
- [0323] U.S. patent application Ser. No. 16/172,219, titled METHOD OF HUB COMMUNICATION WITH SURGICAL INSTRUMENT SYSTEMS;
- [0324] U.S. patent application Ser. No. 16/172,248, titled METHOD OF HUB COMMUNICATION WITH SURGICAL INSTRUMENT SYSTEMS;
- [0325] U.S. patent application Ser. No. 16/172,198, titled METHOD OF HUB COMMUNICATION WITH SURGICAL INSTRUMENT SYSTEMS; and
- [0326] U.S. patent application Ser. No. 16/172,155, titled METHOD OF HUB COMMUNICATION WITH SURGICAL INSTRUMENT SYSTEMS.
- [0327] Applicant of the present application owns the following U.S. Patent Applications, filed on Aug. 28, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0328] U.S. patent application Ser. No. 16/115,214, titled ESTIMATING STATE OF ULTRASONIC END EFFECTOR AND CONTROL SYSTEM THEREFOR;
- [0329] U.S. patent application Ser. No. 16/115,205, titled TEMPERATURE CONTROL OF ULTRASONIC END EFFECTOR AND CONTROL SYSTEM THEREFOR;
- [0330] U.S. patent application Ser. No. 16/115,233, titled RADIO FREQUENCY ENERGY DEVICE FOR DELIVERING COMBINED ELECTRICAL SIGNALS;
- [0331] U.S. patent application Ser. No. 16/115,208, titled CONTROLLING AN ULTRASONIC SURGICAL INSTRUMENT ACCORDING TO TISSUE LOCATION;
- [0332] U.S. patent application Ser. No. 16/115,220, titled CONTROLLING ACTIVATION OF AN ULTRASONIC SURGICAL INSTRUMENT ACCORDING TO THE PRESENCE OF TISSUE;
- [0333] U.S. patent application Ser. No. 16/115,232, titled DETERMINING TISSUE COMPOSITION VIA AN ULTRASONIC SYSTEM;
- [0334] U.S. patent application Ser. No. 16/115,239, titled DETERMINING THE STATE OF AN ULTRASONIC ELECTROMECHANICAL SYSTEM ACCORDING TO FREQUENCY SHIFT;
- [0335] U.S. patent application Ser. No. 16/115,247, titled DETERMINING THE STATE OF AN ULTRASONIC END EFFECTOR;
- [0336] U.S. patent application Ser. No. 16/115,211, titled SITUATIONAL AWARENESS OF ELECTRO-SURGICAL SYSTEMS;
- [0337] U.S. patent application Ser. No. 16/115,226, titled MECHANISMS FOR CONTROLLING DIFFERENT ELECTROMECHANICAL SYSTEMS OF AN ELECTROSURGICAL INSTRUMENT;
- [0338] U.S. patent application Ser. No. 16/115,240, titled DETECTION OF END EFFECTOR EMERSION IN LIQUID;
- [0339] U.S. patent application Ser. No. 16/115,249, titled INTERRUPTION OF ENERGY DUE TO INADVERTENT CAPACITIVE COUPLING;
- [0340] U.S. patent application Ser. No. 16/115,256, titled INCREASING RADIO FREQUENCY TO CREATE PAD-LESS MONOPOLAR LOOP;
- [0341] U.S. patent application Ser. No. 16/115,223, titled BIPOLAR COMBINATION DEVICE THAT AUTOMATICALLY ADJUSTS PRESSURE BASED ON ENERGY MODALITY; and
- [0342] U.S. patent application Ser. No. 16/115,238, titled ACTIVATION OF ENERGY DEVICES.
- [0343] Applicant of the present application owns the following U.S. Patent Applications, filed on Aug. 24, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0344] U.S. patent application Ser. No. 16/112,129, titled SURGICAL SUTURING INSTRUMENT CON-

- FIGURED TO MANIPULATE TISSUE USING MECHANICAL AND ELECTRICAL POWER;
- [0345] U.S. patent application Ser. No. 16/112,155, titled SURGICAL SUTURING INSTRUMENT COMPRISING A CAPTURE WIDTH WHICH IS LARGER THAN TROCAR DIAMETER;
- [0346] U.S. patent application Ser. No. 16/112,168, titled SURGICAL SUTURING INSTRUMENT COMPRISING A NON-CIRCULAR NEEDLE;
- [0347] U.S. patent application Ser. No. 16/112,180, titled ELECTRICAL POWER OUTPUT CONTROL BASED ON MECHANICAL FORCES;
- [0348] U.S. patent application Ser. No. 16/112,193, titled REACTIVE ALGORITHM FOR SURGICAL SYSTEM;
- [0349] U.S. patent application Ser. No. 16/112,099, titled SURGICAL INSTRUMENT COMPRISING AN ADAPTIVE ELECTRICAL SYSTEM;
- [0350] U.S. patent application Ser. No. 16/112,112, titled CONTROL SYSTEM ARRANGEMENTS FOR A MODULAR SURGICAL INSTRUMENT;
- [0351] U.S. patent application Ser. No. 16/112,119, titled ADAPTIVE CONTROL PROGRAMS FOR A SURGICAL SYSTEM COMPRISING MORE THAN ONE TYPE OF CARTRIDGE;
- [0352] U.S. patent application Ser. No. 16/112,097, titled SURGICAL INSTRUMENT SYSTEMS COMPRISING BATTERY ARRANGEMENTS;
- [0353] U.S. patent application Ser. No. 16/112,109, titled SURGICAL INSTRUMENT SYSTEMS COMPRISING HANDLE ARRANGEMENTS;
- [0354] U.S. patent application Ser. No. 16/112,114, titled SURGICAL INSTRUMENT SYSTEMS COMPRISING FEEDBACK MECHANISMS;
- [0355] U.S. patent application Ser. No. 16/112,117, titled SURGICAL INSTRUMENT SYSTEMS COMPRISING LOCKOUT MECHANISMS;
- [0356] U.S. patent application Ser. No. 16/112,095, titled SURGICAL INSTRUMENTS COMPRISING A LOCKABLE END EFFECTOR SOCKET;
- [0357] U.S. patent application Ser. No. 16/112,121, titled SURGICAL INSTRUMENTS COMPRISING A SHIFTING MECHANISM;
- [0358] U.S. patent application Ser. No. 16/112,151, titled SURGICAL INSTRUMENTS COMPRISING A SYSTEM FOR ARTICULATION AND ROTATION COMPENSATION;
- [0359] U.S. patent application Ser. No. 16/112,154, titled SURGICAL INSTRUMENTS COMPRISING A BIASED SHIFTING MECHANISM;
- [0360] U.S. patent application Ser. No. 16/112,226, titled SURGICAL INSTRUMENTS COMPRISING AN ARTICULATION DRIVE THAT PROVIDES FOR HIGH ARTICULATION ANGLES;
- [0361] U.S. patent application Ser. No. 16/112,062, titled SURGICAL DISSECTORS AND MANUFACTURING TECHNIQUES;
- [0362] U.S. patent application Ser. No. 16/112,098, titled SURGICAL DISSECTORS CONFIGURED TO APPLY MECHANICAL AND ELECTRICAL ENERGY;
- [0363] U.S. patent application Ser. No. 16/112,237, titled SURGICAL CLIP APPLIER CONFIGURED TO STORE CLIPS IN A STORED STATE;
- [0364] U.S. patent application Ser. No. 16/112,245, titled SURGICAL CLIP APPLIER COMPRISING AN EMPTY CLIP CARTRIDGE LOCKOUT;
- [0365] U.S. patent application Ser. No. 16/112,249, titled SURGICAL CLIP APPLIER COMPRISING AN AUTOMATIC CLIP FEEDING SYSTEM;
- [0366] U.S. patent application Ser. No. 16/112,253, titled SURGICAL CLIP APPLIER COMPRISING ADAPTIVE FIRING CONTROL; and
- [0367] U.S. patent application Ser. No. 16/112,257, titled SURGICAL CLIP APPLIER COMPRISING ADAPTIVE CONTROL IN RESPONSE TO A STRAIN GAUGE CIRCUIT.
- [0368] Applicant of the present application owns the following U.S. Patent Applications, filed on Jun. 29, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0369] U.S. patent application Ser. No. 16/024,090, titled CAPACITIVE COUPLED RETURN PATH PAD WITH SEPARABLE ARRAY ELEMENTS;
- [0370] U.S. patent application Ser. No. 16/024,057, titled CONTROLLING A SURGICAL INSTRUMENT ACCORDING TO SENSED CLOSURE PARAMETERS;
- [0371] U.S. patent application Ser. No. 16/024,067, titled SYSTEMS FOR ADJUSTING END EFFECTOR PARAMETERS BASED ON PERIOPERATIVE INFORMATION;
- [0372] U.S. patent application Ser. No. 16/024,075, titled SAFETY SYSTEMS FOR SMART POWERED SURGICAL STAPLING;
- [0373] U.S. patent application Ser. No. 16/024,083, titled SAFETY SYSTEMS FOR SMART POWERED SURGICAL STAPLING;
- [0374] U.S. patent application Ser. No. 16/024,094, titled SURGICAL SYSTEMS FOR DETECTING END EFFECTOR TISSUE DISTRIBUTION IRREGULARITIES;
- [0375] U.S. patent application Ser. No. 16/024,138, titled SYSTEMS FOR DETECTING PROXIMITY OF SURGICAL END EFFECTOR TO CANCEROUS TISSUE;
- [0376] U.S. patent application Ser. No. 16/024,150, titled SURGICAL INSTRUMENT CARTRIDGE SENSOR ASSEMBLIES;
- [0377] U.S. patent application Ser. No. 16/024,160, titled VARIABLE OUTPUT CARTRIDGE SENSOR ASSEMBLY;
- [0378] U.S. patent application Ser. No. 16/024,124, titled SURGICAL INSTRUMENT HAVING A FLEXIBLE ELECTRODE;
- [0379] U.S. patent application Ser. No. 16/024,132, titled SURGICAL INSTRUMENT HAVING A FLEXIBLE CIRCUIT;
- [0380] U.S. patent application Ser. No. 16/024,141, titled SURGICAL INSTRUMENT WITH A TISSUE MARKING ASSEMBLY;
- [0381] U.S. patent application Ser. No. 16/024,162, titled SURGICAL SYSTEMS WITH PRIORITIZED DATA TRANSMISSION CAPABILITIES;
- [0382] U.S. patent application Ser. No. 16/024,066, titled SURGICAL EVACUATION SENSING AND MOTOR CONTROL;

- [0383] U.S. patent application Ser. No. 16/024,096, titled SURGICAL EVACUATION SENSOR ARRANGEMENTS;
- [0384] U.S. patent application Ser. No. 16/024,116, titled SURGICAL EVACUATION FLOW PATHS;
- [0385] U.S. patent application Ser. No. 16/024,149, titled SURGICAL EVACUATION SENSING AND GENERATOR CONTROL;
- [0386] U.S. patent application Ser. No. 16/024,180, titled SURGICAL EVACUATION SENSING AND DISPLAY;
- [0387] U.S. patent application Ser. No. 16/024,245, titled COMMUNICATION OF SMOKE EVACUATION SYSTEM PARAMETERS TO HUB OR CLOUD IN SMOKE EVACUATION MODULE FOR INTERACTIVE SURGICAL PLATFORM;
- [0388] U.S. patent application Ser. No. 16/024,258, titled SMOKE EVACUATION SYSTEM INCLUDING A SEGMENTED CONTROL CIRCUIT FOR INTERACTIVE SURGICAL PLATFORM;
- [0389] U.S. patent application Ser. No. 16/024,265, titled SURGICAL EVACUATION SYSTEM WITH A COMMUNICATION CIRCUIT FOR COMMUNICATION BETWEEN A FILTER AND A SMOKE EVACUATION DEVICE; and
- [0390] U.S. patent application Ser. No. 16/024,273, titled DUAL IN-SERIES LARGE AND SMALL DROPLET FILTERS.
- [0391] Applicant of the present application owns the following U.S. Patent Applications, filed on Mar. 29, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0392] U.S. patent application Ser. No. 15/940,641, titled INTERACTIVE SURGICAL SYSTEMS WITH ENCRYPTED COMMUNICATION CAPABILITIES;
- [0393] U.S. patent application Ser. No. 15/940,648, titled INTERACTIVE SURGICAL SYSTEMS WITH CONDITION HANDLING OF DEVICES AND DATA CAPABILITIES;
- [0394] U.S. patent application Ser. No. 15/940,656, titled SURGICAL HUB COORDINATION OF CONTROL AND COMMUNICATION OF OPERATING ROOM DEVICES;
- [0395] U.S. patent application Ser. No. 15/940,666, titled SPATIAL AWARENESS OF SURGICAL HUBS IN OPERATING ROOMS;
- [0396] U.S. patent application Ser. No. 15/940,670, titled COOPERATIVE UTILIZATION OF DATA DERIVED FROM SECONDARY SOURCES BY INTELLIGENT SURGICAL HUBS;
- [0397] U.S. patent application Ser. No. 15/940,677, titled SURGICAL HUB CONTROL ARRANGEMENTS;
- [0398] U.S. patent application Ser. No. 15/940,632, titled DATA STRIPPING METHOD TO INTERROGATE PATIENT RECORDS AND CREATE ANONYMIZED RECORD;
- [0399] U.S. patent application Ser. No. 15/940,640, titled COMMUNICATION HUB AND STORAGE DEVICE FOR STORING PARAMETERS AND STATUS OF A SURGICAL DEVICE TO BE SHARED WITH CLOUD BASED ANALYTICS SYSTEMS;
- [0400] U.S. patent application Ser. No. 15/940,645, titled SELF DESCRIBING DATA PACKETS GENERATED AT AN ISSUING INSTRUMENT;
- [0401] U.S. patent application Ser. No. 15/940,649, titled DATA PAIRING TO INTERCONNECT A DEVICE MEASURED PARAMETER WITH AN OUTCOME;
- [0402] U.S. patent application Ser. No. 15/940,654, titled SURGICAL HUB SITUATIONAL AWARENESS;
- [0403] U.S. patent application Ser. No. 15/940,663, titled SURGICAL SYSTEM DISTRIBUTED PROCESSING;
- [0404] U.S. patent application Ser. No. 15/940,668, titled AGGREGATION AND REPORTING OF SURGICAL HUB DATA;
- [0405] U.S. patent application Ser. No. 15/940,671, titled SURGICAL HUB SPATIAL AWARENESS TO DETERMINE DEVICES IN OPERATING THEATER;
- [0406] U.S. patent application Ser. No. 15/940,686, titled DISPLAY OF ALIGNMENT OF STAPLE CARTRIDGE TO PRIOR LINEAR STAPLE LINE;
- [0407] U.S. patent application Ser. No. 15/940,700, titled STERILE FIELD INTERACTIVE CONTROL DISPLAYS;
- [0408] U.S. patent application Ser. No. 15/940,629, titled COMPUTER IMPLEMENTED INTERACTIVE SURGICAL SYSTEMS;
- [0409] U.S. patent application Ser. No. 15/940,704, titled USE OF LASER LIGHT AND RED-GREEN-BLUE COLORATION TO DETERMINE PROPERTIES OF BACK SCATTERED LIGHT;
- [0410] U.S. patent application Ser. No. 15/940,722, titled CHARACTERIZATION OF TISSUE IRREGULARITIES THROUGH THE USE OF MONO-CHROMATIC LIGHT REFRACTIVITY;
- [0411] U.S. patent application Ser. No. 15/940,742, titled DUAL CMOS ARRAY IMAGING;
- [0412] U.S. patent application Ser. No. 15/940,636, titled ADAPTIVE CONTROL PROGRAM UPDATES FOR SURGICAL DEVICES;
- [0413] U.S. patent application Ser. No. 15/940,653, titled ADAPTIVE CONTROL PROGRAM UPDATES FOR SURGICAL HUBS;
- [0414] U.S. patent application Ser. No. 15/940,660, titled CLOUD-BASED MEDICAL ANALYTICS FOR CUSTOMIZATION AND RECOMMENDATIONS TO A USER;
- [0415] U.S. patent application Ser. No. 15/940,679, titled CLOUD-BASED MEDICAL ANALYTICS FOR LINKING OF LOCAL USAGE TRENDS WITH THE RESOURCE ACQUISITION BEHAVIORS OF LARGER DATA SET;
- [0416] U.S. patent application Ser. No. 15/940,694, titled CLOUD-BASED MEDICAL ANALYTICS FOR MEDICAL FACILITY SEGMENTED INDIVIDUALIZATION OF INSTRUMENT FUNCTION;
- [0417] U.S. patent application Ser. No. 15/940,634, titled CLOUD-BASED MEDICAL ANALYTICS FOR SECURITY AND AUTHENTICATION TRENDS AND REACTIVE MEASURES;

- [0418] U.S. patent application Ser. No. 15/940,706, titled DATA HANDLING AND PRIORITIZATION IN A CLOUD ANALYTICS NETWORK;
- [0419] U.S. patent application Ser. No. 15/940,675, titled CLOUD INTERFACE FOR COUPLED SURGICAL DEVICES;
- [0420] U.S. patent application Ser. No. 15/940,627, titled DRIVE ARRANGEMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS;
- [0421] U.S. patent application Ser. No. 15/940,637, titled COMMUNICATION ARRANGEMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS;
- [0422] U.S. patent application Ser. No. 15/940,642, titled CONTROLS FOR ROBOT-ASSISTED SURGICAL PLATFORMS;
- [0423] U.S. patent application Ser. No. 15/940,676, titled AUTOMATIC TOOL ADJUSTMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS;
- [0424] U.S. patent application Ser. No. 15/940,680, titled CONTROLLERS FOR ROBOT-ASSISTED SURGICAL PLATFORMS;
- [0425] U.S. patent application Ser. No. 15/940,683, titled COOPERATIVE SURGICAL ACTIONS FOR ROBOT-ASSISTED SURGICAL PLATFORMS;
- [0426] U.S. patent application Ser. No. 15/940,690, titled DISPLAY ARRANGEMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS; and
- [0427] U.S. patent application Ser. No. 15/940,711, titled SENSING ARRANGEMENTS FOR ROBOT-ASSISTED SURGICAL PLATFORMS.
- [0428] Applicant of the present application owns the following U.S. Provisional Patent Applications, filed on Mar. 8, 2018, the disclosure of each of which is herein incorporated by reference in its entirety:
- [0429] U.S. Provisional Patent Application No. 62/640,417, titled TEMPERATURE CONTROL IN ULTRASONIC DEVICE AND CONTROL SYSTEM THEREFOR; and
- [0430] U.S. Provisional Patent Application No. 62/640,415, titled ESTIMATING STATE OF ULTRASONIC END Effector and Control System Therefor.
- [0431] Before explaining various aspects of surgical devices and generators in detail, it should be noted that the illustrative examples are not limited in application or use to the details of construction and arrangement of parts illustrated in the accompanying drawings and description. The illustrative examples may be implemented or incorporated in other aspects, variations and modifications, and may be practiced or carried out in various ways. Further, unless otherwise indicated, the terms and expressions employed herein have been chosen for the purpose of describing the illustrative examples for the convenience of the reader and are not for the purpose of limitation thereof. Also, it will be appreciated that one or more of the following-described aspects, expressions of aspects, and/or examples, can be combined with any one or more of the other following-described aspects, expressions of aspects and/or examples.
- [0432] Aspects of the present disclosure are presented for a comprehensive digital medical system capable of spanning multiple medical facilities and configured to provide integrated and comprehensive improved medical care to a vast number of patients. The comprehensive digital medical system includes a cloud-based medical analytics system that is configured to interconnect to multiple surgical hubs

located across many different medical facilities. The surgical hubs are configured to interconnect with one or more surgical devices that are used to conduct medical procedures on patients. The surgical hubs provide a wide array of functionality to improve the outcomes of medical procedures. The data generated by the various surgical devices and medical hubs about the patient and the medical procedure may be transmitted to the cloud-based medical analytics system. This data may then be aggregated with similar data gathered from many other surgical hubs and surgical devices located at other medical facilities. Various patterns and correlations may be found through the cloud-based analytics system analyzing the collected data. Improvements in the techniques used to generate the data may be generated as a result, and these improvements may then be disseminated to the various surgical hubs and surgical devices. Due to the interconnectedness of all of the aforementioned components, improvements in medical procedures and practices may be found that otherwise may not be found if the many components were not so interconnected. Various examples of structures and functions of these various components will be described in more detail in the following description.

[0433] Referring to FIG. 1, a computer-implemented interactive surgical system 100 includes one or more surgical systems 102 and a cloud-based system (e.g., the cloud 104 that may include a remote server 113 coupled to a storage device 105). Each surgical system 102 includes at least one surgical hub 106 in communication with the cloud 104 that may include a remote server 113. In one example, as illustrated in FIG. 1, the surgical system 102 includes a visualization system 108, a robotic system 110, and a handheld intelligent surgical instrument 112, which are configured to communicate with one another and/or the hub 106. In some aspects, a surgical system 102 may include an M number of hubs 106, an N number of visualization systems 108, an O number of robotic systems 110, and a P number of handheld intelligent surgical instruments 112, where M, N, O, and P are integers greater than or equal to one.

[0434] FIG. 3 depicts an example of a surgical system 102 being used to perform a surgical procedure on a patient who is lying down on an operating table 114 in a surgical operating room 116. A robotic system 110 is used in the surgical procedure as a part of the surgical system 102. The robotic system 110 includes a surgeon's console 118, a patient side cart 120 (surgical robot), and a surgical robotic hub 122. The patient side cart 120 can manipulate at least one removably coupled surgical tool 117 through a minimally invasive incision in the body of the patient while the surgeon views the surgical site through the surgeon's console 118. An image of the surgical site can be obtained by a medical imaging device 124, which can be manipulated by the patient side cart 120 to orient the imaging device 124. The robotic hub 122 can be used to process the images of the surgical site for subsequent display to the surgeon through the surgeon's console 118.

[0435] Other types of robotic systems can be readily adapted for use with the surgical system 102. Various examples of robotic systems and surgical tools that are suitable for use with the present disclosure are described in U.S. Provisional Patent Application Ser. No. 62/611,339, titled ROBOT ASSISTED SURGICAL PLATFORM, filed Dec. 28, 2017, the disclosure of which is herein incorporated by reference in its entirety.



**[0436]** Various examples of cloud-based analytics that are performed by the cloud **104**, and are suitable for use with the present disclosure, are described in U.S. Provisional Patent Application Ser. No. 62/611,340, titled CLOUD-BASED MEDICAL ANALYTICS, filed Dec. 28, 2017, the disclosure of which is herein incorporated by reference in its entirety.

**[0437]** In various aspects, the imaging device **124** includes at least one image sensor and one or more optical components. Suitable image sensors include, but are not limited to, Charge-Coupled Device (CCD) sensors and Complementary Metal-Oxide Semiconductor (CMOS) sensors.

**[0438]** The optical components of the imaging device **124** may include one or more illumination sources and/or one or more lenses. The one or more illumination sources may be directed to illuminate portions of the surgical field. The one or more image sensors may receive light reflected or refracted from the surgical field, including light reflected or refracted from tissue and/or surgical instruments.

**[0439]** The one or more illumination sources may be configured to radiate electromagnetic energy in the visible spectrum as well as the invisible spectrum. The visible spectrum, sometimes referred to as the optical spectrum or luminous spectrum, is that portion of the electromagnetic spectrum that is visible to (i.e., can be detected by) the human eye and may be referred to as visible light or simply light. A typical human eye will respond to wavelengths in air that are from about 380 nm to about 750 nm.

**[0440]** The invisible spectrum (i.e., the non-luminous spectrum) is that portion of the electromagnetic spectrum that lies below and above the visible spectrum (i.e., wavelengths below about 380 nm and above about 750 nm). The invisible spectrum is not detectable by the human eye. Wavelengths greater than about 750 nm are longer than the red visible spectrum, and they become invisible infrared (IR), microwave, and radio electromagnetic radiation. Wavelengths less than about 380 nm are shorter than the violet spectrum, and they become invisible ultraviolet, x-ray, and gamma ray electromagnetic radiation.

**[0441]** In various aspects, the imaging device **124** is configured for use in a minimally invasive procedure. Examples of imaging devices suitable for use with the present disclosure include, but not limited to, an arthroscope, angioscope, bronchoscope, choledochoscope, colonoscope, cytoscope, duodenoscope, enteroscope, esophago-gastro-duodenoscope (gastroscope), endoscope, laryngoscope, nasopharyngo-neproscope, sigmoidoscope, thoracoscope, and ureteroscopy.

**[0442]** In one aspect, the imaging device employs multi-spectrum monitoring to discriminate topography and underlying structures. A multi-spectral image is one that captures image data within specific wavelength ranges across the electromagnetic spectrum. The wavelengths may be separated by filters or by the use of instruments that are sensitive to particular wavelengths, including light from frequencies beyond the visible light range, e.g., IR and ultraviolet. Spectral imaging can allow extraction of additional information the human eye fails to capture with its receptors for red, green, and blue. The use of multi-spectral imaging is described in greater detail under the heading "Advanced Imaging Acquisition Module" in U.S. Provisional Patent Application Ser. No. 62/611,341, titled INTERACTIVE SURGICAL PLATFORM, filed Dec. 28, 2017, the disclosure of which is herein incorporated by reference in its

entirety. Multi-spectrum monitoring can be a useful tool in relocating a surgical field after a surgical task is completed to perform one or more of the previously described tests on the treated tissue.

**[0443]** It is axiomatic that strict sterilization of the operating room and surgical equipment is required during any surgery. The strict hygiene and sterilization conditions required in a "surgical theater," i.e., an operating or treatment room, necessitate the highest possible sterility of all medical devices and equipment. Part of that sterilization process is the need to sterilize anything that comes in contact with the patient or penetrates the sterile field, including the imaging device **124** and its attachments and components. It will be appreciated that the sterile field may be considered a specified area, such as within a tray or on a sterile towel, that is considered free of microorganisms, or the sterile field may be considered an area, immediately around a patient, who has been prepared for a surgical procedure. The sterile field may include the scrubbed team members, who are properly attired, and all furniture and fixtures in the area.

**[0444]** In various aspects, the visualization system **108** includes one or more imaging sensors, one or more image-processing units, one or more storage arrays, and one or more displays that are strategically arranged with respect to the sterile field, as illustrated in FIG. 2. In one aspect, the visualization system **108** includes an interface for HL7, PACS, and EMR. Various components of the visualization system **108** are described under the heading "Advanced Imaging Acquisition Module" in U.S. Provisional Patent Application Ser. No. 62/611,341, titled INTERACTIVE SURGICAL PLATFORM, filed Dec. 28, 2017, the disclosure of which is herein incorporated by reference in its entirety.

**[0445]** As illustrated in FIG. 2, a primary display **119** is positioned in the sterile field to be visible to an operator at the operating table **114**. In addition, a visualization tower **111** is positioned outside the sterile field. The visualization tower **111** includes a first non-sterile display **107** and a second non-sterile display **109**, which face away from each other. The visualization system **108**, guided by the hub **106**, is configured to utilize the displays **107**, **109**, and **119** to coordinate information flow to operators inside and outside the sterile field. For example, the hub **106** may cause the visualization system **108** to display a snapshot of a surgical site, as recorded by an imaging device **124**, on a non-sterile display **107** or **109**, while maintaining a live feed of the surgical site on the primary display **119**. The snapshot on the non-sterile display **107** or **109** can permit a non-sterile operator to perform a diagnostic step relevant to the surgical procedure, for example.

**[0446]** In one aspect, the hub **106** is also configured to route a diagnostic input or feedback entered by a non-sterile operator at the visualization tower **111** to the primary display **119** within the sterile field, where it can be viewed by a sterile operator at the operating table. In one example, the input can be in the form of a modification to the snapshot displayed on the non-sterile display **107** or **109**, which can be routed to the primary display **119** by the hub **106**.

**[0447]** Referring to FIG. 2, a surgical instrument **112** is being used in the surgical procedure as part of the surgical system **102**. The hub **106** is also configured to coordinate information flow to a display of the surgical instrument **112**. For example, in U.S. Provisional Patent Application Ser. No. 62/611,341, titled INTERACTIVE SURGICAL PLAT-

FORM, filed Dec. 28, 2017, the disclosure of which is herein incorporated by reference in its entirety. A diagnostic input or feedback entered by a non-sterile operator at the visualization tower **111** can be routed by the hub **106** to the surgical instrument display **115** within the sterile field, where it can be viewed by the operator of the surgical instrument **112**. Example surgical instruments that are suitable for use with the surgical system **102** are described under the heading “Surgical Instrument Hardware” and in U.S. Provisional Patent Application Ser. No. 62/611,341, titled INTERACTIVE SURGICAL PLATFORM, filed Dec. 28, 2017, the disclosure of which is herein incorporated by reference in its entirety, for example.

[0448] Referring now to FIG. 3, a hub **106** is depicted in communication with a visualization system **108**, a robotic system **110**, and a handheld intelligent surgical instrument **112**. The hub **106** includes a hub display **135**, an imaging module **138**, a generator module **140**, a communication module **130**, a processor module **132**, and a storage array **134**. In certain aspects, as illustrated in FIG. 3, the hub **106** further includes a smoke evacuation module **126** and/or a suction/irrigation module **128**.

[0449] During a surgical procedure, energy application to tissue, for sealing and/or cutting, is generally associated with smoke evacuation, suction of excess fluid, and/or irrigation of the tissue. Fluid, power, and/or data lines from different sources are often entangled during the surgical procedure. Valuable time can be lost addressing this issue during a surgical procedure. Detangling the lines may necessitate disconnecting the lines from their respective modules, which may require resetting the modules. The hub modular enclosure **136** offers a unified environment for managing the power, data, and fluid lines, which reduces the frequency of entanglement between such lines.

[0450] Aspects of the present disclosure present a surgical hub for use in a surgical procedure that involves energy application to tissue at a surgical site. The surgical hub includes a hub enclosure and a combo generator module slidably receivable in a docking station of the hub enclosure. The docking station includes data and power contacts. The combo generator module includes two or more of an ultrasonic energy generator component, a bipolar RF energy generator component, and a monopolar RF energy generator component that are housed in a single unit. In one aspect, the combo generator module also includes a smoke evacuation component, at least one energy delivery cable for connecting the combo generator module to a surgical instrument, at least one smoke evacuation component configured to evacuate smoke, fluid, and/or particulates generated by the application of therapeutic energy to the tissue, and a fluid line extending from the remote surgical site to the smoke evacuation component.

[0451] In one aspect, the fluid line is a first fluid line and a second fluid line extends from the remote surgical site to a suction and irrigation module slidably received in the hub enclosure. In one aspect, the hub enclosure comprises a fluid interface.

[0452] Certain surgical procedures may require the application of more than one energy type to the tissue. One energy type may be more beneficial for cutting the tissue, while another different energy type may be more beneficial for sealing the tissue. For example, a bipolar generator can be used to seal the tissue while an ultrasonic generator can be used to cut the sealed tissue. Aspects of the present

disclosure present a solution where a hub modular enclosure **136** is configured to accommodate different generators, and facilitate an interactive communication therebetween. One of the advantages of the hub modular enclosure **136** is enabling the quick removal and/or replacement of various modules.

[0453] Aspects of the present disclosure present a modular surgical enclosure for use in a surgical procedure that involves energy application to tissue. The modular surgical enclosure includes a first energy-generator module, configured to generate a first energy for application to the tissue, and a first docking station comprising a first docking port that includes first data and power contacts, wherein the first energy-generator module is slidably movable into an electrical engagement with the power and data contacts and wherein the first energy-generator module is slidably movable out of the electrical engagement with the first power and data contacts,

[0454] Further to the above, the modular surgical enclosure also includes a second energy-generator module configured to generate a second energy, different than the first energy, for application to the tissue, and a second docking station comprising a second docking port that includes second data and power contacts, wherein the second energy-generator module is slidably movable into an electrical engagement with the power and data contacts, and wherein the second energy-generator module is slidably movable out of the electrical engagement with the second power and data contacts.

[0455] In addition, the modular surgical enclosure also includes a communication bus between the first docking port and the second docking port, configured to facilitate communication between the first energy-generator module and the second energy-generator module.

[0456] Referring to FIGS. 3-7, aspects of the present disclosure are presented for a hub modular enclosure **136** that allows the modular integration of a generator module **140**, a smoke evacuation module **126**, and a suction/irrigation module **128**. The hub modular enclosure **136** further facilitates interactive communication between the modules **140**, **126**, **128**. As illustrated in FIG. 5, the generator module **140** can be a generator module with integrated monopolar, bipolar, and ultrasonic components supported in a single housing unit **139** slidably insertable into the hub modular enclosure **136**. As illustrated in FIG. 5, the generator module **140** can be configured to connect to a monopolar device **146**, a bipolar device **147**, and an ultrasonic device **148**. Alternatively, the generator module **140** may comprise a series of monopolar, bipolar, and/or ultrasonic generator modules that interact through the hub modular enclosure **136**. The hub modular enclosure **136** can be configured to facilitate the insertion of multiple generators and interactive communication between the generators docked into the hub modular enclosure **136** so that the generators would act as a single generator.

[0457] In one aspect, the hub modular enclosure **136** comprises a modular power and communication backplane **149** with external and wireless communication headers to enable the removable attachment of the modules **140**, **126**, **128** and interactive communication therebetween.

[0458] In one aspect, the hub modular enclosure **136** includes docking stations, or drawers, **151**, herein also referred to as drawers, which are configured to slidably receive the modules **140**, **126**, **128**. FIG. 4 illustrates a partial

perspective view of a surgical hub enclosure 136, and a combo generator module 145 slidably receivable in a docking station 151 of the surgical hub enclosure 136. A docking port 152 with power and data contacts on a rear side of the combo generator module 145 is configured to engage a corresponding docking port 150 with power and data contacts of a corresponding docking station 151 of the hub modular enclosure 136 as the combo generator module 145 is slid into position within the corresponding docking station 151 of the hub module enclosure 136. In one aspect, the combo generator module 145 includes a bipolar, ultrasonic, and monopolar module and a smoke evacuation module integrated together into a single housing unit 139, as illustrated in FIG. 5.

[0459] In various aspects, the smoke evacuation module 126 includes a fluid line 154 that conveys captured/collected smoke and/or fluid away from a surgical site and to, for example, the smoke evacuation module 126. Vacuum suction originating from the smoke evacuation module 126 can draw the smoke into an opening of a utility conduit at the surgical site. The utility conduit, coupled to the fluid line, can be in the form of a flexible tube terminating at the smoke evacuation module 126. The utility conduit and the fluid line define a fluid path extending toward the smoke evacuation module 126 that is received in the hub enclosure 136.

[0460] In various aspects, the suction/irrigation module 128 is coupled to a surgical tool comprising an aspiration fluid line and a suction fluid line. In one example, the aspiration and suction fluid lines are in the form of flexible tubes extending from the surgical site toward the suction/irrigation module 128. One or more drive systems can be configured to cause irrigation and aspiration of fluids to and from the surgical site.

[0461] In one aspect, the surgical tool includes a shaft having an end effector at a distal end thereof and at least one energy treatment associated with the end effector, an aspiration tube, and an irrigation tube. The aspiration tube can have an inlet port at a distal end thereof and the aspiration tube extends through the shaft. Similarly, an irrigation tube can extend through the shaft and can have an inlet port in proximity to the energy deliver implement. The energy deliver implement is configured to deliver ultrasonic and/or RF energy to the surgical site and is coupled to the generator module 140 by a cable extending initially through the shaft.

[0462] The irrigation tube can be in fluid communication with a fluid source, and the aspiration tube can be in fluid communication with a vacuum source. The fluid source and/or the vacuum source can be housed in the suction/irrigation module 128. In one example, the fluid source and/or the vacuum source can be housed in the hub enclosure 136 separately from the suction/irrigation module 128. In such example, a fluid interface can be configured to connect the suction/irrigation module 128 to the fluid source and/or the vacuum source.

[0463] In one aspect, the modules 140, 126, 128 and/or their corresponding docking stations on the hub modular enclosure 136 may include alignment features that are configured to align the docking ports of the modules into engagement with their counterparts in the docking stations of the hub modular enclosure 136. For example, as illustrated in FIG. 4, the combo generator module 145 includes side brackets 155 that are configured to slidably engage with corresponding brackets 156 of the corresponding docking station 151 of the hub modular enclosure 136. The brackets

cooperate to guide the docking port contacts of the combo generator module 145 into an electrical engagement with the docking port contacts of the hub modular enclosure 136.

[0464] In some aspects, the drawers 151 of the hub modular enclosure 136 are the same, or substantially the same size, and the modules are adjusted in size to be received in the drawers 151. For example, the side brackets 155 and/or 156 can be larger or smaller depending on the size of the module. In other aspects, the drawers 151 are different in size and are each designed to accommodate a particular module.

[0465] Furthermore, the contacts of a particular module can be keyed for engagement with the contacts of a particular drawer to avoid inserting a module into a drawer with mismatching contacts.

[0466] As illustrated in FIG. 4, the docking port 150 of one drawer 151 can be coupled to the docking port 150 of another drawer 151 through a communications link 157 to facilitate an interactive communication between the modules housed in the hub modular enclosure 136. The docking ports 150 of the hub modular enclosure 136 may alternatively, or additionally, facilitate a wireless interactive communication between the modules housed in the hub modular enclosure 136. Any suitable wireless communication can be employed, such as for example Air Titan-Bluetooth.

[0467] FIG. 6 illustrates individual power bus attachments for a plurality of lateral docking ports of a lateral modular housing 160 configured to receive a plurality of modules of a surgical hub 206. The lateral modular housing 160 is configured to laterally receive and interconnect the modules 161. The modules 161 are slidably inserted into docking stations 162 of lateral modular housing 160, which includes a backplane for interconnecting the modules 161. As illustrated in FIG. 6, the modules 161 are arranged laterally in the lateral modular housing 160. Alternatively, the modules 161 may be arranged vertically in a lateral modular housing.

[0468] FIG. 7 illustrates a vertical modular housing 164 configured to receive a plurality of modules 165 of the surgical hub 106. The modules 165 are slidably inserted into docking stations, or drawers, 167 of vertical modular housing 164, which includes a backplane for interconnecting the modules 165. Although the drawers 167 of the vertical modular housing 164 are arranged vertically, in certain instances, a vertical modular housing 164 may include drawers that are arranged laterally. Furthermore, the modules 165 may interact with one another through the docking ports of the vertical modular housing 164. In the example of FIG. 7, a display 177 is provided for displaying data relevant to the operation of the modules 165. In addition, the vertical modular housing 164 includes a master module 178 housing a plurality of sub-modules that are slidably received in the master module 178.

[0469] In various aspects, the imaging module 138 comprises an integrated video processor and a modular light source and is adapted for use with various imaging devices. In one aspect, the imaging device is comprised of a modular housing that can be assembled with a light source module and a camera module. The housing can be a disposable housing. In at least one example, the disposable housing is removably coupled to a reusable controller, a light source module, and a camera module. The light source module and/or the camera module can be selectively chosen depending on the type of surgical procedure. In one aspect, the camera module comprises a CCD sensor. In another aspect,

the camera module comprises a CMOS sensor. In another aspect, the camera module is configured for scanned beam imaging. Likewise, the light source module can be configured to deliver a white light or a different light, depending on the surgical procedure.

[0470] During a surgical procedure, removing a surgical device from the surgical field and replacing it with another surgical device that includes a different camera or a different light source can be inefficient. Temporarily losing sight of the surgical field may lead to undesirable consequences. The module imaging device of the present disclosure is configured to permit the replacement of a light source module or a camera module midstream during a surgical procedure, without having to remove the imaging device from the surgical field.

[0471] In one aspect, the imaging device comprises a tubular housing that includes a plurality of channels. A first channel is configured to slidably receive the camera module, which can be configured for a snap-fit engagement with the first channel. A second channel is configured to slidably receive the light source module, which can be configured for a snap-fit engagement with the second channel. In another example, the camera module and/or the light source module can be rotated into a final position within their respective channels. A threaded engagement can be employed in lieu of the snap-fit engagement.

[0472] In various examples, multiple imaging devices are placed at different positions in the surgical field to provide multiple views. The imaging module 138 can be configured to switch between the imaging devices to provide an optimal view. In various aspects, the imaging module 138 can be configured to integrate the images from the different imaging device.

[0473] Various image processors and imaging devices suitable for use with the present disclosure are described in U.S. Pat. No. 7,995,045, titled COMBINED SBI AND CONVENTIONAL IMAGE PROCESSOR, which issued on Aug. 9, 2011, which is herein incorporated by reference in its entirety. In addition, U.S. Pat. No. 7,982,776, titled SBI MOTION ARTIFACT REMOVAL APPARATUS AND METHOD, which issued on Jul. 19, 2011, which is herein incorporated by reference in its entirety, describes various systems for removing motion artifacts from image data. Such systems can be integrated with the imaging module 138. Furthermore, U.S. Patent Application Publication No. 2011/0306840, titled CONTROLLABLE MAGNETIC SOURCE TO FIXTURE INTRACORPOREAL APPARATUS, which published on Dec. 15, 2011, and U.S. Patent Application Publication No. 2014/0243597, titled SYSTEM FOR PERFORMING A MINIMALLY INVASIVE SURGICAL PROCEDURE, which published on Aug. 28, 2014, each of which is herein incorporated by reference in its entirety.

[0474] FIG. 8 illustrates a surgical data network 201 comprising a modular communication hub 203 configured to connect modular devices located in one or more operating theaters of a healthcare facility, or any room in a healthcare facility specially equipped for surgical operations, to a cloud-based system (e.g., the cloud 204 that may include a remote server 213 coupled to a storage device 205). In one aspect, the modular communication hub 203 comprises a network hub 207 and/or a network switch 209 in communication with a network router. The modular communication hub 203 also can be coupled to a local computer system 210

to provide local computer processing and data manipulation. The surgical data network 201 may be configured as passive, intelligent, or switching. A passive surgical data network serves as a conduit for the data, enabling it to go from one device (or segment) to another and to the cloud computing resources. An intelligent surgical data network includes additional features to enable the traffic passing through the surgical data network to be monitored and to configure each port in the network hub 207 or network switch 209. An intelligent surgical data network may be referred to as a manageable hub or switch. A switching hub reads the destination address of each packet and then forwards the packet to the correct port.

[0475] Modular devices 1a-1n located in the operating theater may be coupled to the modular communication hub 203. The network hub 207 and/or the network switch 209 may be coupled to a network router 211 to connect the devices 1a-1n to the cloud 204 or the local computer system 210. Data associated with the devices 1a-1n may be transferred to cloud-based computers via the router for remote data processing and manipulation. Data associated with the devices 1a-1n may also be transferred to the local computer system 210 for local data processing and manipulation. Modular devices 2a-2m located in the same operating theater also may be coupled to a network switch 209. The network switch 209 may be coupled to the network hub 207 and/or the network router 211 to connect to the devices 2a-2m to the cloud 204. Data associated with the devices 2a-2m may be transferred to the cloud 204 via the network router 211 for data processing and manipulation. Data associated with the devices 2a-2m may also be transferred to the local computer system 210 for local data processing and manipulation.

[0476] It will be appreciated that the surgical data network 201 may be expanded by interconnecting multiple network hubs 207 and/or multiple network switches 209 with multiple network routers 211. The modular communication hub 203 may be contained in a modular control tower configured to receive multiple devices 1a-1n/2a-2m. The local computer system 210 also may be contained in a modular control tower. The modular communication hub 203 is connected to a display 212 to display images obtained by some of the devices 1a-1n/2a-2m, for example during surgical procedures. In various aspects, the devices 1a-1n/2a-2m may include, for example, various modules such as an imaging module 138 coupled to an endoscope, a generator module 140 coupled to an energy-based surgical device, a smoke evacuation module 126, a suction/irrigation module 128, a communication module 130, a processor module 132, a storage array 134, a surgical device coupled to a display, and/or a non-contact sensor module, among other modular devices that may be connected to the modular communication hub 203 of the surgical data network 201.

[0477] In one aspect, the surgical data network 201 may comprise a combination of network hub(s), network switch(es), and network router(s) connecting the devices 1a-1n/2a-2m to the cloud. Any one of or all of the devices 1a-1n/2a-2m coupled to the network hub or network switch may collect data in real time and transfer the data to cloud computers for data processing and manipulation. It will be appreciated that cloud computing relies on sharing computing resources rather than having local servers or personal devices to handle software applications. The word “cloud” may be used as a metaphor for “the Internet,” although the

term is not limited as such. Accordingly, the term “cloud computing” may be used herein to refer to “a type of Internet-based computing,” where different services—such as servers, storage, and applications—are delivered to the modular communication hub **203** and/or computer system **210** located in the surgical theater (e.g., a fixed, mobile, temporary, or field operating room or space) and to devices connected to the modular communication hub **203** and/or computer system **210** through the Internet. The cloud infrastructure may be maintained by a cloud service provider. In this context, the cloud service provider may be the entity that coordinates the usage and control of the devices **1a-1n/2a-2m** located in one or more operating theaters. The cloud computing services can perform a large number of calculations based on the data gathered by smart surgical instruments, robots, and other computerized devices located in the operating theater. The hub hardware enables multiple devices or connections to be connected to a computer that communicates with the cloud computing resources and storage.

[0478] Applying cloud computer data processing techniques on the data collected by the devices **1a-1n/2a-2m**, the surgical data network provides improved surgical outcomes, reduced costs, and improved patient satisfaction. At least some of the devices **1a-1n/2a-2m** may be employed to view tissue states to assess leaks or perfusion of sealed tissue after a tissue sealing and cutting procedure. At least some of the devices **1a-1n/2a-2m** may be employed to identify pathology, such as the effects of diseases, using the cloud-based computing to examine data including images of samples of body tissue for diagnostic purposes. This includes localization and margin confirmation of tissue and phenotypes. At least some of the devices **1a-1n/2a-2m** may be employed to identify anatomical structures of the body using a variety of sensors integrated with imaging devices and techniques such as overlaying images captured by multiple imaging devices. The data gathered by the devices **1a-1n/2a-2m**, including image data, may be transferred to the cloud **204** or the local computer system **210** or both for data processing and manipulation including image processing and manipulation. The data may be analyzed to improve surgical procedure outcomes by determining if further treatment, such as the application of endoscopic intervention, emerging technologies, a targeted radiation, targeted intervention, and precise robotics to tissue-specific sites and conditions, may be pursued. Such data analysis may further employ outcome analytics processing, and using standardized approaches may provide beneficial feedback to either confirm surgical treatments and the behavior of the surgeon or suggest modifications to surgical treatments and the behavior of the surgeon.

[0479] In one implementation, the operating theater devices **1a-1n** may be connected to the modular communication hub **203** over a wired channel or a wireless channel depending on the configuration of the devices **1a-1n** to a network hub. The network hub **207** may be implemented, in one aspect, as a local network broadcast device that works on the physical layer of the Open System Interconnection (OSI) model. The network hub provides connectivity to the devices **1a-1n** located in the same operating theater network. The network hub **207** collects data in the form of packets and sends them to the router in half duplex mode. The network hub **207** does not store any media access control/Internet Protocol (MAC/IP) to transfer the device data. Only one of

the devices **1a-1n** can send data at a time through the network hub **207**. The network hub **207** has no routing tables or intelligence regarding where to send information and broadcasts all network data across each connection and to a remote server **213** (FIG. 9) over the cloud **204**. The network hub **207** can detect basic network errors such as collisions, but having all information broadcast to multiple ports can be a security risk and cause bottlenecks.

[0480] In another implementation, the operating theater devices **2a-2m** may be connected to a network switch **209** over a wired channel or a wireless channel. The network switch **209** works in the data link layer of the OSI model. The network switch **209** is a multicast device for connecting the devices **2a-2m** located in the same operating theater to the network. The network switch **209** sends data in the form of frames to the network router **211** and works in full duplex mode. Multiple devices **2a-2m** can send data at the same time through the network switch **209**. The network switch **209** stores and uses MAC addresses of the devices **2a-2m** to transfer data.

[0481] The network hub **207** and/or the network switch **209** are coupled to the network router **211** for connection to the cloud **204**. The network router **211** works in the network layer of the OSI model. The network router **211** creates a route for transmitting data packets received from the network hub **207** and/or network switch **211** to cloud-based computer resources for further processing and manipulation of the data collected by any one of or all the devices **1a-1n/2a-2m**. The network router **211** may be employed to connect two or more different networks located in different locations, such as, for example, different operating theaters of the same healthcare facility or different networks located in different operating theaters of different healthcare facilities. The network router **211** sends data in the form of packets to the cloud **204** and works in full duplex mode. Multiple devices can send data at the same time. The network router **211** uses IP addresses to transfer data.

[0482] In one example, the network hub **207** may be implemented as a USB hub, which allows multiple USB devices to be connected to a host computer. The USB hub may expand a single USB port into several tiers so that there are more ports available to connect devices to the host system computer. The network hub **207** may include wired or wireless capabilities to receive information over a wired channel or a wireless channel. In one aspect, a wireless USB short-range, high-bandwidth wireless radio communication protocol may be employed for communication between the devices **1a-1n** and devices **2a-2m** located in the operating theater.

[0483] In other examples, the operating theater devices **1a-1n/2a-2m** may communicate to the modular communication hub **203** via Bluetooth wireless technology standard for exchanging data over short distances (using short-wavelength UHF radio waves in the ISM band from 2.4 to 2.485 GHz) from fixed and mobile devices and building personal area networks (PANs). In other aspects, the operating theater devices **1a-1n/2a-2m** may communicate to the modular communication hub **203** via a number of wireless or wired communication standards or protocols, including but not limited to Wi-Fi (IEEE 802.11 family), WiMAX (IEEE 802.16 family), IEEE 802.20, long-term evolution (LTE), and Ev-DO, HSPA+, HSDPA+, HSUPA+, EDGE, GSM, GPRS, CDMA, TDMA, DECT, and Ethernet derivatives thereof, as well as any other wireless and wired protocols

that are designated as 3G, 4G, 5G, and beyond. The computing module may include a plurality of communication modules. For instance, a first communication module may be dedicated to shorter-range wireless communications such as Wi-Fi and Bluetooth, and a second communication module may be dedicated to longer-range wireless communications such as GPS, EDGE, GPRS, CDMA, WiMAX, LTE, Ev-DO, and others.

**[0484]** The modular communication hub **203** may serve as a central connection for one or all of the operating theater devices **1a-1n/2a-2m** and handles a data type known as frames. Frames carry the data generated by the devices **1a-1n/2a-2m**. When a frame is received by the modular communication hub **203**, it is amplified and transmitted to the network router **211**, which transfers the data to the cloud computing resources by using a number of wireless or wired communication standards or protocols, as described herein.

**[0485]** The modular communication hub **203** can be used as a standalone device or be connected to compatible network hubs and network switches to form a larger network. The modular communication hub **203** is generally easy to install, configure, and maintain, making it a good option for networking the operating theater devices **1a-1n/2a-2m**.

**[0486]** FIG. 9 illustrates a computer-implemented interactive surgical system **200**. The computer-implemented interactive surgical system **200** is similar in many respects to the computer-implemented interactive surgical system **100**. For example, the computer-implemented interactive surgical system **200** includes one or more surgical systems **202**, which are similar in many respects to the surgical systems **102**. Each surgical system **202** includes at least one surgical hub **206** in communication with a cloud **204** that may include a remote server **213**. In one aspect, the computer-implemented interactive surgical system **200** comprises a modular control tower **236** connected to multiple operating theater devices such as, for example, intelligent surgical instruments, robots, and other computerized devices located in the operating theater. As shown in FIG. 10, the modular control tower **236** comprises a modular communication hub **203** coupled to a computer system **210**. As illustrated in the example of FIG. 9, the modular control tower **236** is coupled to an imaging module **238** that is coupled to an endoscope **239**, a generator module **240** that is coupled to an energy device **241**, a smoke evacuator module **226**, a suction/irrigation module **228**, a communication module **230**, a processor module **232**, a storage array **234**, a smart device/instrument **235** optionally coupled to a display **237**, and a non-contact sensor module **242**. The operating theater devices are coupled to cloud computing resources and data storage via the modular control tower **236**. A robot hub **222** also may be connected to the modular control tower **236** and to the cloud computing resources. The devices/instruments **235**, visualization systems **208**, among others, may be coupled to the modular control tower **236** via wired or wireless communication standards or protocols, as described herein. The modular control tower **236** may be coupled to a hub display **215** (e.g., monitor, screen) to display and overlay images received from the imaging module, device/instrument display, and/or other visualization systems **208**. The hub display also may display data received from devices connected to the modular control tower in conjunction with images and overlaid images.

**[0487]** FIG. 10 illustrates a surgical hub **206** comprising a plurality of modules coupled to the modular control tower **236**. The modular control tower **236** comprises a modular communication hub **203**, e.g., a network connectivity device, and a computer system **210** to provide local processing, visualization, and imaging, for example. As shown in FIG. 10, the modular communication hub **203** may be connected in a tiered configuration to expand the number of modules (e.g., devices) that may be connected to the modular communication hub **203** and transfer data associated with the modules to the computer system **210**, cloud computing resources, or both. As shown in FIG. 10, each of the network hubs/switches in the modular communication hub **203** includes three downstream ports and one upstream port. The upstream network hub/switch is connected to a processor to provide a communication connection to the cloud computing resources and a local display **217**. Communication to the cloud **204** may be made either through a wired or a wireless communication channel.

**[0488]** The surgical hub **206** employs a non-contact sensor module **242** to measure the dimensions of the operating theater and generate a map of the surgical theater using either ultrasonic or laser-type non-contact measurement devices. An ultrasound-based non-contact sensor module scans the operating theater by transmitting a burst of ultrasound and receiving the echo when it bounces off the perimeter walls of an operating theater as described under the heading “Surgical Hub Spatial Awareness Within an Operating Room” in U.S. Provisional Patent Application Ser. No. 62/611,341, titled INTERACTIVE SURGICAL PLATFORM, filed Dec. 28, 2017, which is herein incorporated by reference in its entirety, in which the sensor module is configured to determine the size of the operating theater and to adjust Bluetooth-pairing distance limits. A laser-based non-contact sensor module scans the operating theater by transmitting laser light pulses, receiving laser light pulses that bounce off the perimeter walls of the operating theater, and comparing the phase of the transmitted pulse to the received pulse to determine the size of the operating theater and to adjust Bluetooth pairing distance limits, for example.

**[0489]** The computer system **210** comprises a processor **244** and a network interface **245**. The processor **244** is coupled to a communication module **247**, storage **248**, memory **249**, non-volatile memory **250**, and input/output interface **251** via a system bus. The system bus can be any of several types of bus structure(s) including the memory bus or memory controller, a peripheral bus or external bus, and/or a local bus using any variety of available bus architectures including, but not limited to, 9-bit bus, Industrial Standard Architecture (ISA), Micro-Channel Architecture (MSA), Extended ISA (EISA), Intelligent Drive Electronics (IDE), VESA Local Bus (VLB), Peripheral Component Interconnect (PCI), USB, Advanced Graphics Port (AGP), Personal Computer Memory Card International Association bus (PCMCIA), Small Computer Systems Interface (SCSI), or any other proprietary bus.

**[0490]** The processor **244** may be any single-core or multicore processor such as those known under the trade name ARM Cortex by Texas Instruments. In one aspect, the processor may be an LM4F230H5QR ARM Cortex-M4F Processor Core, available from Texas Instruments, for example, comprising an on-chip memory of 256 KB single-cycle flash memory, or other non-volatile memory, up to 40 MHz, a prefetch buffer to improve performance above 40

MHz, a 32 KB single-cycle serial random access memory (SRAM), an internal read-only memory (ROM) loaded with StellarisWare® software, a 2 KB electrically erasable programmable read-only memory (EEPROM), and/or one or more pulse width modulation (PWM) modules, one or more quadrature encoder inputs (QEI) analogs, one or more 12-bit analog-to-digital converters (ADCs) with 12 analog input channels, details of which are available for the product datasheet.

**[0491]** In one aspect, the processor **244** may comprise a safety controller comprising two controller-based families such as TMS570 and RM4x, known under the trade name Hercules ARM Cortex R4, also by Texas Instruments. The safety controller may be configured specifically for IEC 61508 and ISO 26262 safety critical applications, among others, to provide advanced integrated safety features while delivering scalable performance, connectivity, and memory options.

**[0492]** The system memory includes volatile memory and non-volatile memory. The basic input/output system (BIOS), containing the basic routines to transfer information between elements within the computer system, such as during start-up, is stored in non-volatile memory. For example, the non-volatile memory can include ROM, programmable ROM (PROM), electrically programmable ROM (EPROM), EEPROM, or flash memory. Volatile memory includes random-access memory (RAM), which acts as external cache memory. Moreover, RAM is available in many forms such as SRAM, dynamic RAM (DRAM), synchronous DRAM (SDRAM), double data rate SDRAM (DDR SDRAM), enhanced SDRAM (ESDRAM), Synchlink DRAM (SL-DRAM), and direct Rambus RAM (DRRAM).

**[0493]** The computer system **210** also includes removable/non-removable, volatile/non-volatile computer storage media, such as for example disk storage. The disk storage includes, but is not limited to, devices like a magnetic disk drive, floppy disk drive, tape drive, Jaz drive, Zip drive, LS-60 drive, flash memory card, or memory stick. In addition, the disk storage can include storage media separately or in combination with other storage media including, but not limited to, an optical disc drive such as a compact disc ROM device (CD-ROM), compact disc recordable drive (CD-R Drive), compact disc rewritable drive (CD-RW Drive), or a digital versatile disc ROM drive (DVD-ROM). To facilitate the connection of the disk storage devices to the system bus, a removable or non-removable interface may be employed.

**[0494]** It is to be appreciated that the computer system **210** includes software that acts as an intermediary between users and the basic computer resources described in a suitable operating environment. Such software includes an operating system. The operating system, which can be stored on the disk storage, acts to control and allocate resources of the computer system. System applications take advantage of the management of resources by the operating system through program modules and program data stored either in the system memory or on the disk storage. It is to be appreciated that various components described herein can be implemented with various operating systems or combinations of operating systems.

**[0495]** A user enters commands or information into the computer system **210** through input device(s) coupled to the I/O interface **251**. The input devices include, but are not limited to, a pointing device such as a mouse, trackball, stylus, touch pad, keyboard, microphone, joystick, game

pad, satellite dish, scanner, TV tuner card, digital camera, digital video camera, web camera, and the like. These and other input devices connect to the processor through the system bus via interface port(s). The interface port(s) include, for example, a serial port, a parallel port, a game port, and a USB. The output device(s) use some of the same types of ports as input device(s). Thus, for example, a USB port may be used to provide input to the computer system and to output information from the computer system to an output device. An output adapter is provided to illustrate that there are some output devices like monitors, displays, speakers, and printers, among other output devices that require special adapters. The output adapters include, by way of illustration and not limitation, video and sound cards that provide a means of connection between the output device and the system bus. It should be noted that other devices and/or systems of devices, such as remote computer(s), provide both input and output capabilities.

**[0496]** The computer system **210** can operate in a networked environment using logical connections to one or more remote computers, such as cloud computer(s), or local computers. The remote cloud computer(s) can be a personal computer, server, router, network PC, workstation, micro-processor-based appliance, peer device, or other common network node, and the like, and typically includes many or all of the elements described relative to the computer system. For purposes of brevity, only a memory storage device is illustrated with the remote computer(s). The remote computer(s) is logically connected to the computer system through a network interface and then physically connected via a communication connection. The network interface encompasses communication networks such as local area networks (LANs) and wide area networks (WANs). LAN technologies include Fiber Distributed Data Interface (FDDI), Copper Distributed Data Interface (CDDI), Ethernet/IEEE 802.3, Token Ring/IEEE 802.5 and the like. WAN technologies include, but are not limited to, point-to-point links, circuit-switching networks like Integrated Services Digital Networks (ISDN) and variations thereon, packet-switching networks, and Digital Subscriber Lines (DSL).

**[0497]** In various aspects, the computer system **210** of FIG. 10, the imaging module **238** and/or visualization system **208**, and/or the processor module **232** of FIGS. 9-10, may comprise an image processor, image-processing engine, media processor, or any specialized digital signal processor (DSP) used for the processing of digital images. The image processor may employ parallel computing with single instruction, multiple data (SIMD) or multiple instruction, multiple data (MIMD) technologies to increase speed and efficiency. The digital image-processing engine can perform a range of tasks. The image processor may be a system on a chip with multicore processor architecture.

**[0498]** The communication connection(s) refers to the hardware/software employed to connect the network interface to the bus. While the communication connection is shown for illustrative clarity inside the computer system, it can also be external to the computer system **210**. The hardware/software necessary for connection to the network interface includes, for illustrative purposes only, internal and external technologies such as modems, including regular telephone-grade modems, cable modems, and DSL modems, ISDN adapters, and Ethernet cards.

[0499] FIG. 11 illustrates a functional block diagram of one aspect of a USB network hub 300 device, according to one aspect of the present disclosure. In the illustrated aspect, the USB network hub device 300 employs a TUSB2036 integrated circuit hub by Texas Instruments. The USB network hub 300 is a CMOS device that provides an upstream USB transceiver port 302 and up to three downstream USB transceiver ports 304, 306, 308 in compliance with the USB 2.0 specification. The upstream USB transceiver port 302 is a differential root data port comprising a differential data minus (DM0) input paired with a differential data plus (DP0) input. The three downstream USB transceiver ports 304, 306, 308 are differential data ports where each port includes differential data plus (DP1-DP3) outputs paired with differential data minus (DM1-DM3) outputs.

[0500] The USB network hub 300 device is implemented with a digital state machine instead of a microcontroller, and no firmware programming is required. Fully compliant USB transceivers are integrated into the circuit for the upstream USB transceiver port 302 and all downstream USB transceiver ports 304, 306, 308. The downstream USB transceiver ports 304, 306, 308 support both full-speed and low-speed devices by automatically setting the slew rate according to the speed of the device attached to the ports. The USB network hub 300 device may be configured either in bus-powered or self-powered mode and includes a hub power logic 312 to manage power.

[0501] The USB network hub 300 device includes a serial interface engine 310 (SIE). The SIE 310 is the front end of the USB network hub 300 hardware and handles most of the protocol described in chapter 8 of the USB specification. The SIE 310 typically comprehends signaling up to the transaction level. The functions that it handles could include: packet recognition, transaction sequencing, SOP, EOP, RESET, and RESUME signal detection/generation, clock/data separation, non-return-to-zero invert (NRZI) data encoding/decoding and bit-stuffing, CRC generation and checking (token and data), packet ID (PID) generation and checking/decoding, and/or serial-parallel/parallel-serial conversion. The 310 receives a clock input 314 and is coupled to a suspend/resume logic and frame timer 316 circuit and a hub repeater circuit 318 to control communication between the upstream USB transceiver port 302 and the downstream USB transceiver ports 304, 306, 308 through port logic circuits 320, 322, 324. The SIE 310 is coupled to a command decoder 326 via interface logic to control commands from a serial EEPROM via a serial EEPROM interface 330.

[0502] In various aspects, the USB network hub 300 can connect 127 functions configured in up to six logical layers (tiers) to a single computer. Further, the USB network hub 300 can connect to all peripherals using a standardized four-wire cable that provides both communication and power distribution. The power configurations are bus-powered and self-powered modes. The USB network hub 300 may be configured to support four modes of power management: a bus-powered hub, with either individual-port power management or ganged-port power management, and the self-powered hub, with either individual-port power management or ganged-port power management. In one aspect, using a USB cable, the USB network hub 300, the upstream USB transceiver port 302 is plugged into a USB host controller, and the downstream USB transceiver ports 304, 306, 308 are exposed for connecting USB compatible devices, and so forth.

#### Surgical Instrument Hardware

[0503] FIG. 12 illustrates a logic diagram of a control system 470 of a surgical instrument or tool in accordance with one or more aspects of the present disclosure. The system 470 comprises a control circuit. The control circuit includes a microcontroller 461 comprising a processor 462 and a memory 468. One or more of sensors 472, 474, 476, for example, provide real-time feedback to the processor 462. A motor 482, driven by a motor driver 492, operably couples a longitudinally movable displacement member to drive the I-beam knife element. A tracking system 480 is configured to determine the position of the longitudinally movable displacement member. The position information is provided to the processor 462, which can be programmed or configured to determine the position of the longitudinally movable drive member as well as the position of a firing member, firing bar, and I-beam knife element. Additional motors may be provided at the tool driver interface to control I-beam firing, closure tube travel, shaft rotation, and articulation. A display 473 displays a variety of operating conditions of the instruments and may include touch screen functionality for data input. Information displayed on the display 473 may be overlaid with images acquired via endoscopic imaging modules.

[0504] In one aspect, the microcontroller 461 may be any single-core or multicore processor such as those known under the trade name ARM Cortex by Texas Instruments. In one aspect, the main microcontroller 461 may be an LM4F230H5QR ARM Cortex-M4F Processor Core, available from Texas Instruments, for example, comprising an on-chip memory of 256 KB single-cycle flash memory, or other non-volatile memory, up to 40 MHz, a prefetch buffer to improve performance above 40 MHz, a 32 KB single-cycle SRAM, and internal ROM loaded with StellarisWare® software, a 2 KB EEPROM, one or more PWM modules, one or more QEI analogs, and/or one or more 12-bit ADCs with 12 analog input channels, details of which are available for the product datasheet.

[0505] In one aspect, the microcontroller 461 may comprise a safety controller comprising two controller-based families such as TMS570 and RM4x, known under the trade name Hercules ARM Cortex R4, also by Texas Instruments. The safety controller may be configured specifically for IEC 61508 and ISO 26262 safety critical applications, among others, to provide advanced integrated safety features while delivering scalable performance, connectivity, and memory options.

[0506] The microcontroller 461 may be programmed to perform various functions such as precise control over the speed and position of the knife and articulation systems. In one aspect, the microcontroller 461 includes a processor 462 and a memory 468. The electric motor 482 may be a brushed direct current (DC) motor with a gearbox and mechanical links to an articulation or knife system. In one aspect, a motor driver 492 may be an A3941 available from Allegro Microsystems, Inc. Other motor drivers may be readily substituted for use in the tracking system 480 comprising an absolute positioning system. A detailed description of an absolute positioning system is described in U.S. Patent Application Publication No. 2017/0296213, titled SYSTEMS AND METHODS FOR CONTROLLING A SURGICAL STAPLING AND CUTTING INSTRUMENT, which published on Oct. 19, 2017, which is herein incorporated by reference in its entirety.



[0507] The microcontroller 461 may be programmed to provide precise control over the speed and position of displacement members and articulation systems. The microcontroller 461 may be configured to compute a response in the software of the microcontroller 461. The computed response is compared to a measured response of the actual system to obtain an “observed” response, which is used for actual feedback decisions. The observed response is a favorable, tuned value that balances the smooth, continuous nature of the simulated response with the measured response, which can detect outside influences on the system.

[0508] In one aspect, the motor 482 may be controlled by the motor driver 492 and can be employed by the firing system of the surgical instrument or tool. In various forms, the motor 482 may be a brushed DC driving motor having a maximum rotational speed of approximately 25,000 RPM. In other arrangements, the motor 482 may include a brushless motor, a cordless motor, a synchronous motor, a stepper motor, or any other suitable electric motor. The motor driver 492 may comprise an H-bridge driver comprising field-effect transistors (FETs), for example. The motor 482 can be powered by a power assembly releasably mounted to the handle assembly or tool housing for supplying control power to the surgical instrument or tool. The power assembly may comprise a battery which may include a number of battery cells connected in series that can be used as the power source to power the surgical instrument or tool. In certain circumstances, the battery cells of the power assembly may be replaceable and/or rechargeable. In at least one example, the battery cells can be lithium-ion batteries which can be coupleable to and separable from the power assembly.

[0509] The motor driver 492 may be an A3941 available from Allegro Microsystems, Inc. The A3941 492 is a full-bridge controller for use with external N-channel power metal-oxide semiconductor field-effect transistors (MOSFETs) specifically designed for inductive loads, such as brush DC motors. The driver 492 comprises a unique charge pump regulator that provides full (>10 V) gate drive for battery voltages down to 7 V and allows the A3941 to operate with a reduced gate drive, down to 5.5 V. A bootstrap capacitor may be employed to provide the above battery supply voltage required for N-channel MOSFETs. An internal charge pump for the high-side drive allows DC (100% duty cycle) operation. The full bridge can be driven in fast or slow decay modes using diode or synchronous rectification. In the slow decay mode, current recirculation can be through the high-side or the lowside FETs. The power FETs are protected from shoot-through by resistor-adjustable dead time. Integrated diagnostics provide indications of under-voltage, overtemperature, and power bridge faults and can be configured to protect the power MOSFETs under most short circuit conditions. Other motor drivers may be readily substituted for use in the tracking system 480 comprising an absolute positioning system.

[0510] The tracking system 480 comprises a controlled motor drive circuit arrangement comprising a position sensor 472 according to one aspect of this disclosure. The position sensor 472 for an absolute positioning system provides a unique position signal corresponding to the location of a displacement member. In one aspect, the displacement member represents a longitudinally movable drive member comprising a rack of drive teeth for meshing engagement with a corresponding drive gear of a gear reducer assembly. In other aspects, the displacement mem-

ber represents the firing member, which could be adapted and configured to include a rack of drive teeth. In yet another aspect, the displacement member represents a firing bar or the I-beam, each of which can be adapted and configured to include a rack of drive teeth. Accordingly, as used herein, the term displacement member is used generically to refer to any movable member of the surgical instrument or tool such as the drive member, the firing member, the firing bar, the I-beam, or any element that can be displaced. In one aspect, the longitudinally movable drive member is coupled to the firing member, the firing bar, and the I-beam. Accordingly, the absolute positioning system can, in effect, track the linear displacement of the I-beam by tracking the linear displacement of the longitudinally movable drive member. In various other aspects, the displacement member may be coupled to any position sensor 472 suitable for measuring linear displacement. Thus, the longitudinally movable drive member, the firing member, the firing bar, or the I-beam, or combinations thereof, may be coupled to any suitable linear displacement sensor. Linear displacement sensors may include contact or non-contact displacement sensors. Linear displacement sensors may comprise linear variable differential transformers (LVDT), differential variable reluctance transducers (DVRT), a slide potentiometer, a magnetic sensing system comprising a movable magnet and a series of linearly arranged Hall effect sensors, a magnetic sensing system comprising a fixed magnet and a series of movable, linearly arranged Hall effect sensors, an optical sensing system comprising a movable light source and a series of linearly arranged photo diodes or photo detectors, an optical sensing system comprising a fixed light source and a series of movable linearly, arranged photo diodes or photo detectors, or any combination thereof.

[0511] The electric motor 482 can include a rotatable shaft that operably interfaces with a gear assembly that is mounted in meshing engagement with a set, or rack, of drive teeth on the displacement member. A sensor element may be operably coupled to a gear assembly such that a single revolution of the position sensor 472 element corresponds to some linear longitudinal translation of the displacement member. An arrangement of gearing and sensors can be connected to the linear actuator, via a rack and pinion arrangement, or a rotary actuator, via a spur gear or other connection. A power source supplies power to the absolute positioning system and an output indicator may display the output of the absolute positioning system. The displacement member represents the longitudinally movable drive member comprising a rack of drive teeth formed thereon for meshing engagement with a corresponding drive gear of the gear reducer assembly. The displacement member represents the longitudinally movable firing member, firing bar, I-beam, or combinations thereof.

[0512] A single revolution of the sensor element associated with the position sensor 472 is equivalent to a longitudinal linear displacement  $d1$  of the of the displacement member, where  $d1$  is the longitudinal linear distance that the displacement member moves from point “a” to point “b” after a single revolution of the sensor element coupled to the displacement member. The sensor arrangement may be connected via a gear reduction that results in the position sensor 472 completing one or more revolutions for the full stroke of the displacement member. The position sensor 472 may complete multiple revolutions for the full stroke of the displacement member.

[0513] A series of switches, where  $n$  is an integer greater than one, may be employed alone or in combination with a gear reduction to provide a unique position signal for more than one revolution of the position sensor 472. The state of the switches are fed back to the microcontroller 461 that applies logic to determine a unique position signal corresponding to the longitudinal linear displacement  $d_1+d_2+ \dots +d_n$  of the displacement member. The output of the position sensor 472 is provided to the microcontroller 461. The position sensor 472 of the sensor arrangement may comprise a magnetic sensor, an analog rotary sensor like a potentiometer, or an array of analog Hall-effect elements, which output a unique combination of position signals or values.

[0514] The position sensor 472 may comprise any number of magnetic sensing elements, such as, for example, magnetic sensors classified according to whether they measure the total magnetic field or the vector components of the magnetic field. The techniques used to produce both types of magnetic sensors encompass many aspects of physics and electronics. The technologies used for magnetic field sensing include search coil, fluxgate, optically pumped, nuclear precession, SQUID, Hall-effect, anisotropic magnetoresistance, giant magnetoresistance, magnetic tunnel junctions, giant magnetoimpedance, magnetostrictive/piezoelectric composites, magnetodiode, magnetotransistor, fiber-optic, magneto-optic, and microelectromechanical systems-based magnetic sensors, among others.

[0515] In one aspect, the position sensor 472 for the tracking system 480 comprising an absolute positioning system comprises a magnetic rotary absolute positioning system. The position sensor 472 may be implemented as an AS5055EQFT single-chip magnetic rotary position sensor available from Austria Microsystems, AG. The position sensor 472 is interfaced with the microcontroller 461 to provide an absolute positioning system. The position sensor 472 is a low-voltage and low-power component and includes four Hall-effect elements in an area of the position sensor 472 that is located above a magnet. A high-resolution ADC and a smart power management controller are also provided on the chip. A coordinate rotation digital computer (CORDIC) processor, also known as the digit-by-digit method and Volder's algorithm, is provided to implement a simple and efficient algorithm to calculate hyperbolic and trigonometric functions that require only addition, subtraction, bitshift, and table lookup operations. The angle position, alarm bits, and magnetic field information are transmitted over a standard serial communication interface, such as a serial peripheral interface (SPI) interface, to the microcontroller 461. The position sensor 472 provides 12 or 14 bits of resolution. The position sensor 472 may be an AS5055 chip provided in a small QFN 16-pin  $4 \times 4 \times 0.85$  mm package.

[0516] The tracking system 480 comprising an absolute positioning system may comprise and/or be programmed to implement a feedback controller, such as a PID, state feedback, and adaptive controller. A power source converts the signal from the feedback controller into a physical input to the system: in this case the voltage. Other examples include a PWM of the voltage, current, and force. Other sensor(s) may be provided to measure physical parameters of the physical system in addition to the position measured by the position sensor 472. In some aspects, the other sensor(s) can include sensor arrangements such as those described in U.S. Pat. No. 9,345,481, titled STAPLE CAR-

TRIDGE TISSUE THICKNESS SENSOR SYSTEM, which issued on May 24, 2016, which is herein incorporated by reference in its entirety; U.S. Patent Application Publication No. 2014/0263552, titled STAPLE CARTRIDGE TISSUE THICKNESS SENSOR SYSTEM, which published on Sep. 18, 2014, which is herein incorporated by reference in its entirety; and U.S. patent application Ser. No. 15/628,175, titled TECHNIQUES FOR ADAPTIVE CONTROL OF MOTOR VELOCITY OF A SURGICAL STAPLING AND CUTTING INSTRUMENT, filed Jun. 20, 2017, which is herein incorporated by reference in its entirety. In a digital signal processing system, an absolute positioning system is coupled to a digital data acquisition system where the output of the absolute positioning system will have a finite resolution and sampling frequency. The absolute positioning system may comprise a compare-and-combine circuit to combine a computed response with a measured response using algorithms, such as a weighted average and a theoretical control loop, that drive the computed response towards the measured response. The computed response of the physical system takes into account properties like mass, inertial, viscous friction, inductance resistance, etc., to predict what the states and outputs of the physical system will be by knowing the input.

[0517] The absolute positioning system provides an absolute position of the displacement member upon power-up of the instrument, without retracting or advancing the displacement member to a reset (zero or home) position as may be required with conventional rotary encoders that merely count the number of steps forwards or backwards that the motor 482 has taken to infer the position of a device actuator, drive bar, knife, or the like.

[0518] A sensor 474, such as, for example, a strain gauge or a micro-strain gauge, is configured to measure one or more parameters of the end effector, such as, for example, the amplitude of the strain exerted on the anvil during a clamping operation, which can be indicative of the closure forces applied to the anvil. The measured strain is converted to a digital signal and provided to the processor 462. Alternatively, or in addition to the sensor 474, a sensor 476, such as, for example, a load sensor, can measure the closure force applied by the closure drive system to the anvil. The sensor 476, such as, for example, a load sensor, can measure the firing force applied to an I-beam in a firing stroke of the surgical instrument or tool. The I-beam is configured to engage a wedge sled, which is configured to upwardly cam staple drivers to force out staples into deforming contact with an anvil. The I-beam also includes a sharpened cutting edge that can be used to sever tissue as the I-beam is advanced distally by the firing bar. Alternatively, a current sensor 478 can be employed to measure the current drawn by the motor 482. The force required to advance the firing member can correspond to the current drawn by the motor 482, for example. The measured force is converted to a digital signal and provided to the processor 462.

[0519] In one form, the strain gauge sensor 474 can be used to measure the force applied to the tissue by the end effector. A strain gauge can be coupled to the end effector to measure the force on the tissue being treated by the end effector. A system for measuring forces applied to the tissue grasped by the end effector comprises a strain gauge sensor 474, such as, for example, a micro-strain gauge, that is configured to measure one or more parameters of the end effector, for example. In one aspect, the strain gauge sensor

**474** can measure the amplitude or magnitude of the strain exerted on a jaw member of an end effector during a clamping operation, which can be indicative of the tissue compression. The measured strain is converted to a digital signal and provided to a processor **462** of the microcontroller **461**. A load sensor **476** can measure the force used to operate the knife element, for example, to cut the tissue captured between the anvil and the staple cartridge. A magnetic field sensor can be employed to measure the thickness of the captured tissue. The measurement of the magnetic field sensor also may be converted to a digital signal and provided to the processor **462**.

[**0520**] The measurements of the tissue compression, the tissue thickness, and/or the force required to close the end effector on the tissue, as respectively measured by the sensors **474**, **476**, can be used by the microcontroller **461** to characterize the selected position of the firing member and/or the corresponding value of the speed of the firing member. In one instance, a memory **468** may store a technique, an equation, and/or a lookup table which can be employed by the microcontroller **461** in the assessment.

[**0521**] The control system **470** of the surgical instrument or tool also may comprise wired or wireless communication circuits to communicate with the modular communication hub as shown in FIGS. **8-11**.

[**0522**] FIG. **13** illustrates a control circuit **500** configured to control aspects of the surgical instrument or tool according to one aspect of this disclosure. The control circuit **500** can be configured to implement various processes described herein. The control circuit **500** may comprise a microcontroller comprising one or more processors **502** (e.g., microprocessor, microcontroller) coupled to at least one memory circuit **504**. The memory circuit **504** stores machine-executable instructions that, when executed by the processor **502**, cause the processor **502** to execute machine instructions to implement various processes described herein. The processor **502** may be any one of a number of single-core or multicore processors known in the art. The memory circuit **504** may comprise volatile and non-volatile storage media. The processor **502** may include an instruction processing unit **506** and an arithmetic unit **508**. The instruction processing unit may be configured to receive instructions from the memory circuit **504** of this disclosure.

[**0523**] FIG. **14** illustrates a combinational logic circuit **510** configured to control aspects of the surgical instrument or tool according to one aspect of this disclosure. The combinational logic circuit **510** can be configured to implement various processes described herein. The combinational logic circuit **510** may comprise a finite state machine comprising a combinational logic **512** configured to receive data associated with the surgical instrument or tool at an input **514**, process the data by the combinational logic **512**, and provide an output **516**.

[**0524**] FIG. **15** illustrates a sequential logic circuit **520** configured to control aspects of the surgical instrument or tool according to one aspect of this disclosure. The sequential logic circuit **520** or the combinational logic **522** can be configured to implement various processes described herein. The sequential logic circuit **520** may comprise a finite state machine. The sequential logic circuit **520** may comprise a combinational logic **522**, at least one memory circuit **524**, and a clock **529**, for example. The at least one memory circuit **524** can store a current state of the finite state machine. In certain instances, the sequential logic circuit

**520** may be synchronous or asynchronous. The combinational logic **522** is configured to receive data associated with the surgical instrument or tool from an input **526**, process the data by the combinational logic **522**, and provide an output **528**. In other aspects, the circuit may comprise a combination of a processor (e.g., processor **502**, FIG. **13**) and a finite state machine to implement various processes herein. In other aspects, the finite state machine may comprise a combination of a combinational logic circuit (e.g., combinational logic circuit **510**, FIG. **14**) and the sequential logic circuit **520**.

[**0525**] FIG. **16** illustrates a surgical instrument or tool comprising a plurality of motors which can be activated to perform various functions. In certain instances, a first motor can be activated to perform a first function, a second motor can be activated to perform a second function, a third motor can be activated to perform a third function, a fourth motor can be activated to perform a fourth function, and so on. In certain instances, the plurality of motors of robotic surgical instrument **600** can be individually activated to cause firing, closure, and/or articulation motions in the end effector. The firing, closure, and/or articulation motions can be transmitted to the end effector through a shaft assembly, for example.

[**0526**] In certain instances, the surgical instrument system or tool may include a firing motor **602**. The firing motor **602** may be operably coupled to a firing motor drive assembly **604** which can be configured to transmit firing motions, generated by the motor **602** to the end effector, in particular to displace the I-beam element. In certain instances, the firing motions generated by the motor **602** may cause the staples to be deployed from the staple cartridge into tissue captured by the end effector and/or the cutting edge of the I-beam element to be advanced to cut the captured tissue, for example. The I-beam element may be retracted by reversing the direction of the motor **602**.

[**0527**] In certain instances, the surgical instrument or tool may include a closure motor **603**. The closure motor **603** may be operably coupled to a closure motor drive assembly **605** which can be configured to transmit closure motions, generated by the motor **603** to the end effector, in particular to displace a closure tube to close the anvil and compress tissue between the anvil and the staple cartridge. The closure motions may cause the end effector to transition from an open configuration to an approximated configuration to capture tissue, for example. The end effector may be transitioned to an open position by reversing the direction of the motor **603**.

[**0528**] In certain instances, the surgical instrument or tool may include one or more articulation motors **606a**, **606b**, for example. The motors **606a**, **606b** may be operably coupled to respective articulation motor drive assemblies **608a**, **608b**, which can be configured to transmit articulation motions generated by the motors **606a**, **606b** to the end effector. In certain instances, the articulation motions may cause the end effector to articulate relative to the shaft, for example.

[**0529**] As described above, the surgical instrument or tool may include a plurality of motors which may be configured to perform various independent functions. In certain instances, the plurality of motors of the surgical instrument or tool can be individually or separately activated to perform one or more functions while the other motors remain inactive. For example, the articulation motors **606a**, **606b** can be activated to cause the end effector to be articulated while the

firing motor **602** remains inactive. Alternatively, the firing motor **602** can be activated to fire the plurality of staples, and/or to advance the cutting edge, while the articulation motor **606** remains inactive. Furthermore, the closure motor **603** may be activated simultaneously with the firing motor **602** to cause the closure tube and the I-beam element to advance distally as described in more detail hereinbelow.

[0530] In certain instances, the surgical instrument or tool may include a common control module **610** which can be employed with a plurality of motors of the surgical instrument or tool. In certain instances, the common control module **610** may accommodate one of the plurality of motors at a time. For example, the common control module **610** can be couplable to and separable from the plurality of motors of the robotic surgical instrument individually. In certain instances, a plurality of the motors of the surgical instrument or tool may share one or more common control modules such as the common control module **610**. In certain instances, a plurality of motors of the surgical instrument or tool can be individually and selectively engaged with the common control module **610**. In certain instances, the common control module **610** can be selectively switched from interfacing with one of a plurality of motors of the surgical instrument or tool to interfacing with another one of the plurality of motors of the surgical instrument or tool.

[0531] In at least one example, the common control module **610** can be selectively switched between operable engagement with the articulation motors **606a**, **606b** and operable engagement with either the firing motor **602** or the closure motor **603**. In at least one example, as illustrated in FIG. 16, a switch **614** can be moved or transitioned between a plurality of positions and/or states. In a first position **616**, the switch **614** may electrically couple the common control module **610** to the firing motor **602**; in a second position **617**, the switch **614** may electrically couple the common control module **610** to the closure motor **603**; in a third position **618a**, the switch **614** may electrically couple the common control module **610** to the first articulation motor **606a**; and in a fourth position **618b**, the switch **614** may electrically couple the common control module **610** to the second articulation motor **606b**, for example. In certain instances, separate common control modules **610** can be electrically coupled to the firing motor **602**, the closure motor **603**, and the articulations motor **606a**, **606b** at the same time. In certain instances, the switch **614** may be a mechanical switch, an electromechanical switch, a solid-state switch, or any suitable switching mechanism.

[0532] Each of the motors **602**, **603**, **606a**, **606b** may comprise a torque sensor to measure the output torque on the shaft of the motor. The force on an end effector may be sensed in any conventional manner, such as by force sensors on the outer sides of the jaws or by a torque sensor for the motor actuating the jaws.

[0533] In various instances, as illustrated in FIG. 16, the common control module **610** may comprise a motor driver **626** which may comprise one or more H-Bridge FETs. The motor driver **626** may modulate the power transmitted from a power source **628** to a motor coupled to the common control module **610** based on input from a microcontroller **620** (the “controller”), for example. In certain instances, the microcontroller **620** can be employed to determine the current drawn by the motor, for example, while the motor is coupled to the common control module **610**, as described above.

[0534] In certain instances, the microcontroller **620** may include a microprocessor **622** (the “processor”) and one or more non-transitory computer-readable mediums or memory units **624** (the “memory”). In certain instances, the memory **624** may store various program instructions, which when executed may cause the processor **622** to perform a plurality of functions and/or calculations described herein. In certain instances, one or more of the memory units **624** may be coupled to the processor **622**, for example.

[0535] In certain instances, the power source **628** can be employed to supply power to the microcontroller **620**, for example. In certain instances, the power source **628** may comprise a battery (or “battery pack” or “power pack”), such as a lithium-ion battery, for example. In certain instances, the battery pack may be configured to be releasably mounted to a handle for supplying power to the surgical instrument **600**. A number of battery cells connected in series may be used as the power source **628**. In certain instances, the power source **628** may be replaceable and/or rechargeable, for example.

[0536] In various instances, the processor **622** may control the motor driver **626** to control the position, direction of rotation, and/or velocity of a motor that is coupled to the common control module **610**. In certain instances, the processor **622** can signal the motor driver **626** to stop and/or disable a motor that is coupled to the common control module **610**. It should be understood that the term “processor” as used herein includes any suitable microprocessor, microcontroller, or other basic computing device that incorporates the functions of a computer’s central processing unit (CPU) on an integrated circuit or, at most, a few integrated circuits. The processor is a multipurpose, programmable device that accepts digital data as input, processes it according to instructions stored in its memory, and provides results as output. It is an example of sequential digital logic, as it has internal memory. Processors operate on numbers and symbols represented in the binary numeral system.

[0537] In one instance, the processor **622** may be any single-core or multicore processor such as those known under the trade name ARM Cortex by Texas Instruments. In certain instances, the microcontroller **620** may be an LM4F230H5QR, available from Texas Instruments, for example. In at least one example, the Texas Instruments LM4F230H5QR is an ARM Cortex-M4F Processor Core comprising an on-chip memory of 256 KB single-cycle flash memory, or other non-volatile memory, up to 40 MHz, a prefetch buffer to improve performance above 40 MHz, a 32 KB single-cycle SRAM, an internal ROM loaded with StellarisWare® software, a 2 KB EEPROM, one or more PWM modules, one or more QEI analogs, one or more 12-bit ADCs with 12 analog input channels, among other features that are readily available for the product datasheet. Other microcontrollers may be readily substituted for use with the module **4410**. Accordingly, the present disclosure should not be limited in this context.

[0538] In certain instances, the memory **624** may include program instructions for controlling each of the motors of the surgical instrument **600** that are couplable to the common control module **610**. For example, the memory **624** may include program instructions for controlling the firing motor **602**, the closure motor **603**, and the articulation motors **606a**, **606b**. Such program instructions may cause the processor **622** to control the firing, closure, and articulation

functions in accordance with inputs from algorithms or control programs of the surgical instrument or tool.

[0539] In certain instances, one or more mechanisms and/or sensors such as, for example, sensors 630 can be employed to alert the processor 622 to the program instructions that should be used in a particular setting. For example, the sensors 630 may alert the processor 622 to use the program instructions associated with firing, closing, and articulating the end effector. In certain instances, the sensors 630 may comprise position sensors which can be employed to sense the position of the switch 614, for example. Accordingly, the processor 622 may use the program instructions associated with firing the I-beam of the end effector upon detecting, through the sensors 630 for example, that the switch 614 is in the first position 616; the processor 622 may use the program instructions associated with closing the anvil upon detecting, through the sensors 630 for example, that the switch 614 is in the second position 617; and the processor 622 may use the program instructions associated with articulating the end effector upon detecting, through the sensors 630 for example, that the switch 614 is in the third or fourth position 618a, 618b.

[0540] FIG. 17 is a schematic diagram of a robotic surgical instrument 700 configured to operate a surgical tool described herein according to one aspect of this disclosure. The robotic surgical instrument 700 may be programmed or configured to control distal/proximal translation of a displacement member, distal/proximal displacement of a closure tube, shaft rotation, and articulation, either with single or multiple articulation drive links. In one aspect, the surgical instrument 700 may be programmed or configured to individually control a firing member, a closure member, a shaft member, and/or one or more articulation members. The surgical instrument 700 comprises a control circuit 710 configured to control motor-driven firing members, closure members, shaft members, and/or one or more articulation members.

[0541] In one aspect, the robotic surgical instrument 700 comprises a control circuit 710 configured to control an anvil 716 and an I-beam 714 (including a sharp cutting edge) portion of an end effector 702, a removable staple cartridge 718, a shaft 740, and one or more articulation members 742a, 742b via a plurality of motors 704a-704e. A position sensor 734 may be configured to provide position feedback of the I-beam 714 to the control circuit 710. Other sensors 738 may be configured to provide feedback to the control circuit 710. A timer/counter 731 provides timing and counting information to the control circuit 710. An energy source 712 may be provided to operate the motors 704a-704e, and a current sensor 736 provides motor current feedback to the control circuit 710. The motors 704a-704e can be operated individually by the control circuit 710 in an open-loop or closed-loop feedback control.

[0542] In one aspect, the control circuit 710 may comprise one or more microcontrollers, microprocessors, or other suitable processors for executing instructions that cause the processor or processors to perform one or more tasks. In one aspect, a timer/counter 731 provides an output signal, such as the elapsed time or a digital count, to the control circuit 710 to correlate the position of the I-beam 714 as determined by the position sensor 734 with the output of the timer/counter 731 such that the control circuit 710 can determine the position of the I-beam 714 at a specific time (t) relative to a starting position or the time (t) when the I-beam 714 is

at a specific position relative to a starting position. The timer/counter 731 may be configured to measure elapsed time, count external events, or time external events.

[0543] In one aspect, the control circuit 710 may be programmed to control functions of the end effector 702 based on one or more tissue conditions. The control circuit 710 may be programmed to sense tissue conditions, such as thickness, either directly or indirectly, as described herein. The control circuit 710 may be programmed to select a firing control program or closure control program based on tissue conditions. A firing control program may describe the distal motion of the displacement member. Different firing control programs may be selected to better treat different tissue conditions. For example, when thicker tissue is present, the control circuit 710 may be programmed to translate the displacement member at a lower velocity and/or with lower power. When thinner tissue is present, the control circuit 710 may be programmed to translate the displacement member at a higher velocity and/or with higher power. A closure control program may control the closure force applied to the tissue by the anvil 716. Other control programs control the rotation of the shaft 740 and the articulation members 742a, 742b.

[0544] In one aspect, the control circuit 710 may generate motor set point signals. The motor set point signals may be provided to various motor controllers 708a-708e. The motor controllers 708a-708e may comprise one or more circuits configured to provide motor drive signals to the motors 704a-704e to drive the motors 704a-704e as described herein. In some examples, the motors 704a-704e may be brushed DC electric motors. For example, the velocity of the motors 704a-704e may be proportional to the respective motor drive signals. In some examples, the motors 704a-704e may be brushless DC electric motors, and the respective motor drive signals may comprise a PWM signal provided to one or more stator windings of the motors 704a-704e. Also, in some examples, the motor controllers 708a-708e may be omitted and the control circuit 710 may generate the motor drive signals directly.

[0545] In one aspect, the control circuit 710 may initially operate each of the motors 704a-704e in an open-loop configuration for a first open-loop portion of a stroke of the displacement member. Based on the response of the robotic surgical instrument 700 during the open-loop portion of the stroke, the control circuit 710 may select a firing control program in a closed-loop configuration. The response of the instrument may include a translation distance of the displacement member during the open-loop portion, a time elapsed during the open-loop portion, the energy provided to one of the motors 704a-704e during the open-loop portion, a sum of pulse widths of a motor drive signal, etc. After the open-loop portion, the control circuit 710 may implement the selected firing control program for a second portion of the displacement member stroke. For example, during a closed-loop portion of the stroke, the control circuit 710 may modulate one of the motors 704a-704e based on translation data describing a position of the displacement member in a closed-loop manner to translate the displacement member at a constant velocity.

[0546] In one aspect, the motors 704a-704e may receive power from an energy source 712. The energy source 712 may be a DC power supply driven by a main alternating current power source, a battery, a super capacitor, or any other suitable energy source. The motors 704a-704e may be

mechanically coupled to individual movable mechanical elements such as the I-beam 714, anvil 716, shaft 740, articulation 742a, and articulation 742b via respective transmissions 706a-706e. The transmissions 706a-706e may include one or more gears or other linkage components to couple the motors 704a-704e to movable mechanical elements. A position sensor 734 may sense a position of the I-beam 714. The position sensor 734 may be or include any type of sensor that is capable of generating position data that indicate a position of the I-beam 714. In some examples, the position sensor 734 may include an encoder configured to provide a series of pulses to the control circuit 710 as the I-beam 714 translates distally and proximally. The control circuit 710 may track the pulses to determine the position of the I-beam 714. Other suitable position sensors may be used, including, for example, a proximity sensor. Other types of position sensors may provide other signals indicating motion of the I-beam 714. Also, in some examples, the position sensor 734 may be omitted. Where any of the motors 704a-704e is a stepper motor, the control circuit 710 may track the position of the I-beam 714 by aggregating the number and direction of steps that the motor 704 has been instructed to execute. The position sensor 734 may be located in the end effector 702 or at any other portion of the instrument. The outputs of each of the motors 704a-704e include a torque sensor 744a-744e to sense force and have an encoder to sense rotation of the drive shaft.

[0547] In one aspect, the control circuit 710 is configured to drive a firing member such as the I-beam 714 portion of the end effector 702. The control circuit 710 provides a motor set point to a motor control 708a, which provides a drive signal to the motor 704a. The output shaft of the motor 704a is coupled to a torque sensor 744a. The torque sensor 744a is coupled to a transmission 706a which is coupled to the I-beam 714. The transmission 706a comprises movable mechanical elements such as rotating elements and a firing member to control the movement of the I-beam 714 distally and proximally along a longitudinal axis of the end effector 702. In one aspect, the motor 704a may be coupled to the knife gear assembly, which includes a knife gear reduction set that includes a first knife drive gear and a second knife drive gear. A torque sensor 744a provides a firing force feedback signal to the control circuit 710. The firing force signal represents the force required to fire or displace the I-beam 714. A position sensor 734 may be configured to provide the position of the I-beam 714 along the firing stroke or the position of the firing member as a feedback signal to the control circuit 710. The end effector 702 may include additional sensors 738 configured to provide feedback signals to the control circuit 710. When ready to use, the control circuit 710 may provide a firing signal to the motor control 708a. In response to the firing signal, the motor 704a may drive the firing member distally along the longitudinal axis of the end effector 702 from a proximal stroke start position to a stroke end position distal to the stroke start position. As the firing member translates distally, an I-beam 714, with a cutting element positioned at a distal end, advances distally to cut tissue located between the staple cartridge 718 and the anvil 716.

[0548] In one aspect, the control circuit 710 is configured to drive a closure member such as the anvil 716 portion of the end effector 702. The control circuit 710 provides a motor set point to a motor control 708b, which provides a drive signal to the motor 704b. The output shaft of the motor

704b is coupled to a torque sensor 744b. The torque sensor 744b is coupled to a transmission 706b which is coupled to the anvil 716. The transmission 706b comprises movable mechanical elements such as rotating elements and a closure member to control the movement of the anvil 716 from the open and closed positions. In one aspect, the motor 704b is coupled to a closure gear assembly, which includes a closure reduction gear set that is supported in meshing engagement with the closure spur gear. The torque sensor 744b provides a closure force feedback signal to the control circuit 710. The closure force feedback signal represents the closure force applied to the anvil 716. The position sensor 734 may be configured to provide the position of the closure member as a feedback signal to the control circuit 710. Additional sensors 738 in the end effector 702 may provide the closure force feedback signal to the control circuit 710. The pivotable anvil 716 is positioned opposite the staple cartridge 718. When ready to use, the control circuit 710 may provide a closure signal to the motor control 708b. In response to the closure signal, the motor 704b advances a closure member to grasp tissue between the anvil 716 and the staple cartridge 718.

[0549] In one aspect, the control circuit 710 is configured to rotate a shaft member such as the shaft 740 to rotate the end effector 702. The control circuit 710 provides a motor set point to a motor control 708c, which provides a drive signal to the motor 704c. The output shaft of the motor 704c is coupled to a torque sensor 744c. The torque sensor 744c is coupled to a transmission 706c which is coupled to the shaft 740. The transmission 706c comprises movable mechanical elements such as rotating elements to control the rotation of the shaft 740 clockwise or counterclockwise up to and over 360°. In one aspect, the motor 704c is coupled to the rotational transmission assembly, which includes a tube gear segment that is formed on (or attached to) the proximal end of the proximal closure tube for operable engagement by a rotational gear assembly that is operably supported on the tool mounting plate. The torque sensor 744c provides a rotation force feedback signal to the control circuit 710. The rotation force feedback signal represents the rotation force applied to the shaft 740. The position sensor 734 may be configured to provide the position of the closure member as a feedback signal to the control circuit 710. Additional sensors 738 such as a shaft encoder may provide the rotational position of the shaft 740 to the control circuit 710.

[0550] In one aspect, the control circuit 710 is configured to articulate the end effector 702. The control circuit 710 provides a motor set point to a motor control 708d, which provides a drive signal to the motor 704d. The output shaft of the motor 704d is coupled to a torque sensor 744d. The torque sensor 744d is coupled to a transmission 706d which is coupled to an articulation member 742a. The transmission 706d comprises movable mechanical elements such as articulation elements to control the articulation of the end effector 702±65°. In one aspect, the motor 704d is coupled to an articulation nut, which is rotatably journaled on the proximal end portion of the distal spine portion and is rotatably driven thereon by an articulation gear assembly. The torque sensor 744d provides an articulation force feedback signal to the control circuit 710. The articulation force feedback signal represents the articulation force applied to the end effector 702. Sensors 738, such as an articulation

encoder, may provide the articulation position of the end effector **702** to the control circuit **710**.

[0551] In another aspect, the articulation function of the robotic surgical system **700** may comprise two articulation members, or links, **742a**, **742b**. These articulation members **742a**, **742b** are driven by separate disks on the robot interface (the rack) which are driven by the two motors **708d**, **708e**. When the separate firing motor **704a** is provided, each of articulation links **742a**, **742b** can be antagonistically driven with respect to the other link in order to provide a resistive holding motion and a load to the head when it is not moving and to provide an articulation motion as the head is articulated. The articulation members **742a**, **742b** attach to the head at a fixed radius as the head is rotated. Accordingly, the mechanical advantage of the push-and-pull link changes as the head is rotated. This change in the mechanical advantage may be more pronounced with other articulation link drive systems.

[0552] In one aspect, the one or more motors **704a-704e** may comprise a brushed DC motor with a gearbox and mechanical links to a firing member, closure member, or articulation member. Another example includes electric motors **704a-704e** that operate the movable mechanical elements such as the displacement member, articulation links, closure tube, and shaft. An outside influence is an unmeasured, unpredictable influence of things like tissue, surrounding bodies, and friction on the physical system. Such outside influence can be referred to as drag, which acts in opposition to one of electric motors **704a-704e**. The outside influence, such as drag, may cause the operation of the physical system to deviate from a desired operation of the physical system.

[0553] In one aspect, the position sensor **734** may be implemented as an absolute positioning system. In one aspect, the position sensor **734** may comprise a magnetic rotary absolute positioning system implemented as an AS5055EQFT single-chip magnetic rotary position sensor available from Austria Microsystems, AG. The position sensor **734** may interface with the control circuit **710** to provide an absolute positioning system. The position may include multiple Hall-effect elements located above a magnet and coupled to a CORDIC processor, also known as the digit-by-digit method and Volder's algorithm, that is provided to implement a simple and efficient algorithm to calculate hyperbolic and trigonometric functions that require only addition, subtraction, bitshift, and table lookup operations.

[0554] In one aspect, the control circuit **710** may be in communication with one or more sensors **738**. The sensors **738** may be positioned on the end effector **702** and adapted to operate with the robotic surgical instrument **700** to measure the various derived parameters such as the gap distance versus time, tissue compression versus time, and anvil strain versus time. The sensors **738** may comprise a magnetic sensor, a magnetic field sensor, a strain gauge, a load cell, a pressure sensor, a force sensor, a torque sensor, an inductive sensor such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor for measuring one or more parameters of the end effector **702**. The sensors **738** may include one or more sensors. The sensors **738** may be located on the staple cartridge **718** deck to determine tissue location using segmented electrodes. The torque sensors **744a-744e** may be configured to sense force such as firing force, closure force,

and/or articulation force, among others. Accordingly, the control circuit **710** can sense (1) the closure load experienced by the distal closure tube and its position, (2) the firing member at the rack and its position, (3) what portion of the staple cartridge **718** has tissue on it, and (4) the load and position on both articulation rods.

[0555] In one aspect, the one or more sensors **738** may comprise a strain gauge, such as a micro-strain gauge, configured to measure the magnitude of the strain in the anvil **716** during a clamped condition. The strain gauge provides an electrical signal whose amplitude varies with the magnitude of the strain. The sensors **738** may comprise a pressure sensor configured to detect a pressure generated by the presence of compressed tissue between the anvil **716** and the staple cartridge **718**. The sensors **738** may be configured to detect impedance of a tissue section located between the anvil **716** and the staple cartridge **718** that is indicative of the thickness and/or fullness of tissue located therebetween.

[0556] In one aspect, the sensors **738** may be implemented as one or more limit switches, electromechanical devices, solid-state switches, Hall-effect devices, magneto-resistive (MR) devices, giant magneto-resistive (GMR) devices, magnetometers, among others. In other implementations, the sensors **738** may be implemented as solid-state switches that operate under the influence of light, such as optical sensors, IR sensors, ultraviolet sensors, among others. Still, the switches may be solid-state devices such as transistors (e.g., FET, junction FET, MOSFET, bipolar, and the like). In other implementations, the sensors **738** may include electrical conductorless switches, ultrasonic switches, accelerometers, and inertial sensors, among others.

[0557] In one aspect, the sensors **738** may be configured to measure forces exerted on the anvil **716** by the closure drive system. For example, one or more sensors **738** can be at an interaction point between the closure tube and the anvil **716** to detect the closure forces applied by the closure tube to the anvil **716**. The forces exerted on the anvil **716** can be representative of the tissue compression experienced by the tissue section captured between the anvil **716** and the staple cartridge **718**. The one or more sensors **738** can be positioned at various interaction points along the closure drive system to detect the closure forces applied to the anvil **716** by the closure drive system. The one or more sensors **738** may be sampled in real time during a clamping operation by the processor of the control circuit **710**. The control circuit **710** receives real-time sample measurements to provide and analyze time-based information and assess, in real time, closure forces applied to the anvil **716**.

[0558] In one aspect, a current sensor **736** can be employed to measure the current drawn by each of the motors **704a-704e**. The force required to advance any of the movable mechanical elements such as the I-beam **714** corresponds to the current drawn by one of the motors **704a-704e**. The force is converted to a digital signal and provided to the control circuit **710**. The control circuit **710** can be configured to simulate the response of the actual system of the instrument in the software of the controller. A displacement member can be actuated to move an I-beam **714** in the end effector **702** at or near a target velocity. The robotic surgical instrument **700** can include a feedback controller, which can be one of any feedback controllers, including, but not limited to a PID, a state feedback, a linear-quadratic (LQR), and/or an adaptive controller, for example. The robotic surgical instrument **700** can include a power source

to convert the signal from the feedback controller into a physical input such as case voltage, PWM voltage, frequency modulated voltage, current, torque, and/or force, for example. Additional details are disclosed in U.S. patent application Ser. No. 15/636,829, titled CLOSED LOOP VELOCITY CONTROL TECHNIQUES FOR ROBOTIC SURGICAL INSTRUMENT, filed Jun. 29, 2017, which is herein incorporated by reference in its entirety.

[0559] FIG. 18 illustrates a block diagram of a surgical instrument 750 programmed to control the distal translation of a displacement member according to one aspect of this disclosure. In one aspect, the surgical instrument 750 is programmed to control the distal translation of a displacement member such as the I-beam 764. The surgical instrument 750 comprises an end effector 752 that may comprise an anvil 766, an I-beam 764 (including a sharp cutting edge), and a removable staple cartridge 768.

[0560] The position, movement, displacement, and/or translation of a linear displacement member, such as the I-beam 764, can be measured by an absolute positioning system, sensor arrangement, and position sensor 784. Because the I-beam 764 is coupled to a longitudinally movable drive member, the position of the I-beam 764 can be determined by measuring the position of the longitudinally movable drive member employing the position sensor 784. Accordingly, in the following description, the position, displacement, and/or translation of the I-beam 764 can be achieved by the position sensor 784 as described herein. A control circuit 760 may be programmed to control the translation of the displacement member, such as the I-beam 764. The control circuit 760, in some examples, may comprise one or more microcontrollers, microprocessors, or other suitable processors for executing instructions that cause the processor or processors to control the displacement member, e.g., the I-beam 764, in the manner described. In one aspect, a timer/counter 781 provides an output signal, such as the elapsed time or a digital count, to the control circuit 760 to correlate the position of the I-beam 764 as determined by the position sensor 784 with the output of the timer/counter 781 such that the control circuit 760 can determine the position of the I-beam 764 at a specific time (t) relative to a starting position. The timer/counter 781 may be configured to measure elapsed time, count external events, or time external events.

[0561] The control circuit 760 may generate a motor set point signal 772. The motor set point signal 772 may be provided to a motor controller 758. The motor controller 758 may comprise one or more circuits configured to provide a motor drive signal 774 to the motor 754 to drive the motor 754 as described herein. In some examples, the motor 754 may be a brushed DC electric motor. For example, the velocity of the motor 754 may be proportional to the motor drive signal 774. In some examples, the motor 754 may be a brushless DC electric motor and the motor drive signal 774 may comprise a PWM signal provided to one or more stator windings of the motor 754. Also, in some examples, the motor controller 758 may be omitted, and the control circuit 760 may generate the motor drive signal 774 directly.

[0562] The motor 754 may receive power from an energy source 762. The energy source 762 may be or include a battery, a super capacitor, or any other suitable energy source. The motor 754 may be mechanically coupled to the I-beam 764 via a transmission 756. The transmission 756 may include one or more gears or other linkage components

to couple the motor 754 to the I-beam 764. A position sensor 784 may sense a position of the I-beam 764. The position sensor 784 may be or include any type of sensor that is capable of generating position data that indicate a position of the I-beam 764. In some examples, the position sensor 784 may include an encoder configured to provide a series of pulses to the control circuit 760 as the I-beam 764 translates distally and proximally. The control circuit 760 may track the pulses to determine the position of the I-beam 764. Other suitable position sensors may be used, including, for example, a proximity sensor. Other types of position sensors may provide other signals indicating motion of the I-beam 764. Also, in some examples, the position sensor 784 may be omitted. Where the motor 754 is a stepper motor, the control circuit 760 may track the position of the I-beam 764 by aggregating the number and direction of steps that the motor 754 has been instructed to execute. The position sensor 784 may be located in the end effector 752 or at any other portion of the instrument.

[0563] The control circuit 760 may be in communication with one or more sensors 788. The sensors 788 may be positioned on the end effector 752 and adapted to operate with the surgical instrument 750 to measure the various derived parameters such as gap distance versus time, tissue compression versus time, and anvil strain versus time. The sensors 788 may comprise a magnetic sensor, a magnetic field sensor, a strain gauge, a pressure sensor, a force sensor, an inductive sensor such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor for measuring one or more parameters of the end effector 752. The sensors 788 may include one or more sensors.

[0564] The one or more sensors 788 may comprise a strain gauge, such as a micro-strain gauge, configured to measure the magnitude of the strain in the anvil 766 during a clamped condition. The strain gauge provides an electrical signal whose amplitude varies with the magnitude of the strain. The sensors 788 may comprise a pressure sensor configured to detect a pressure generated by the presence of compressed tissue between the anvil 766 and the staple cartridge 768. The sensors 788 may be configured to detect impedance of a tissue section located between the anvil 766 and the staple cartridge 768 that is indicative of the thickness and/or fullness of tissue located therebetween.

[0565] The sensors 788 may be configured to measure forces exerted on the anvil 766 by a closure drive system. For example, one or more sensors 788 can be at an interaction point between a closure tube and the anvil 766 to detect the closure forces applied by a closure tube to the anvil 766. The forces exerted on the anvil 766 can be representative of the tissue compression experienced by the tissue section captured between the anvil 766 and the staple cartridge 768. The one or more sensors 788 can be positioned at various interaction points along the closure drive system to detect the closure forces applied to the anvil 766 by the closure drive system. The one or more sensors 788 may be sampled in real time during a clamping operation by a processor of the control circuit 760. The control circuit 760 receives real-time sample measurements to provide and analyze time-based information and assess, in real time, closure forces applied to the anvil 766.

[0566] A current sensor 786 can be employed to measure the current drawn by the motor 754. The force required to advance the I-beam 764 corresponds to the current drawn by



the motor 754. The force is converted to a digital signal and provided to the control circuit 760.

[0567] The control circuit 760 can be configured to simulate the response of the actual system of the instrument in the software of the controller. A displacement member can be actuated to move an I-beam 764 in the end effector 752 at or near a target velocity. The surgical instrument 750 can include a feedback controller, which can be one of any feedback controllers, including, but not limited to a PID, a state feedback, LQR, and/or an adaptive controller, for example. The surgical instrument 750 can include a power source to convert the signal from the feedback controller into a physical input such as case voltage, PWM voltage, frequency modulated voltage, current, torque, and/or force, for example.

[0568] The actual drive system of the surgical instrument 750 is configured to drive the displacement member, cutting member, or I-beam 764, by a brushed DC motor with gearbox and mechanical links to an articulation and/or knife system. Another example is the electric motor 754 that operates the displacement member and the articulation driver, for example, of an interchangeable shaft assembly. An outside influence is an unmeasured, unpredictable influence of things like tissue, surrounding bodies and friction on the physical system. Such outside influence can be referred to as drag which acts in opposition to the electric motor 754. The outside influence, such as drag, may cause the operation of the physical system to deviate from a desired operation of the physical system.

[0569] Various example aspects are directed to a surgical instrument 750 comprising an end effector 752 with motor-driven surgical stapling and cutting implements. For example, a motor 754 may drive a displacement member distally and proximally along a longitudinal axis of the end effector 752. The end effector 752 may comprise a pivotable anvil 766 and, when configured for use, a staple cartridge 768 positioned opposite the anvil 766. A clinician may grasp tissue between the anvil 766 and the staple cartridge 768, as described herein. When ready to use the instrument 750, the clinician may provide a firing signal, for example by depressing a trigger of the instrument 750. In response to the firing signal, the motor 754 may drive the displacement member distally along the longitudinal axis of the end effector 752 from a proximal stroke begin position to a stroke end position distal of the stroke begin position. As the displacement member translates distally, an I-beam 764 with a cutting element positioned at a distal end, may cut the tissue between the staple cartridge 768 and the anvil 766.

[0570] In various examples, the surgical instrument 750 may comprise a control circuit 760 programmed to control the distal translation of the displacement member, such as the I-beam 764, for example, based on one or more tissue conditions. The control circuit 760 may be programmed to sense tissue conditions, such as thickness, either directly or indirectly, as described herein. The control circuit 760 may be programmed to select a firing control program based on tissue conditions. A firing control program may describe the distal motion of the displacement member. Different firing control programs may be selected to better treat different tissue conditions. For example, when thicker tissue is present, the control circuit 760 may be programmed to translate the displacement member at a lower velocity and/or with lower power. When thinner tissue is present, the control

circuit 760 may be programmed to translate the displacement member at a higher velocity and/or with higher power.

[0571] In some examples, the control circuit 760 may initially operate the motor 754 in an open loop configuration for a first open loop portion of a stroke of the displacement member. Based on a response of the instrument 750 during the open loop portion of the stroke, the control circuit 760 may select a firing control program. The response of the instrument may include, a translation distance of the displacement member during the open loop portion, a time elapsed during the open loop portion, energy provided to the motor 754 during the open loop portion, a sum of pulse widths of a motor drive signal, etc. After the open loop portion, the control circuit 760 may implement the selected firing control program for a second portion of the displacement member stroke. For example, during the closed loop portion of the stroke, the control circuit 760 may modulate the motor 754 based on translation data describing a position of the displacement member in a closed loop manner to translate the displacement member at a constant velocity. Additional details are disclosed in U.S. patent application Ser. No. 15/720,852, titled SYSTEM AND METHODS FOR CONTROLLING A DISPLAY OF A SURGICAL INSTRUMENT, filed Sep. 29, 2017, which is herein incorporated by reference in its entirety.

[0572] FIG. 19 is a schematic diagram of a surgical instrument 790 configured to control various functions according to one aspect of this disclosure. In one aspect, the surgical instrument 790 is programmed to control distal translation of a displacement member such as the I-beam 764. The surgical instrument 790 comprises an end effector 792 that may comprise an anvil 766, an I-beam 764, and a removable staple cartridge 768 which may be interchanged with an RF cartridge 796 (shown in dashed line).

[0573] In one aspect, sensors 788 may be implemented as a limit switch, electromechanical device, solid-state switches, Hall-effect devices, MR devices, GMR devices, magnetometers, among others. In other implementations, the sensors 638 may be solid-state switches that operate under the influence of light, such as optical sensors, IR sensors, ultraviolet sensors, among others. Still, the switches may be solid-state devices such as transistors (e.g., FET, junction FET, MOSFET, bipolar, and the like). In other implementations, the sensors 788 may include electrical conductorless switches, ultrasonic switches, accelerometers, and inertial sensors, among others.

[0574] In one aspect, the position sensor 784 may be implemented as an absolute positioning system comprising a magnetic rotary absolute positioning system implemented as an AS5055EQFT single-chip magnetic rotary position sensor available from Austria Microsystems, AG. The position sensor 784 may interface with the control circuit 760 to provide an absolute positioning system. The position may include multiple Hall-effect elements located above a magnet and coupled to a CORDIC processor, also known as the digit-by-digit method and Volder's algorithm, that is provided to implement a simple and efficient algorithm to calculate hyperbolic and trigonometric functions that require only addition, subtraction, bitshift, and table lookup operations.

[0575] In one aspect, the I-beam 764 may be implemented as a knife member comprising a knife body that operably supports a tissue cutting blade thereon and may further include anvil engagement tabs or features and channel

engagement features or a foot. In one aspect, the staple cartridge **768** may be implemented as a standard (mechanical) surgical fastener cartridge. In one aspect, the RF cartridge **796** may be implemented as an RF cartridge. These and other sensors arrangements are described in commonly-owned U.S. patent application Ser. No. 15/628,175, titled TECHNIQUES FOR ADAPTIVE CONTROL OF MOTOR VELOCITY OF A SURGICAL STAPLING AND CUTTING INSTRUMENT, filed Jun. 20, 2017, which is herein incorporated by reference in its entirety.

[0576] The position, movement, displacement, and/or translation of a linear displacement member, such as the I-beam **764**, can be measured by an absolute positioning system, sensor arrangement, and position sensor represented as position sensor **784**. Because the I-beam **764** is coupled to the longitudinally movable drive member, the position of the I-beam **764** can be determined by measuring the position of the longitudinally movable drive member employing the position sensor **784**. Accordingly, in the following description, the position, displacement, and/or translation of the I-beam **764** can be achieved by the position sensor **784** as described herein. A control circuit **760** may be programmed to control the translation of the displacement member, such as the I-beam **764**, as described herein. The control circuit **760**, in some examples, may comprise one or more micro-controllers, microprocessors, or other suitable processors for executing instructions that cause the processor or processors to control the displacement member, e.g., the I-beam **764**, in the manner described. In one aspect, a timer/counter **781** provides an output signal, such as the elapsed time or a digital count, to the control circuit **760** to correlate the position of the I-beam **764** as determined by the position sensor **784** with the output of the timer/counter **781** such that the control circuit **760** can determine the position of the I-beam **764** at a specific time (t) relative to a starting position. The timer/counter **781** may be configured to measure elapsed time, count external events, or time external events.

[0577] The control circuit **760** may generate a motor set point signal **772**. The motor set point signal **772** may be provided to a motor controller **758**. The motor controller **758** may comprise one or more circuits configured to provide a motor drive signal **774** to the motor **754** to drive the motor **754** as described herein. In some examples, the motor **754** may be a brushed DC electric motor. For example, the velocity of the motor **754** may be proportional to the motor drive signal **774**. In some examples, the motor **754** may be a brushless DC electric motor and the motor drive signal **774** may comprise a PWM signal provided to one or more stator windings of the motor **754**. Also, in some examples, the motor controller **758** may be omitted, and the control circuit **760** may generate the motor drive signal **774** directly.

[0578] The motor **754** may receive power from an energy source **762**. The energy source **762** may be or include a battery, a super capacitor, or any other suitable energy source. The motor **754** may be mechanically coupled to the I-beam **764** via a transmission **756**. The transmission **756** may include one or more gears or other linkage components to couple the motor **754** to the I-beam **764**. A position sensor **784** may sense a position of the I-beam **764**. The position sensor **784** may be or include any type of sensor that is capable of generating position data that indicate a position of the I-beam **764**. In some examples, the position sensor **784** may include an encoder configured to provide a series of

pulses to the control circuit **760** as the I-beam **764** translates distally and proximally. The control circuit **760** may track the pulses to determine the position of the I-beam **764**. Other suitable position sensors may be used, including, for example, a proximity sensor. Other types of position sensors may provide other signals indicating motion of the I-beam **764**. Also, in some examples, the position sensor **784** may be omitted. Where the motor **754** is a stepper motor, the control circuit **760** may track the position of the I-beam **764** by aggregating the number and direction of steps that the motor has been instructed to execute. The position sensor **784** may be located in the end effector **792** or at any other portion of the instrument.

[0579] The control circuit **760** may be in communication with one or more sensors **788**. The sensors **788** may be positioned on the end effector **792** and adapted to operate with the surgical instrument **790** to measure the various derived parameters such as gap distance versus time, tissue compression versus time, and anvil strain versus time. The sensors **788** may comprise a magnetic sensor, a magnetic field sensor, a strain gauge, a pressure sensor, a force sensor, an inductive sensor such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor for measuring one or more parameters of the end effector **792**. The sensors **788** may include one or more sensors.

[0580] The one or more sensors **788** may comprise a strain gauge, such as a micro-strain gauge, configured to measure the magnitude of the strain in the anvil **766** during a clamped condition. The strain gauge provides an electrical signal whose amplitude varies with the magnitude of the strain. The sensors **788** may comprise a pressure sensor configured to detect a pressure generated by the presence of compressed tissue between the anvil **766** and the staple cartridge **768**. The sensors **788** may be configured to detect impedance of a tissue section located between the anvil **766** and the staple cartridge **768** that is indicative of the thickness and/or fullness of tissue located therebetween.

[0581] The sensors **788** may be is configured to measure forces exerted on the anvil **766** by the closure drive system. For example, one or more sensors **788** can be at an interaction point between a closure tube and the anvil **766** to detect the closure forces applied by a closure tube to the anvil **766**. The forces exerted on the anvil **766** can be representative of the tissue compression experienced by the tissue section captured between the anvil **766** and the staple cartridge **768**. The one or more sensors **788** can be positioned at various interaction points along the closure drive system to detect the closure forces applied to the anvil **766** by the closure drive system. The one or more sensors **788** may be sampled in real time during a clamping operation by a processor portion of the control circuit **760**. The control circuit **760** receives real-time sample measurements to provide and analyze time-based information and assess, in real time, closure forces applied to the anvil **766**.

[0582] A current sensor **786** can be employed to measure the current drawn by the motor **754**. The force required to advance the I-beam **764** corresponds to the current drawn by the motor **754**. The force is converted to a digital signal and provided to the control circuit **760**.

[0583] An RF energy source **794** is coupled to the end effector **792** and is applied to the RF cartridge **796** when the RF cartridge **796** is loaded in the end effector **792** in place

of the staple cartridge **768**. The control circuit **760** controls the delivery of the RF energy to the RF cartridge **796**.

[0584] Additional details are disclosed in U.S. patent application Ser. No. 15/636,096, titled SURGICAL SYSTEM COUPLABLE WITH STAPLE CARTRIDGE AND RADIO FREQUENCY CARTRIDGE, AND METHOD OF USING SAME, filed Jun. 28, 2017, which is herein incorporated by reference in its entirety.

#### Generator Hardware

[0585] FIG. 20 is a simplified block diagram of a generator **800** configured to provide inductorless tuning, among other benefits. Additional details of the generator **800** are described in U.S. Pat. No. 9,060,775, titled SURGICAL GENERATOR FOR ULTRASONIC AND ELECTROSURGICAL DEVICES, which issued on Jun. 23, 2015, which is herein incorporated by reference in its entirety. The generator **800** may comprise a patient isolated stage **802** in communication with a non-isolated stage **804** via a power transformer **806**. A secondary winding **808** of the power transformer **806** is contained in the isolated stage **802** and may comprise a tapped configuration (e.g., a center-tapped or a non-center-tapped configuration) to define drive signal outputs **810a**, **810b**, **810c** for delivering drive signals to different surgical instruments, such as, for example, an ultrasonic surgical instrument, an RF electro-surgical instrument, and a multifunction surgical instrument which includes ultrasonic and RF energy modes that can be delivered alone or simultaneously. In particular, drive signal outputs **810a**, **810c** may output an ultrasonic drive signal (e.g., a 420V root-mean-square (RMS) drive signal) to an ultrasonic surgical instrument, and drive signal outputs **810b**, **810c** may output an RF electro-surgical drive signal (e.g., a 100V RMS drive signal) to an RF electro-surgical instrument, with the drive signal output **810b** corresponding to the center tap of the power transformer **806**.

[0586] In certain forms, the ultrasonic and electro-surgical drive signals may be provided simultaneously to distinct surgical instruments and/or to a single surgical instrument, such as the multifunction surgical instrument, having the capability to deliver both ultrasonic and electro-surgical energy to tissue. It will be appreciated that the electro-surgical signal, provided either to a dedicated electro-surgical instrument and/or to a combined multifunction ultrasonic/electro-surgical instrument may be either a therapeutic or sub-therapeutic level signal where the sub-therapeutic signal can be used, for example, to monitor tissue or instrument conditions and provide feedback to the generator. For example, the ultrasonic and RF signals can be delivered separately or simultaneously from a generator with a single output port in order to provide the desired output signal to the surgical instrument, as will be discussed in more detail below. Accordingly, the generator can combine the ultrasonic and electro-surgical RF energies and deliver the combined energies to the multifunction ultrasonic/electro-surgical instrument. Bipolar electrodes can be placed on one or both jaws of the end effector. One jaw may be driven by ultrasonic energy in addition to electro-surgical RF energy, working simultaneously. The ultrasonic energy may be employed to dissect tissue, while the electro-surgical RF energy may be employed for vessel sealing.

[0587] The non-isolated stage **804** may comprise a power amplifier **812** having an output connected to a primary winding **814** of the power transformer **806**. In certain forms,

the power amplifier **812** may comprise a push-pull amplifier. For example, the non-isolated stage **804** may further comprise a logic device **816** for supplying a digital output to a digital-to-analog converter (DAC) circuit **818**, which in turn supplies a corresponding analog signal to an input of the power amplifier **812**. In certain forms, the logic device **816** may comprise a programmable gate array (PGA), a FPGA, programmable logic device (PLD), among other logic circuits, for example. The logic device **816**, by virtue of controlling the input of the power amplifier **812** via the DAC circuit **818**, may therefore control any of a number of parameters (e.g., frequency, waveform shape, waveform amplitude) of drive signals appearing at the drive signal outputs **810a**, **810b**, **810c**. In certain forms and as discussed below, the logic device **816**, in conjunction with a processor (e.g., a DSP discussed below), may implement a number of DSP-based and/or other control algorithms to control parameters of the drive signals output by the generator **800**.

[0588] Power may be supplied to a power rail of the power amplifier **812** by a switch-mode regulator **820**, e.g., a power converter. In certain forms, the switch-mode regulator **820** may comprise an adjustable buck regulator, for example. The non-isolated stage **804** may further comprise a first processor **822**, which in one form may comprise a DSP processor such as an Analog Devices ADSP-21469 SHARC DSP, available from Analog Devices, Norwood, Mass., for example, although in various forms any suitable processor may be employed. In certain forms the DSP processor **822** may control the operation of the switch-mode regulator **820** responsive to voltage feedback data received from the power amplifier **812** by the DSP processor **822** via an ADC circuit **824**. In one form, for example, the DSP processor **822** may receive as input, via the ADC circuit **824**, the waveform envelope of a signal (e.g., an RF signal) being amplified by the power amplifier **812**. The DSP processor **822** may then control the switch-mode regulator **820** (e.g., via a PWM output) such that the rail voltage supplied to the power amplifier **812** tracks the waveform envelope of the amplified signal. By dynamically modulating the rail voltage of the power amplifier **812** based on the waveform envelope, the efficiency of the power amplifier **812** may be significantly improved relative to a fixed rail voltage amplifier schemes.

[0589] In certain forms, the logic device **816**, in conjunction with the DSP processor **822**, may implement a digital synthesis circuit such as a direct digital synthesizer control scheme to control the waveform shape, frequency, and/or amplitude of drive signals output by the generator **800**. In one form, for example, the logic device **816** may implement a DDS control algorithm by recalling waveform samples stored in a dynamically updated lookup table (LUT), such as a RAM LUT, which may be embedded in an FPGA. This control algorithm is particularly useful for ultrasonic applications in which an ultrasonic transducer, such as an ultrasonic transducer, may be driven by a clean sinusoidal current at its resonant frequency. Because other frequencies may excite parasitic resonances, minimizing or reducing the total distortion of the motional branch current may correspondingly minimize or reduce undesirable resonance effects. Because the waveform shape of a drive signal output by the generator **800** is impacted by various sources of distortion present in the output drive circuit (e.g., the power transformer **806**, the power amplifier **812**), voltage and current feedback data based on the drive signal may be input into an algorithm, such as an error control algorithm implemented

by the DSP processor **822**, which compensates for distortion by suitably pre-distorting or modifying the waveform samples stored in the LUT on a dynamic, ongoing basis (e.g., in real time). In one form, the amount or degree of pre-distortion applied to the LUT samples may be based on the error between a computed motional branch current and a desired current waveform shape, with the error being determined on a sample-by-sample basis. In this way, the pre-distorted LUT samples, when processed through the drive circuit, may result in a motional branch drive signal having the desired waveform shape (e.g., sinusoidal) for optimally driving the ultrasonic transducer. In such forms, the LUT waveform samples will therefore not represent the desired waveform shape of the drive signal, but rather the waveform shape that is required to ultimately produce the desired waveform shape of the motional branch drive signal when distortion effects are taken into account.

**[0590]** The non-isolated stage **804** may further comprise a first ADC circuit **826** and a second ADC circuit **828** coupled to the output of the power transformer **806** via respective isolation transformers **830**, **832** for respectively sampling the voltage and current of drive signals output by the generator **800**. In certain forms, the ADC circuits **826**, **828** may be configured to sample at high speeds (e.g., 80 mega samples per second (MSPS)) to enable oversampling of the drive signals. In one form, for example, the sampling speed of the ADC circuits **826**, **828** may enable approximately 200× (depending on frequency) oversampling of the drive signals. In certain forms, the sampling operations of the ADC circuit **826**, **828** may be performed by a single ADC circuit receiving input voltage and current signals via a two-way multiplexer. The use of high-speed sampling in forms of the generator **800** may enable, among other things, calculation of the complex current flowing through the motional branch (which may be used in certain forms to implement DDS-based waveform shape control described above), accurate digital filtering of the sampled signals, and calculation of real power consumption with a high degree of precision. Voltage and current feedback data output by the ADC circuits **826**, **828** may be received and processed (e.g., first-in-first-out (FIFO) buffer, multiplexer) by the logic device **816** and stored in data memory for subsequent retrieval by, for example, the DSP processor **822**. As noted above, voltage and current feedback data may be used as input to an algorithm for pre-distorting or modifying LUT waveform samples on a dynamic and ongoing basis. In certain forms, this may require each stored voltage and current feedback data pair to be indexed based on, or otherwise associated with, a corresponding LUT sample that was output by the logic device **816** when the voltage and current feedback data pair was acquired. Synchronization of the LUT samples and the voltage and current feedback data in this manner contributes to the correct timing and stability of the pre-distortion algorithm.

**[0591]** In certain forms, the voltage and current feedback data may be used to control the frequency and/or amplitude (e.g., current amplitude) of the drive signals. In one form, for example, voltage and current feedback data may be used to determine impedance phase. The frequency of the drive signal may then be controlled to minimize or reduce the difference between the determined impedance phase and an impedance phase setpoint (e.g., 0°), thereby minimizing or reducing the effects of harmonic distortion and correspondingly enhancing impedance phase measurement accuracy.

The determination of phase impedance and a frequency control signal may be implemented in the DSP processor **822**, for example, with the frequency control signal being supplied as input to a DDS control algorithm implemented by the logic device **816**.

**[0592]** In another form, for example, the current feedback data may be monitored in order to maintain the current amplitude of the drive signal at a current amplitude setpoint. The current amplitude setpoint may be specified directly or determined indirectly based on specified voltage amplitude and power setpoints. In certain forms, control of the current amplitude may be implemented by control algorithm, such as, for example, a proportional-integral-derivative (PID) control algorithm, in the DSP processor **822**. Variables controlled by the control algorithm to suitably control the current amplitude of the drive signal may include, for example, the scaling of the LUT waveform samples stored in the logic device **816** and/or the full-scale output voltage of the DAC circuit **818** (which supplies the input to the power amplifier **812**) via a DAC circuit **834**.

**[0593]** The non-isolated stage **804** may further comprise a second processor **836** for providing, among other things user interface (UI) functionality. In one form, the UI processor **836** may comprise an Atmel AT91SAM9263 processor having an ARM 926EJ-S core, available from Atmel Corporation, San Jose, Calif., for example. Examples of UI functionality supported by the UI processor **836** may include audible and visual user feedback, communication with peripheral devices (e.g., via a USB interface), communication with a foot switch, communication with an input device (e.g., a touch screen display) and communication with an output device (e.g., a speaker). The UI processor **836** may communicate with the DSP processor **822** and the logic device **816** (e.g., via SPI buses). Although the UI processor **836** may primarily support UI functionality, it may also coordinate with the DSP processor **822** to implement hazard mitigation in certain forms. For example, the UI processor **836** may be programmed to monitor various aspects of user input and/or other inputs (e.g., touch screen inputs, foot switch inputs, temperature sensor inputs) and may disable the drive output of the generator **800** when an erroneous condition is detected.

**[0594]** In certain forms, both the DSP processor **822** and the UI processor **836**, for example, may determine and monitor the operating state of the generator **800**. For the DSP processor **822**, the operating state of the generator **800** may dictate, for example, which control and/or diagnostic processes are implemented by the DSP processor **822**. For the UI processor **836**, the operating state of the generator **800** may dictate, for example, which elements of a UI (e.g., display screens, sounds) are presented to a user. The respective DSP and UI processors **822**, **836** may independently maintain the current operating state of the generator **800** and recognize and evaluate possible transitions out of the current operating state. The DSP processor **822** may function as the master in this relationship and determine when transitions between operating states are to occur. The UI processor **836** may be aware of valid transitions between operating states and may confirm if a particular transition is appropriate. For example, when the DSP processor **822** instructs the UI processor **836** to transition to a specific state, the UI processor **836** may verify that requested transition is valid. In the event that a requested transition between states is deter-

mined to be invalid by the UI processor **836**, the UI processor **836** may cause the generator **800** to enter a failure mode.

[0595] The non-isolated stage **804** may further comprise a controller **838** for monitoring input devices (e.g., a capacitive touch sensor used for turning the generator **800** on and off, a capacitive touch screen). In certain forms, the controller **838** may comprise at least one processor and/or other controller device in communication with the UI processor **836**. In one form, for example, the controller **838** may comprise a processor (e.g., a Meg168 8-bit controller available from Atmel) configured to monitor user input provided via one or more capacitive touch sensors. In one form, the controller **838** may comprise a touch screen controller (e.g., a QT5480 touch screen controller available from Atmel) to control and manage the acquisition of touch data from a capacitive touch screen.

[0596] In certain forms, when the generator **800** is in a “power off” state, the controller **838** may continue to receive operating power (e.g., via a line from a power supply of the generator **800**, such as the power supply **854** discussed below). In this way, the controller **838** may continue to monitor an input device (e.g., a capacitive touch sensor located on a front panel of the generator **800**) for turning the generator **800** on and off. When the generator **800** is in the power off state, the controller **838** may wake the power supply (e.g., enable operation of one or more DC/DC voltage converters **856** of the power supply **854**) if activation of the “on/off” input device by a user is detected. The controller **838** may therefore initiate a sequence for transitioning the generator **800** to a “power on” state. Conversely, the controller **838** may initiate a sequence for transitioning the generator **800** to the power off state if activation of the “on/off” input device is detected when the generator **800** is in the power on state. In certain forms, for example, the controller **838** may report activation of the “on/off” input device to the UI processor **836**, which in turn implements the necessary process sequence for transitioning the generator **800** to the power off state. In such forms, the controller **838** may have no independent ability for causing the removal of power from the generator **800** after its power on state has been established.

[0597] In certain forms, the controller **838** may cause the generator **800** to provide audible or other sensory feedback for alerting the user that a power on or power off sequence has been initiated. Such an alert may be provided at the beginning of a power on or power off sequence and prior to the commencement of other processes associated with the sequence.

[0598] In certain forms, the isolated stage **802** may comprise an instrument interface circuit **840** to, for example, provide a communication interface between a control circuit of a surgical instrument (e.g., a control circuit comprising handpiece switches) and components of the non-isolated stage **804**, such as, for example, the logic device **816**, the DSP processor **822**, and/or the UI processor **836**. The instrument interface circuit **840** may exchange information with components of the non-isolated stage **804** via a communication link that maintains a suitable degree of electrical isolation between the isolated and non-isolated stages **802**, **804**, such as, for example, an IR-based communication link. Power may be supplied to the instrument interface circuit

**840** using, for example, a low-dropout voltage regulator powered by an isolation transformer driven from the non-isolated stage **804**.

[0599] In one form, the instrument interface circuit **840** may comprise a logic circuit **842** (e.g., logic circuit, programmable logic circuit, PGA, FPGA, PLD) in communication with a signal conditioning circuit **844**. The signal conditioning circuit **844** may be configured to receive a periodic signal from the logic circuit **842** (e.g., a 2 kHz square wave) to generate a bipolar interrogation signal having an identical frequency. The interrogation signal may be generated, for example, using a bipolar current source fed by a differential amplifier. The interrogation signal may be communicated to a surgical instrument control circuit (e.g., by using a conductive pair in a cable that connects the generator **800** to the surgical instrument) and monitored to determine a state or configuration of the control circuit. The control circuit may comprise a number of switches, resistors, and/or diodes to modify one or more characteristics (e.g., amplitude, rectification) of the interrogation signal such that a state or configuration of the control circuit is uniquely discernable based on the one or more characteristics. In one form, for example, the signal conditioning circuit **844** may comprise an ADC circuit for generating samples of a voltage signal appearing across inputs of the control circuit resulting from passage of interrogation signal therethrough. The logic circuit **842** (or a component of the non-isolated stage **804**) may then determine the state or configuration of the control circuit based on the ADC circuit samples.

[0600] In one form, the instrument interface circuit **840** may comprise a first data circuit interface **846** to enable information exchange between the logic circuit **842** (or other element of the instrument interface circuit **840**) and a first data circuit disposed in or otherwise associated with a surgical instrument. In certain forms, for example, a first data circuit may be disposed in a cable integrally attached to a surgical instrument handpiece or in an adaptor for interfacing a specific surgical instrument type or model with the generator **800**. The first data circuit may be implemented in any suitable manner and may communicate with the generator according to any suitable protocol, including, for example, as described herein with respect to the first data circuit. In certain forms, the first data circuit may comprise a non-volatile storage device, such as an EEPROM device. In certain forms, the first data circuit interface **846** may be implemented separately from the logic circuit **842** and comprise suitable circuitry (e.g., discrete logic devices, a processor) to enable communication between the logic circuit **842** and the first data circuit. In other forms, the first data circuit interface **846** may be integral with the logic circuit **842**.

[0601] In certain forms, the first data circuit may store information pertaining to the particular surgical instrument with which it is associated. Such information may include, for example, a model number, a serial number, a number of operations in which the surgical instrument has been used, and/or any other type of information. This information may be read by the instrument interface circuit **840** (e.g., by the logic circuit **842**), transferred to a component of the non-isolated stage **804** (e.g., to logic device **816**, DSP processor **822**, and/or UI processor **836**) for presentation to a user via an output device and/or for controlling a function or operation of the generator **800**. Additionally, any type of information may be communicated to the first data circuit for

storage therein via the first data circuit interface **846** (e.g., using the logic circuit **842**). Such information may comprise, for example, an updated number of operations in which the surgical instrument has been used and/or dates and/or times of its usage.

[0602] As discussed previously, a surgical instrument may be detachable from a handpiece (e.g., the multifunction surgical instrument may be detachable from the handpiece) to promote instrument interchangeability and/or disposability. In such cases, conventional generators may be limited in their ability to recognize particular instrument configurations being used and to optimize control and diagnostic processes accordingly. The addition of readable data circuits to surgical instruments to address this issue is problematic from a compatibility standpoint, however. For example, designing a surgical instrument to remain backwardly compatible with generators that lack the requisite data reading functionality may be impractical due to, for example, differing signal schemes, design complexity, and cost. Forms of instruments discussed herein address these concerns by using data circuits that may be implemented in existing surgical instruments economically and with minimal design changes to preserve compatibility of the surgical instruments with current generator platforms.

[0603] Additionally, forms of the generator **800** may enable communication with instrument-based data circuits. For example, the generator **800** may be configured to communicate with a second data circuit contained in an instrument (e.g., the multifunction surgical instrument). In some forms, the second data circuit may be implemented in a many similar to that of the first data circuit described herein. The instrument interface circuit **840** may comprise a second data circuit interface **848** to enable this communication. In one form, the second data circuit interface **848** may comprise a tri-state digital interface, although other interfaces may also be used. In certain forms, the second data circuit may generally be any circuit for transmitting and/or receiving data. In one form, for example, the second data circuit may store information pertaining to the particular surgical instrument with which it is associated. Such information may include, for example, a model number, a serial number, a number of operations in which the surgical instrument has been used, and/or any other type of information.

[0604] In some forms, the second data circuit may store information about the electrical and/or ultrasonic properties of an associated ultrasonic transducer, end effector, or ultrasonic drive system. For example, the first data circuit may indicate a burn-in frequency slope, as described herein. Additionally or alternatively, any type of information may be communicated to second data circuit for storage therein via the second data circuit interface **848** (e.g., using the logic circuit **842**). Such information may comprise, for example, an updated number of operations in which the instrument has been used and/or dates and/or times of its usage. In certain forms, the second data circuit may transmit data acquired by one or more sensors (e.g., an instrument-based temperature sensor). In certain forms, the second data circuit may receive data from the generator **800** and provide an indication to a user (e.g., a light emitting diode indication or other visible indication) based on the received data.

[0605] In certain forms, the second data circuit and the second data circuit interface **848** may be configured such that communication between the logic circuit **842** and the

second data circuit can be effected without the need to provide additional conductors for this purpose (e.g., dedicated conductors of a cable connecting a handpiece to the generator **800**). In one form, for example, information may be communicated to and from the second data circuit using a one-wire bus communication scheme implemented on existing cabling, such as one of the conductors used transmit interrogation signals from the signal conditioning circuit **844** to a control circuit in a handpiece. In this way, design changes or modifications to the surgical instrument that might otherwise be necessary are minimized or reduced. Moreover, because different types of communications implemented over a common physical channel can be frequency-band separated, the presence of a second data circuit may be “invisible” to generators that do not have the requisite data reading functionality, thus enabling backward compatibility of the surgical instrument.

[0606] In certain forms, the isolated stage **802** may comprise at least one blocking capacitor **850-1** connected to the drive signal output **810b** to prevent passage of DC current to a patient. A single blocking capacitor may be required to comply with medical regulations or standards, for example. While failure in single-capacitor designs is relatively uncommon, such failure may nonetheless have negative consequences. In one form, a second blocking capacitor **850-2** may be provided in series with the blocking capacitor **850-1**, with current leakage from a point between the blocking capacitors **850-1**, **850-2** being monitored by, for example, an ADC circuit **852** for sampling a voltage induced by leakage current. The samples may be received by the logic circuit **842**, for example. Based changes in the leakage current (as indicated by the voltage samples), the generator **800** may determine when at least one of the blocking capacitors **850-1**, **850-2** has failed, thus providing a benefit over single-capacitor designs having a single point of failure.

[0607] In certain forms, the non-isolated stage **804** may comprise a power supply **854** for delivering DC power at a suitable voltage and current. The power supply may comprise, for example, a 400 W power supply for delivering a 48 VDC system voltage. The power supply **854** may further comprise one or more DC/DC voltage converters **856** for receiving the output of the power supply to generate DC outputs at the voltages and currents required by the various components of the generator **800**. As discussed above in connection with the controller **838**, one or more of the DC/DC voltage converters **856** may receive an input from the controller **838** when activation of the “on/off” input device by a user is detected by the controller **838** to enable operation of, or wake, the DC/DC voltage converters **856**.

[0608] FIG. 21 illustrates an example of a generator **900**, which is one form of the generator **800** (FIG. 20). The generator **900** is configured to deliver multiple energy modalities to a surgical instrument. The generator **900** provides RF and ultrasonic signals for delivering energy to a surgical instrument either independently or simultaneously. The RF and ultrasonic signals may be provided alone or in combination and may be provided simultaneously. As noted above, at least one generator output can deliver multiple energy modalities (e.g., ultrasonic, bipolar or monopolar RF, irreversible and/or reversible electroporation, and/or microwave energy, among others) through a single port, and these signals can be delivered separately or simultaneously to the end effector to treat tissue. The gen-

erator **900** comprises a processor **902** coupled to a waveform generator **904**. The processor **902** and waveform generator **904** are configured to generate a variety of signal waveforms based on information stored in a memory coupled to the processor **902**, not shown for clarity of disclosure. The digital information associated with a waveform is provided to the waveform generator **904** which includes one or more DAC circuits to convert the digital input into an analog output. The analog output is fed to an amplifier **1106** for signal conditioning and amplification. The conditioned and amplified output of the amplifier **906** is coupled to a power transformer **908**. The signals are coupled across the power transformer **908** to the secondary side, which is in the patient isolation side. A first signal of a first energy modality is provided to the surgical instrument between the terminals labeled ENERGY1 and RETURN. A second signal of a second energy modality is coupled across a capacitor **910** and is provided to the surgical instrument between the terminals labeled ENERGY2 and RETURN. It will be appreciated that more than two energy modalities may be output and thus the subscript “n” may be used to designate that up to n ENERGYn terminals may be provided, where n is a positive integer greater than 1. It also will be appreciated that up to “n” return paths RETURNn may be provided without departing from the scope of the present disclosure.

**[0609]** A first voltage sensing circuit **912** is coupled across the terminals labeled ENERGY1 and the RETURN path to measure the output voltage therebetween. A second voltage sensing circuit **924** is coupled across the terminals labeled ENERGY2 and the RETURN path to measure the output voltage therebetween. A current sensing circuit **914** is disposed in series with the RETURN leg of the secondary side of the power transformer **908** as shown to measure the output current for either energy modality. If different return paths are provided for each energy modality, then a separate current sensing circuit should be provided in each return leg. The outputs of the first and second voltage sensing circuits **912**, **924** are provided to respective isolation transformers **916**, **922** and the output of the current sensing circuit **914** is provided to another isolation transformer **918**. The outputs of the isolation transformers **916**, **928**, **922** in the on the primary side of the power transformer **908** (non-patient isolated side) are provided to a one or more ADC circuit **926**. The digitized output of the ADC circuit **926** is provided to the processor **902** for further processing and computation. The output voltages and output current feedback information can be employed to adjust the output voltage and current provided to the surgical instrument and to compute output impedance, among other parameters. Input/output communications between the processor **902** and patient isolated circuits is provided through an interface circuit **920**. Sensors also may be in electrical communication with the processor **902** by way of the interface circuit **920**.

**[0610]** In one aspect, the impedance may be determined by the processor **902** by dividing the output of either the first voltage sensing circuit **912** coupled across the terminals labeled ENERGY1/RETURN or the second voltage sensing circuit **924** coupled across the terminals labeled ENERGY2/RETURN by the output of the current sensing circuit **914** disposed in series with the RETURN leg of the secondary side of the power transformer **908**. The outputs of the first and second voltage sensing circuits **912**, **924** are provided to separate isolations transformers **916**, **922** and the output of the current sensing circuit **914** is provided to another isola-

tion transformer **916**. The digitized voltage and current sensing measurements from the ADC circuit **926** are provided the processor **902** for computing impedance. As an example, the first energy modality ENERGY1 may be ultrasonic energy and the second energy modality ENERGY2 may be RF energy. Nevertheless, in addition to ultrasonic and bipolar or monopolar RF energy modalities, other energy modalities include irreversible and/or reversible electroporation and/or microwave energy, among others. Also, although the example illustrated in FIG. **21** shows a single return path RETURN may be provided for two or more energy modalities, in other aspects, multiple return paths RETURNn may be provided for each energy modality ENERGYn. Thus, as described herein, the ultrasonic transducer impedance may be measured by dividing the output of the first voltage sensing circuit **912** by the current sensing circuit **914** and the tissue impedance may be measured by dividing the output of the second voltage sensing circuit **924** by the current sensing circuit **914**.

**[0611]** As shown in FIG. **21**, the generator **900** comprising at least one output port can include a power transformer **908** with a single output and with multiple taps to provide power in the form of one or more energy modalities, such as ultrasonic, bipolar or monopolar RF, irreversible and/or reversible electroporation, and/or microwave energy, among others, for example, to the end effector depending on the type of treatment of tissue being performed. For example, the generator **900** can deliver energy with higher voltage and lower current to drive an ultrasonic transducer, with lower voltage and higher current to drive RF electrodes for sealing tissue, or with a coagulation waveform for spot coagulation using either monopolar or bipolar RF electrosurgical electrodes. The output waveform from the generator **900** can be steered, switched, or filtered to provide the frequency to the end effector of the surgical instrument. The connection of an ultrasonic transducer to the generator **900** output would be preferably located between the output labeled ENERGY1 and RETURN as shown in FIG. **21**. In one example, a connection of RF bipolar electrodes to the generator **900** output would be preferably located between the output labeled ENERGY2 and RETURN. In the case of monopolar output, the preferred connections would be active electrode (e.g., pencil or other probe) to the ENERGY2 output and a suitable return pad connected to the RETURN output.

**[0612]** Additional details are disclosed in U.S. Patent Application Publication No. 2017/0086914, titled TECHNIQUES FOR OPERATING GENERATOR FOR DIGITALLY GENERATING ELECTRICAL SIGNAL WAVEFORMS AND SURGICAL INSTRUMENTS, which published on Mar. 30, 2017, which is herein incorporated by reference in its entirety.

**[0613]** As used throughout this description, the term “wireless” and its derivatives may be used to describe circuits, devices, systems, methods, techniques, communications channels, etc., that may communicate data through the use of modulated electromagnetic radiation through a non-solid medium. The term does not imply that the associated devices do not contain any wires, although in some aspects they might not. The communication module may implement any of a number of wireless or wired communication standards or protocols, including but not limited to Wi-Fi (IEEE 802.11 family), WiMAX (IEEE 802.16 family), IEEE 802.20, long term evolution (LTE), Ev-DO, HSPA+, HSDPA+, HSUPA+, EDGE, GSM, GPRS, CDMA,

TDMA, DECT, Bluetooth, Ethernet derivatives thereof, as well as any other wireless and wired protocols that are designated as 3G, 4G, 5G, and beyond. The computing module may include a plurality of communication modules. For instance, a first communication module may be dedicated to shorter range wireless communications such as Wi-Fi and Bluetooth and a second communication module may be dedicated to longer range wireless communications such as GPS, EDGE, GPRS, CDMA, WiMAX, LTE, Ev-DO, and others.

**[0614]** As used herein a processor or processing unit is an electronic circuit which performs operations on some external data source, usually memory or some other data stream. The term is used herein to refer to the central processor (central processing unit) in a system or computer systems (especially systems on a chip (SoCs)) that combine a number of specialized “processors.”

**[0615]** As used herein, a system on a chip or system on chip (SoC or SOC) is an integrated circuit (also known as an “IC” or “chip”) that integrates all components of a computer or other electronic systems. It may contain digital, analog, mixed-signal, and often radio-frequency functions—all on a single substrate. A SoC integrates a microcontroller (or microprocessor) with advanced peripherals like graphics processing unit (GPU), Wi-Fi module, or coprocessor. A SoC may or may not contain built-in memory.

**[0616]** As used herein, a microcontroller or controller is a system that integrates a microprocessor with peripheral circuits and memory. A microcontroller (or MCU for microcontroller unit) may be implemented as a small computer on a single integrated circuit. It may be similar to a SoC; an SoC may include a microcontroller as one of its components. A microcontroller may contain one or more core processing units (CPUs) along with memory and programmable input/output peripherals. Program memory in the form of Ferroelectric RAM, NOR flash or OTP ROM is also often included on chip, as well as a small amount of RAM. Microcontrollers may be employed for embedded applications, in contrast to the microprocessors used in personal computers or other general purpose applications consisting of various discrete chips.

**[0617]** As used herein, the term controller or microcontroller may be a stand-alone IC or chip device that interfaces with a peripheral device. This may be a link between two parts of a computer or a controller on an external device that manages the operation of (and connection with) that device.

**[0618]** Any of the processors or microcontrollers described herein, may be implemented by any single core or multicore processor such as those known under the trade name ARM Cortex by Texas Instruments. In one aspect, the processor may be an LM4F230H5QR ARM Cortex-M4F Processor Core, available from Texas Instruments, for example, comprising on-chip memory of 256 KB single-cycle flash memory, or other non-volatile memory, up to 40 MHz, a prefetch buffer to improve performance above 40 MHz, a 32 KB single-cycle serial random access memory (SRAM), internal read-only memory (ROM) loaded with StellarisWare® software, 2 KB electrically erasable programmable read-only memory (EEPROM), one or more pulse width modulation (PWM) modules, one or more quadrature encoder inputs (QEI) analog, one or more 12-bit Analog-to-Digital Converters (ADC) with 12 analog input channels, details of which are available for the product datasheet.

**[0619]** In one aspect, the processor may comprise a safety controller comprising two controller-based families such as TMS570 and RM4x known under the trade name Hercules ARM Cortex R4, also by Texas Instruments. The safety controller may be configured specifically for IEC 61508 and ISO 26262 safety critical applications, among others, to provide advanced integrated safety features while delivering scalable performance, connectivity, and memory options.

**[0620]** Modular devices include the modules (as described in connection with FIGS. 3 and 9, for example) that are receivable within a surgical hub and the surgical devices or instruments that can be connected to the various modules in order to connect or pair with the corresponding surgical hub. The modular devices include, for example, intelligent surgical instruments, medical imaging devices, suction/irrigation devices, smoke evacuators, energy generators, ventilators, insufflators, and displays. The modular devices described herein can be controlled by control algorithms. The control algorithms can be executed on the modular device itself, on the surgical hub to which the particular modular device is paired, or on both the modular device and the surgical hub (e.g., via a distributed computing architecture). In some exemplifications, the modular devices’ control algorithms control the devices based on data sensed by the modular device itself (i.e., by sensors in, on, or connected to the modular device). This data can be related to the patient being operated on (e.g., tissue properties or insufflation pressure) or the modular device itself (e.g., the rate at which a knife is being advanced, motor current, or energy levels). For example, a control algorithm for a surgical stapling and cutting instrument can control the rate at which the instrument’s motor drives its knife through tissue according to resistance encountered by the knife as it advances.

#### Long Distance Communication and Condition Handling of Devices and Data

**[0621]** Surgical procedures are performed by different surgeons at different locations, some with much less experience than others. For a given surgical procedure, there are many parameters that can be varied to attempt to realize a desired outcome. For example, for a given surgical procedure which utilizes energy supplied by a generator, the surgeon often relies on experience alone for determining which mode of energy to utilize, which level of output power to utilize, the duration of the application of the energy, etc., in order to attempt to realize the desired outcome. To increase the likelihood of realizing desired outcomes for a plurality of different surgical procedures, each surgeon should be provided with best practice recommendations which are based on important relationships identified within large, accurate data sets of information associated with multiple surgical procedures performed in multiple locations over time. However, there are many ways that such data sets can be rendered compromised, inaccurate, and/or insecure, thereby calling into question the applicability of the best practice recommendations derived therefrom. For example, for data sent from a source to a cloud-based system, the data can be lost while in transit to the cloud-based system, the data can be corrupted while in transit to the cloud-based system, the confidentiality of the data can be comprised while in transit to the cloud-based system, and/or the content of the data can be altered while in transit to the cloud-based system.



**[0622]** A plurality of operating rooms located in multiple locations can each be equipped with a surgical hub. When a given surgical procedure is performed in a given operating room, the surgical hub can receive data associated with the surgical procedure and communicate the data to a cloud-based system. Over time, the cloud-based system will receive large data sets of information associated with the surgeries. The data can be communicated from the surgical hubs to the cloud-based system in a manner which allows for the cloud-based system to (1) verify the authenticity of the communicated data, (2) authenticate each of the respective surgical hubs which communicated the data, and (3) trace the paths the data followed from the respective surgical hubs to the cloud-based system.

**[0623]** Accordingly, in one aspect, the present disclosure provides a surgical hub for transmitting generator data associated with a surgical procedure to a cloud-based system communicatively coupled to a plurality of surgical hubs. The surgical hub comprises a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to receive data from a generator, encrypt the data, generate a message authentication code (MAC) based on the data, generate a datagram comprising the encrypted data, the generated MAC, a source identifier, and a destination identifier, and transmit the datagram to a cloud-based system. The data is structured into a data packet comprising at least two of the following fields: a field that indicates the source of the data, a unique time stamp, a field indicating an energy mode of the generator, a field indicating the power output of the generator, and a field indicating a duration of the power output of the generator. The datagram allows for the cloud-based system to decrypt the encrypted data of the transmitted datagram, verify integrity of the data based on the MAC, authenticate the surgical hub as the source of the datagram, and validate a transmission path followed by the datagram between the surgical hub and the cloud-based system.

In various aspects, the present disclosure provides a control circuit to transmit generator data associated with a surgical procedure to a cloud-based system communicatively coupled to a plurality of surgical hubs, as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer-readable instructions which, when executed, causes a machine to transmit generator data associated with a surgical procedure to a cloud-based system communicatively coupled to a plurality of surgical hubs, as described above.

**[0624]** In another aspect, the present disclosure provides a cloud-based system communicatively coupled to a plurality of surgical hubs. Each surgical hub is configured to transmit generator data associated with a surgical procedure to the cloud-based system. The cloud-based system comprises a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to receive a datagram generated by a surgical hub, decrypt the encrypted generator data of the received datagram, verify integrity of the generator data based on the MAC, authenticate the surgical hub as the source of the datagram, and validate a transmission path followed by the datagram between the surgical hub and the cloud-based system. The datagram comprises generator data captured from a generator associated with the surgical hub, a MAC generated by the surgical hub based on the generator data, a source identifier, and a destination identifier. The generator data has been

encrypted by the surgical hub. The encrypted generator data has been structured into a data packet comprising at least two of the following fields: a field that indicates the source of the data, a unique time stamp, a field indicating an energy mode, a field indicating power output, and a field indicating a duration of applied power.

**[0625]** In various aspects, the present disclosure provides a control circuit to transmit generator data associated with a surgical procedure to the cloud-based system. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer-readable instructions which, when executed, causes a machine to transmit generator data associated with a surgical procedure to the cloud-based system.

**[0626]** In another aspect, the present disclosure provides a method, comprising capturing data from a combination generator of a surgical hub during a surgical procedure, wherein the combination generator is configured to supply two or more different modes of energy. Encrypting the captured generator data, generating a MAC based on the captured generator data, generating a datagram comprising the encrypted generator data, the MAC, a source identifier, and a destination identifier, and communicating the datagram from the surgical hub to a cloud-based system. The datagram allows for the cloud-based system to authenticate integrity of the communicated generator data, authenticate the surgical hub as a source of the datagram, and determine a communication path followed by the datagram between the surgical hub and the cloud-based system.

**[0627]** By sending captured generator data from a plurality of different surgical hubs to a cloud-based system, the cloud-based system is able to quickly build large data sets of information associated with multiple surgical procedures performed in multiple locations over time. Furthermore, due to the composition of the respective datagrams, for a given datagram, the cloud-based system is able to determine whether the datagram was originally sent by one of the surgical hubs (source validation), thereby providing an indication that the generator data received at the cloud-based system is legitimate data. For the given datagram, the cloud-based system is also able to determine whether the generator data received at the cloud-based system is identical to the generator data sent by the given surgical hub (data integrity), thereby allowing for the authenticity of the received generator data to be verified. Additionally, for the given datagram, the cloud-based system is also able to re-trace the communication path followed by the datagram, thereby allowing for enhanced troubleshooting if a datagram received by the cloud-based system was originally sent from a device other than the surgical hubs and/or if the content of the datagram was altered while in transit to the cloud-based system. Notably, the present disclosure references generator data in particular. Here, the present disclosure should not be limited as being able to process only generator data. For example, the surgical hub **206** and/or the cloud-based system **205** may process data received from any component (e.g., imaging module **238**, generator module **240**, smoke evacuator module **226**, suction/irrigation module **228**, communication module **230**, processor module **232**, storage array **234**, smart device/instrument **235**, non-contact sensor module **242**, robot hub **222**, a non-robotic surgical hub **206**, wireless smart device/instrument **235**, visualization system **208**) of the surgical system **202** that is coupled to the surgical hub **206** and/or data from any devices (e.g., endoscope **239**,

energy device **241**) coupled to/through such components (e.g., see FIGS. 9-10), in a similar manner as discussed herein.

**[0628]** Unfortunately, the outcome of a surgical procedure is not always optimal. For example, a failure event such as a surgical device failure, an unwanted tissue perforation, an unwanted post-operative bleeding, or the like can occur. The occurrence of a failure event can be attributed to any of a variety of different people and devices, including one or more surgeons, one or more devices associated with the surgery, a condition of the patient, and combinations thereof. When a given failure event occurs, it is not always clear regarding who or what caused the failure event or how the occurrence of the failure event can be mitigated in connection with a future surgery.

**[0629]** During a given surgical procedure, a large amount of data associated with the surgical procedure can be generated and captured. All of the captured data can be communicated to a surgical hub, and the captured data can be time-stamped either before or after being received at the surgical hub. When a failure event associated with the surgical procedure is detected and/or identified, it can be determined which of the captured data is associated with the failure event and/or which of the captured data is not associated with the failure event. In making this determination, the failure event can be defined to include a period of time prior to the detection/identification of the failure event. Once the determination is made regarding the captured data associated with the failure event, the surgical hub can separate the captured data associated with the failure event from all other captured data, and the captured data can be separated based on tagging, flagging, or the like. The captured data associated with the failure event can then be chronologized based on the time-stamping and the defined time period applicable to the failure event. The chronologized captured data can then be communicated to a cloud-based system on a prioritized basis for analysis, where the prioritized basis is relative to the captured data which is not associated with the failure event. Whether or not the analysis identifies a device associated with the surgical procedure as the causation of the failure event, the surgical hub can tag the device for removal of the device from future use, further analysis of the device, and/or to return the device to the manufacturer.

**[0630]** When a given surgical procedure is performed, a large amount of data associated with the surgical procedure can be generated and captured. All of the captured data can be communicated to a surgical hub, where the information can be stripped of all "personal" associations. The captured data can be time-stamped before being received at the surgical hub, after being received at the surgical hub, before being stripped of the "personal" associations, or after being stripped of the "personal" associations. The surgical hub can communicate the stripped data to the cloud-based system for subsequent analysis. Over time, the cloud-based system will receive large data sets of information associated with the surgeries.

Accordingly, in one aspect, the present disclosure provides a surgical hub for prioritizing surgical data associated with a surgical procedure to a cloud-based system communicatively coupled to a plurality of surgical hubs. The surgical hub comprises a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to capture surgical data, wherein the surgical data

comprises data associated with a surgical device, time-stamp the captured surgical data, identify a failure event, identify a time period associated with the failure event, isolate failure event surgical data from surgical data not associated with the failure event based on the identified time period, chronologize the failure event surgical data by time-stamp, encrypt the chronologized failure event surgical data, generate a datagram comprising the encrypted failure event surgical data, and transmit the datagram to a cloud-based system. The datagram is structured to include a field which includes a flag that prioritizes the encrypted failure event surgical data over other encrypted data of the datagram. The datagram allows for the cloud-based system to decrypt the encrypted failure event surgical data, focus analysis on the failure event surgical data rather than surgical data not associated with the failure event, and flag the surgical device associated with the failure event for at least one of the following: removal from an operating room, return to a manufacturer, or future inoperability in the cloud-based system.

**[0631]** In various aspects, the present disclosure provides a control circuit to prioritize surgical data associated with a surgical procedure to a cloud-based system communicatively coupled to a plurality of surgical hubs. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer-readable instructions which, when executed, causes a machine to prioritize surgical data associated with a surgical procedure to a cloud-based system communicatively coupled to a plurality of surgical hubs.

**[0632]** In another aspect, the present disclosure provides a method, comprising capturing data during a surgical procedure, communicating the captured data to a surgical hub, time-stamping the captured data, identifying a failure event associated with the surgical procedure, determining which of the captured data is associated with the failure event, separating the captured data associated with the failure event from all other captured data, chronologizing the captured data associated with the failure event, and communicating the chronologized captured data to a cloud-based system on a prioritized basis.

**[0633]** By capturing the large amount of data associated with the surgical procedure, and with having the captured data time-stamped, the portion of the captured data which is relevant to the detected/identified failure event can be more easily isolated from all of the other captured data, thereby allowing for a more focused subsequent analysis on just the relevant captured data. The data associated with the failure event can then be chronologized (this requires less processing power than chronologizing all of the captured data), thereby allowing for the events leading up to the detection/identification of the failure event to be more easily considered during the subsequent analysis of the failure event. The chronologized data can then be communicated to the cloud-based system (this requires less communication resources than communicating all of the captured data at the same time) on a prioritized basis, thereby allowing for the focused subsequent analysis of the fault event to be performed by the cloud-based system in a more time-sensitive manner.

**[0634]** To help ensure that the best practice recommendations are developed based on accurate data, it would be desirable to ensure that the generator data received at the cloud-based system is the same as the generator data communicated to the cloud-based system. Also, to help to be able to determine the cause of a failure event as quickly as

possible, it would be desirable to ensure that surgical data associated with the failure event is communicated to the cloud-based system in a prioritized manner (relative to surgical data not associated with the failure event) so that analysis of the surgical data can be performed in an expedited manner.

[0635] Aspects of a system and method for communicating data associated with a surgical procedure are described herein. As shown in FIG. 9, various aspects of the computer implemented interactive surgical system 200 includes a device/instrument 235, a generator module 240, a modular control tower 236, and a cloud-based system 205. As shown in FIG. 10, the device/instrument 235, the generator module 240, and the modular control tower 236 are components/portions of a surgical hub 206.

[0636] In various aspects, the generator module 240 of the surgical hub 206 can supply radio-frequency energy such as monopolar radio-frequency energy, bipolar radio-frequency energy, and advanced bipolar energy and/or ultrasonic energy to a device/instrument 235 for use in a surgical procedure. Thus, the generator module 240 may be referred to as a combination generator. An example of such a combination generator is shown in FIG. 22, where the combination generator 3700 is shown as including a monopolar module 3702, a bipolar module 3704, an advanced bipolar module 3706, and an ultrasound module 3708. When utilized during a surgical procedure, the respective energy modules (e.g., 3702, 3704, 3706, and/or 3708) of the combination generator 3700 can provide generator data such as type of energy supplied to the device instrument (e.g., radio-frequency energy, ultrasound energy, radio-frequency energy and ultrasound energy), type of radio-frequency energy (e.g., monopolar, bipolar, advanced bipolar), frequency, power output, duration, etc., to the data communication module 3710 of the combination generator 3700.

[0637] FIG. 23 illustrates various aspects of a method of capturing data from a combination generator 3700 and communicating the captured generator data to a cloud-based system 205. Notably, as discussed herein, the present disclosure should not be limited to processing generator data. As such, the method of FIG. 23 similarly extends to other types of data received from other components coupled to the surgical hub 206 (e.g., imaging module data, smoke evacuator data, suction/irrigation data, device/instrument data). The method comprises (1) capturing 3712 data from a combination generator 3700 of a surgical hub 206 during a surgical procedure, wherein the combination generator 3700 is configured to supply two or more different modes of energy; (2) encrypting 3714 the captured generator data; (3) generating 3716 a MAC based on the captured generator data; (4) generating 3718 a datagram comprising the encrypted generator data, the MAC, a source identifier, and a destination identifier; and (5) communicating 3720 the datagram from the surgical hub 206 to a cloud-based system 205, wherein the datagram allows for the cloud-based system 205 to (i) authenticate integrity of the communicated generator data, (ii) authenticate the surgical hub as a source of the datagram, and (iii) determine a communication path followed by the datagram between the surgical hub 206 and the cloud-based system 205.

[0638] More specifically, once the generator data is received at the data communication module 3710 of the combination generator 3700, the generator data can be communicated to the modular communication hub 203 of

the surgical hub 206 for subsequent communication to the cloud-based system 205. The data communication module 3710 can communicate the generator data to the modular communication hub 203 serially over a single communication line or in parallel over a plurality of communication lines, and such communication can be performed in real time or near real time. Alternatively, such communication can be performed in batches.

[0639] According to various aspects, prior to communicating the generator data to the modular communication hub 203, a component of the combination generator 3700 (e.g., the data communication module 3710) can organize the generator data into data packets. An example of such a data packet is shown in FIG. 24, where the data packet 3722 includes a preamble 3724 or self-describing data header which defines what the data is (e.g., combination generator data—CGD) and fields which indicate where the generator data came from [e.g., combination generator ID number 3726—(e.g., 017), a unique time stamp 3728 (e.g., 08:27:16), the energy mode utilized 3730 (e.g., RF, U, RF+U), the type of radio-frequency energy or radio frequency mode 3732 (e.g., MP, BP, ABP), the frequency 3734 (e.g., 500 KHz), the power output 3736 (e.g., 30 watts), the duration of applied power 3738 (e.g., 45 milliseconds), and an authentication/identification certificate of the data point 3740 (e.g., 01101011001011)]. The example data packet 3722 may be considered a self-describing data packet, and the combination generator 3700 and other intelligent devices (e.g., the surgical hub 206) can use the self-describing data packets to minimize data size and data-handling resources. Again, as discussed herein, the present disclosure should not be limited to processing generator data received from a combination generator 3700. As such, the data packet 3722 of FIG. 24 similarly extends to other types of data received from other components coupled to the surgical hub 206. In one aspect, the data packet 3722 may comprise data associated with endoscope 239 (e.g., image data) received from a component of the imaging module 238. In another aspect, the data packet 3722 may comprise data associated with an evacuation system (e.g., pressures, particle counts, flow rates, motor speeds) received from a component of the smoke evacuator module 226. In yet another aspect, the data packet 3722 may comprise data associated with a device/instrument (e.g., temperature sensor data, firing data, sealing data) received from a component of the device/instrument 235. In various other aspects, the data packet 3722 may similarly comprise data received from other components coupled to the surgical hub 206 (e.g., suction/irrigation module 228, non-contact sensor module 242)

[0640] Additionally, the data communication module 3710 can compress the generator data and/or encrypt the generator data prior to communicating the generator data to the modular communication hub 203. The specific method of compressing and/or encrypting can be the same as or different from the compressing and/or encrypting which may be performed by the surgical hub 206 as described in more detail below.

[0641] The modular communication hub 203 can receive the generator data communicated from the combination generator 3700 (e.g., via the data communication module 3710), and the generator data can be subsequently communicated to the cloud-based system 205 (e.g., through the Internet). According to various aspects, the modular communication hub 203 can receive the generator data through

a hub/switch 207/209 of the modular communication hub 203 (See FIG. 10), and the generator data can be communicated to the cloud-based system 205 by a router 211 of the modular communication hub 203 (See FIG. 10). The generator data may be communicated in real time, near real time, or in batches to the cloud-based system 205 or may be stored at the surgical hub 206 prior to being communicated to the cloud-based system 205. The generator data can be stored, for example, at the storage array 234 or at the memory 249 of the computer system 210 of the surgical hub 206.

[0642] In various aspects, for instances where the generator data received at the modular communication hub 203 is not encrypted, prior to the received generator data being communicated to the cloud-based system 205, the generator data is encrypted to help ensure the confidentiality of the generator data, either while it is being stored at the surgical hub 206 or while it is being transmitted to the cloud 204 using the Internet or other computer networks. According to various aspects, a component of the surgical hub 206 utilizes an encryption algorithm to convert the generator data from a readable version to an encoded version, thereby forming the encrypted generator data. The component of the surgical hub 206 which utilizes/executes the encryption algorithm can be, for example, the processor module 232, the processor 244 of the computer system 210, and/or combinations thereof. The utilized/executed encryption algorithm can be a symmetric encryption algorithm and/or an asymmetric encryption algorithm.

[0643] Using a symmetric encryption algorithm, the surgical hub 206 would encrypt the generator data using a shared secret (e.g., private key, passphrase, password). In such an aspect, a recipient of the encrypted generator data (e.g., cloud-based system 205) would then decrypt the encrypted generator data using the same shared secret. In such an aspect, the surgical hub 206 and the recipient would need access to and/or knowledge of the same shared secret. In one aspect, a shared secret can be generated/chosen by the surgical hub 206 and securely delivered (e.g., physically) to the recipient before encrypted communications to the recipient.

[0644] Alternatively, using an asymmetric encryption algorithm, the surgical hub 206 would encrypt the generator data using a public key associated with a recipient (e.g., cloud-based system 205). This public key could be received by the surgical hub 206 from a certificate authority that issues a digital certificate certifying the public key as owned by the recipient. The certificate authority can be any entity trusted by the surgical hub 206 and the recipient. In such an aspect, the recipient of the encrypted generator data would then decrypt the encrypted generator data using a private key (i.e., known only by the recipient) paired to the public key used by the surgical hub 206 to encrypt the generator data. Notably, in such an aspect, the encrypted generator data can only be decrypted using the recipient's private key.

[0645] According to aspects of the present disclosure, components (e.g., surgical device/instrument 235, energy device 241, endoscope 239) of the surgical system 202 are associated with unique identifiers, which can be in the form of serial numbers. As such, according to various aspects of the present disclosure, when a component is coupled to a surgical hub 206, the component may establish a shared secret with the surgical hub 206 using the unique identifier of the coupled component as the shared secret. Further, in

such an aspect, the component may derive a checksum value by applying a checksum function/algorithm to the unique identifier and/or other data being communicated to the surgical hub 206. Here, the checksum function/algorithm is configured to output a significantly different checksum value if there is a modification to the underlying data.

[0646] In one aspect, the component may initially encrypt the unique identifier of a coupled component using a public key associated with the surgical hub (e.g., received by the component from the surgical hub 206 upon/after connection) and communicate the encrypted unique identifier to the surgical hub 206. In other aspects, the component may encrypt the unique identifier and the derived checksum value of a coupled component using a public key associated with the surgical hub 206 and communicate the encrypted unique identifier and linked/associated checksum value to the surgical hub 206.

[0647] In yet other aspects, the component may encrypt the unique identifier and a checksum function/algorithm using a public key associated with the surgical hub 206 and communicate the encrypted unique identifier and the checksum function/algorithm to the surgical hub 206. In such aspects, the surgical hub 206 would then decrypt the encrypted unique identifier or the encrypted unique identifier and the linked/associated checksum value or the encrypted unique identifier and the checksum function/algorithm using a private key (i.e., known only by the surgical hub 206) paired to the public key used by the component to encrypt the unique identifier.

[0648] Since the encrypted unique identifier can only be decrypted using the surgical hub's 206 private key and the private key is only known by the surgical hub, this is a secure way to communicate a shared secret (e.g., the unique identifier of the coupled component) to the surgical hub 206. Further, in aspects where a checksum value is linked to/associated with the unique identifier, the surgical hub 206 may apply the same checksum function/algorithm to the decrypted unique identifier to generate a validating checksum value. If the validating checksum value matches the decrypted checksum value, the integrity of the decrypted unique identifier is further verified. Further, in such aspects, with a shared secret established, the component can encrypt future communications to the surgical hub 206, and the surgical hub 206 can decrypt the future communications from the component using the shared secret (e.g., the unique identifier of the coupled component). Here, according to various aspects, a checksum value may be derived for and communicated with each communication between the component and the surgical hub 206 (e.g., the checksum value based on the communicated data or at least a designated portion thereof). Here, a checksum function/algorithm (e.g., known by the surgical hub 206 and/or component or communicated when establishing the shared secret between the surgical hub 206 and the component as described above) may be used to generate validating checksum values for comparison with communicated checksum values to further verify the integrity of communicated data in each communication.

[0649] Notably, asymmetric encryption algorithms may be complex and may require significant computational resources to execute each communication. As such, establishing the unique identifier of the coupled component as the shared secret is not only quicker (e.g., no need to generate a shared secret using a pseudorandom key generator) but

also increases computational efficiency (e.g., enables the execution of faster, less complex symmetric encryption algorithms) for all subsequent communications. In various aspects, this established shared secret may be utilized by the component and surgical hub 206 until the component is decoupled from the surgical hub (e.g., surgical procedure ended).

[0650] According to other aspects of the present disclosure, components (e.g., surgical device/instrument 235, energy device 241, endoscope 239) of the surgical system 202 may comprise sub-components (e.g., handle, shaft, end effector, cartridge) each associated with its own unique identifier. As such, according to various aspects of the present disclosure, when a component is coupled to the surgical hub 206, the component may establish a shared secret with the surgical hub 206 using a unique compilation/string (e.g., ordered or random) of the unique identifiers associated with the sub-components that combine to form the coupled component. In one aspect, the component may initially encrypt the unique compilation/string of the coupled component using a public key associated with the surgical hub 206 and communicate the encrypted unique compilation/string to the surgical hub 206. In such an aspect, the surgical hub 206 would then decrypt the encrypted unique compilation/string using a private key (i.e., known only by the surgical hub 206) paired to the public key used by the component to encrypt the unique compilation/string. Since the encrypted unique compilation/string can only be decrypted using the surgical hub's 206 private key and the private key is only known by the surgical hub 206, this is a secure way to communicate a shared secret (e.g., the unique compilation/string of the coupled component) to the surgical hub 206. Further, in such an aspect, with a shared secret established, the component can encrypt future communications to the surgical hub 206, and the surgical hub 206 can decrypt the future communications from the component using the shared secret (e.g., the unique compilation/string of the coupled component).

[0651] Again, asymmetric encryption algorithms may be complex and may require significant computational resources to execute each communication. As such, establishing the unique compilation/string of the coupled component (i.e., readily combinable by the component) as the shared secret is not only quicker (e.g., no need to generate a shared secret using a pseudorandom key generator) but also increases computational efficiency (e.g., enables the execution of faster, less complex symmetric encryption algorithms) for all subsequent communications. In various aspects, this established shared secret may be utilized by the component and surgical hub 206 until the component is decoupled from the surgical hub 206 (e.g., surgical procedure ended). Furthermore, in such an aspect, since various sub-components may be reusable (e.g., handle, shaft, end effector) while other sub-components may not be reusable (e.g., end effector, cartridge) each new combination of sub-components that combine to form the coupled component provide a unique compilation/string usable as a shared secret for component communications to the surgical hub 206.

[0652] According to further aspects of the present disclosure, components (e.g., surgical device/instrument 235, energy device 241, endoscope 239) of the surgical system 202 are associated with unique identifiers. As such, according to various aspects of the present disclosure, when a

component is coupled to the surgical hub 206, the surgical hub 206 may establish a shared secret with a recipient (e.g., cloud-based system 205) using the unique identifier of the coupled component. In one aspect, the surgical hub 206 may initially encrypt the unique identifier of a coupled component using a public key associated with the recipient and communicate the encrypted unique identifier to the recipient. In such an aspect, the recipient would then decrypt the encrypted unique identifier using a private key (i.e., known only by the recipient) paired to the public key used by the surgical hub 206 to encrypt the unique identifier. Since the encrypted unique identifier can only be decrypted using the recipient's private key and the private key is only known by the recipient, this is a secure way to communicate a shared secret (e.g., the unique identifier of the coupled component) to the recipient (e.g., cloud-based system). Further in such an aspect, with a shared secret established, the surgical hub 206 can encrypt future communications to the recipient (e.g., cloud-based system 205), and the recipient can decrypt the future communications from the surgical hub 206 using the shared secret (e.g., the unique identifier of the coupled component).

[0653] Notably, asymmetric encryption algorithms may be complex and may require significant computational resources to execute each communication. As such, establishing the unique identifier of the coupled component (i.e., already available to the surgical hub 206) as the shared secret is not only quicker (e.g., no need to generate a shared secret using a pseudorandom key generator) but also increases computational efficiency by, for example, enabling the execution of faster, less complex symmetric encryption algorithms for all subsequent communications. In various aspects, this established shared secret may be utilized by the surgical hub 206 until the component is decoupled from the surgical hub (e.g., surgical procedure ended).

[0654] According to yet further aspects of the present disclosure, components (e.g., surgical device/instrument 235, energy device 241, endoscope 239) of the surgical system 202 may comprise sub-components (e.g., handle, shaft, end effector, cartridge) each associated with its own unique identifier. As such, according to various aspects of the present disclosure, when a component is coupled to the surgical hub 206, the surgical hub 206 may establish a shared secret with a recipient (e.g., cloud-based system 205) using a unique compilation/string (e.g., ordered or random) of the unique identifiers associated with the sub-components that combine to form the coupled component.

[0655] In one aspect, the surgical hub 206 may initially encrypt the unique compilation/string of the coupled component using a public key associated with the recipient and communicate the encrypted unique compilation/string to the recipient. In such an aspect, the recipient would then decrypt the encrypted unique compilation/string using a private key (i.e., known only by the recipient) paired to the public key used by the surgical hub 206 to encrypt the unique compilation/string. Since the encrypted unique compilation/string can only be decrypted using the recipient's private key and the private key is only known by the recipient, this is a secure way to communicate a shared secret (e.g., the unique compilation/string of the coupled component) to the recipient. With a shared secret established, the surgical hub 206 can encrypt future communications to the recipient (e.g., cloud-based system 205), and the recipient can decrypt the future communications from the surgical hub 206 using the

shared secret (e.g., the unique compilation/string of the coupled component). Again, asymmetric encryption algorithms may be complex and may require significant computational resources to execute each communication. As such, establishing the unique compilation/string of the coupled component (i.e., readily combinable by the surgical hub 206) as the shared secret is not only quicker (e.g., no need to generate a shared secret using a pseudorandom key generator) but also increases computational efficiency (e.g., enables the execution of faster, less complex symmetric encryption algorithms) for all subsequent communications.

[0656] In various aspects, this established shared secret may be utilized by the surgical hub 206 until the component is decoupled from the surgical hub (e.g., surgical procedure ended). Furthermore, in such an aspect, since various sub-components may be reusable (e.g., handle, shaft, end effector) while other sub-components may not be reusable (e.g., end effector, cartridge) each new combination of sub-components that combine to form the coupled component provide a unique compilation/string usable as a shared secret for surgical hub 206 communications to the recipient.

[0657] In some aspects, an encrypt-then-MAC (EtM) approach may be utilized to produce the encrypted generator data. An example of this approach is shown in FIG. 25, where the non-encrypted generator data (i.e., the plaintext 3742, e.g., data packet 3722) is first encrypted 3743 (e.g., via key 3746) to produce a ciphertext 3744 (i.e., the encrypted generator data), then a MAC 3745 is produced based on the resulting ciphertext 3744, the key 3746, and a MAC algorithm (e.g., a hash function 3747). More specifically, the ciphertext 3744 is processed through the MAC algorithm using the key 3746. In one aspect similar to symmetric encryption discussed herein, the key 3746 is a secret key accessible/known by the surgical hub 206 and the recipient (e.g., cloud-based system 205). In such an aspect, the secret key is a shared secret associated with/chosen by the surgical hub 206, a shared secret associated with/chosen by the recipient, or a key selected via a pseudorandom key generator. For this approach, as shown generally at 3748, the encrypted generator data (i.e., the ciphertext 3744) and the MAC 3745 would be communicated together to the cloud-based system 205.

[0658] In other aspects, an encrypt-and-MAC (E&M) approach may be utilized to produce the encrypted generator data. An example of this approach is shown in FIG. 26, where the MAC 3755 is produced based on the non-encrypted generator data (i.e., the plaintext 3752, e.g., data packet 3722), a key 3756, and a MAC algorithm (e.g., a hash function 3757). More specifically, the plaintext 3752 is processed through the MAC algorithm using the key 3756. In one aspect similar to symmetric encryption discussed herein, the key 3756 is a secret key accessible/known by the surgical hub 206 and the recipient (e.g., cloud-based system 205). In such an aspect, the secret key is a shared secret associated with/chosen by the surgical hub 206, a shared secret associated with/chosen by the recipient, or a key selected via a pseudorandom key generator. Further, in such an aspect, the non-encrypted generator data (i.e., the plaintext 3752, e.g., data packet 3722) is encrypted 3753 (e.g., via key 3756) to produce a ciphertext 3754. For this approach, as shown generally at 3758, the MAC 3755 (i.e., produced based on the non-encrypted generator data) and the encrypted generator data (i.e., the ciphertext 3754) would be communicated together to the cloud-based system 205.

[0659] In yet other aspects, a MAC-then-encrypt (MtE) approach may be utilized to produce the encrypted generator data. An example of this approach is shown in FIG. 27, where the MAC 3765 is produced based on the non-encrypted generator data (i.e., the plaintext 3762), a key 3766, and a MAC algorithm (e.g., a hash function 3767). More specifically, the plaintext 3762 is processed through the MAC algorithm using the key 3766. In one aspect similar to symmetric encryption discussed herein, the key 3766 is a secret key accessible/known by the surgical hub 206 and the recipient (e.g., cloud-based system 205). In such an aspect, the secret key is a shared secret associated with/chosen by the surgical hub 206, a shared secret associated with/chosen by the recipient, or a key selected via a pseudorandom key generator. Next, the non-encrypted generator data (i.e., the plaintext 3762) and the MAC 3765 are together encrypted 3763 (e.g., via key 3766) to produce a ciphertext 3764 based on both. For this approach, as shown generally at 3768, the ciphertext 3764 (i.e., which includes the encrypted generator data and the encrypted MAC 3765) would be communicated to the cloud-based system 205.

[0660] In alternative aspects, the key used to encrypt the non-encrypted generator data (e.g., FIG. 25 and FIG. 26) or the non-encrypted generator data and the MAC (e.g., FIG. 27) may be different from the key (e.g., keys 3746, 3756, 3766) used to produce the MAC. For example, the key used to encrypt the non-encrypted generator data (e.g., FIG. 25 and FIG. 26) or the non-encrypted generator data and the MAC (e.g., FIG. 27) may be a different shared secret or a public key associated with the recipient.

[0661] In lieu of utilizing the MAC to provide for a subsequent assurance of data integrity to the cloud-based system 205, according to other aspects, the surgical hub 206 can utilize a digital signature to allow the cloud-based system 205 to subsequently authenticate integrity of the communicated generator data. For example, the processor module 232 and/or the processor 244 of the computer system 210 can utilize one or more algorithms to generate a digital signature associated with the generator data, and the cloud-based system 205 can utilize an algorithm to determine the authenticity of the received generator data. The algorithms utilized by the processor module 232 and/or the processor 244 of the computer system 210 can include: (1) a key generation algorithm that selects a private key uniformly at random from a set of possible private keys, where the key generation algorithm outputs the private key and a corresponding public key; and (2) a signing algorithm that, given the generator data and a private key, produces a digital signature associated with the generator data. The cloud-based system 205 can utilize a signature verifying algorithm that, given the received generator data, public key, and digital signature, can accept the received generator data as authentic if the digital signature is determined to be authentic or consider the generator data to be compromised or altered if the digital signature is not determined to be authentic.

[0662] According to other aspects of the present disclosure, the surgical hub 206 can utilize a commercial authentication program (e.g., Secure Hash Algorithm, SHA-2 comprising SHA-256) to provide for a subsequent assurance of data integrity of the communicated generator data to the cloud-based system 205.

[0663] After the generator data has been encrypted (e.g., via EtM, E&M, MtE), a component of the surgical hub 206

can communicate the encrypted generator data to the cloud-based system 205. The component of the surgical hub 206 which communicates the encrypted generator data to the cloud-based system 205 can be, for example, the processor module 232, a hub/switch 207/209 of the modular communication hub 203, the router 211 of the modular communication hub 203, the communication module 247 of the computer system 210, etc.

[0664] According to various aspects, the communication of the encrypted generator data through the Internet can follow an IP which: (1) defines datagrams that encapsulate the encrypted generator data to be delivered and/or (2) defines addressing methods that are used to label the datagram with source and destination information. A high-level representation of an example datagram 3770 is shown in FIG. 28, where the datagram 3770 includes a header 3772 and a payload 3774, and in other aspects also may include a trailer (not shown). A more detailed representation of an example datagram 3780 is shown in FIG. 29, where the header 3782 can include fields for information such as, for example, the IP address of the source 3786 which is sending the datagram (e.g., the router 211 of the modular communication hub 203), the IP address of the destination 3788 which is to receive the datagram (e.g., the cloud 204 and/or the remote server 213 associated with the cloud-based system 205), a type of service designation (not shown), a header length 3790, a payload length 3792, and a checksum value 3794. In such an aspect, the surgical hub 206 may further apply a checksum function/algorithm to the non-encrypted generator data (i.e., the plaintext 3742, e.g., data packet 3722) or at least a portion of the non-encrypted generator data (e.g., combination generator ID 3726) to derive the checksum value 3794. Here, the checksum function/algorithm is configured to output a significantly different checksum value if there is any modification (e.g., even a slight change) to the underlying data (e.g., generator data). After decryption of the encrypted generator data by its recipient (e.g., cloud-based system 205), the recipient may apply the same checksum function/algorithm to the decrypted generator data to generate a validating checksum value. If the validating checksum value matches the checksum value 3794 (i.e., stored in the header 3782 of the received datagram 3780), the integrity of the received generator data is further verified. The payload 3784 may include the encrypted generator data 3796 and can also include padding 3798 if the encrypted generator data 3796 is less than a specified payload length. Notably, the communicated encrypted generator data 3796 may comprise a MAC as discussed in FIGS. 25, 26, and 27 above (e.g., references 3748, 3758, and 3768, respectively). In some aspects, the header 3782 can further include a specific path the datagram is to follow when the datagram is communicated from the surgical hub 206 to the cloud-based system 205 (e.g., from IP address of the source, to IP address of at least one intermediate network component (e.g., specified routers, specified servers), to IP address of the destination).

[0665] According to various aspects, prior to the generator data being encrypted, the generator data can be time-stamped (if not already time-stamped by the combination generator 3700) and/or the generator data can be compressed (if not already compressed by the combination generator 3700). Time-stamping allows for the cloud-based system 205 to correlate the generator data with other data (e.g., stripped patient data) which may be communicated to the

cloud-based system 205. The compression allows for a smaller representation of the generator data to be subsequently encrypted and communicated to the cloud-based system 205. For the compression, a component of the surgical hub 206 can utilize a compression algorithm to convert a representation of the generator data to a smaller representation of the generator data, thereby allowing for a more efficient and economical encryption of the generator data (e.g., less data to encrypt utilizes less processing resources) and a more efficient and economical communication of the encrypted generator data (e.g., smaller representations of the generator data within the payload of the datagrams (e.g., FIGS. 28 and 29) allow for more generator data to be included in a given datagram, for more generator data to be communicated within a given time period, and/or for generator data to be communicated with fewer communication resources). The component of the surgical hub 206 which utilizes/executes the compression algorithm can be, for example, the processor module 232, the processor 244 of the computer system, and/or combinations thereof. The utilized/executed compression algorithm can be a lossless compression algorithm or a lossy compression algorithm.

[0666] Once the generator data and the MAC for a given datagram has been received at the cloud-based system 205 (e.g., FIG. 25, reference 3748; FIG. 26, reference 3758; and FIG. 27, reference 3768), the cloud-based system 205 can decrypt the encrypted generator data from the payload of the communicated datagram to realize the communicated generator data.

[0667] In one aspect, referring back to FIG. 25, the recipient (e.g., cloud-based system 205) may, similar to the surgical hub 206, process the ciphertext 3744 through the same MAC algorithm using the same known/accessible secret key to produce an authenticating MAC. If the received MAC 3745 matches this authenticating MAC, the recipient (e.g., cloud-based system 205) may safely assume that the ciphertext 3744 has not been altered and is from the surgical hub 206. The recipient (e.g., cloud-based system 205) may then decrypt the ciphertext 3744 (e.g., via key 3746) to realize the plaintext 3742 (e.g., data packet comprising generator data).

[0668] In another aspect, referring back to FIG. 26, the recipient (e.g., cloud-based system 205) may decrypt the ciphertext 3754 (e.g., via key 3756) to realize the plaintext 3752 (e.g., data packet comprising generator data). Next, similar to the surgical hub 206, the recipient (e.g., cloud-based system 205) may process the plaintext 3752 through the same MAC algorithm using the same known/accessible secret key to produce an authenticating MAC. If the received MAC 3755 matches this authenticating MAC, the recipient (e.g., cloud-based system 205) may safely assume that the plaintext 3752 has not been altered and is from the surgical hub 206.

[0669] In yet another aspect, referring back to FIG. 27, the recipient (e.g., cloud-based system 205) may decrypt the ciphertext 3764 (e.g., via key 3766) to realize the plaintext 3762 (e.g., data packet comprising generator data) and the MAC 3765. Next, similar to the surgical hub 206, the recipient (e.g., cloud-based system 205) may process the plaintext 3762 through the same MAC algorithm using the same known/accessible secret key to produce an authenticating MAC. If the received MAC 3765 matches this authenticating MAC, the recipient (e.g., cloud-based system

205) may safely assume that the plaintext 3762 has not been altered and is from the surgical hub 206.

[0670] In alternative aspects, the key used to encrypt the non-encrypted generator data (e.g., FIG. 25 and FIG. 26) or the non-encrypted generator data and the MAC (e.g., FIG. 27) may be different from the key (e.g., keys 3746, 3756, 3766) used to produce the MAC. For example, the key used to encrypt the non-encrypted generator data (e.g., FIG. 25 and FIG. 26) or the non-encrypted generator data and the MAC (e.g., FIG. 27) may be a different shared secret or a public key associated with the recipient. In such aspects, referring to FIG. 25, the recipient (e.g., cloud-based system 205) may, after verifying the authenticating MAC via key 3746 (described above), then decrypt the ciphertext 3744 (e.g., via the different shared secret or private key associated with the recipient) to realize the plaintext 3742 (e.g., data packet comprising generator data). In such aspects, referring to FIG. 26, the recipient may decrypt the ciphertext 3754 (e.g., via the different shared secret or private key associated with the recipient) to realize the plaintext 3752 (e.g., data packet comprising generator data), then verify the authenticating MAC via key 3756 (described above). In such aspects, referring to FIG. 27, the recipient may decrypt the ciphertext 3764 (e.g., via the different shared secret or private key associated with the recipient) to realize the plaintext 3762 (e.g., data packet comprising generator data) and the MAC 3765, then verify the authenticating MAC via key 3766 (described above).

[0671] In sum, referring to FIGS. 25-27, if an authenticating MAC, as determined/calculated by the cloud-based system 205, is the same as the MAC which was received with the datagram, the cloud-based system 205 can have confidence that the received generator data is authentic (i.e., it is the same as the generator data which was communicated by the surgical hub 206) and that the data integrity of the communicated generator data has not been compromised or altered. As described above, the recipient may further apply the plaintext 3742, 3752, 3762, or at least a portion thereof to the same checksum function/algorithm (i.e., used by the surgical hub 206) to generate a validating checksum value to further verify the integrity of the generator data based on the checksum value stored in the header of the communicated datagram.

[0672] Additionally, based on the decrypted datagram, the IP address of the source (e.g., FIG. 29, reference 3786) which originally communicated the datagram to the cloud-based system 205 can be determined from the header of the communicated datagram. If the determined source is a recognized source, the cloud-based system 205 can have confidence that the generator data originated from a trusted source, thereby providing source authentication and even more assurance of the data integrity of the generator data. Furthermore, since each router the datagram passed through in route to the cloud-based system 205 includes its IP address with its forwarded communication, the cloud-based system 205 is able to trace back the path followed by the datagram and identify each router which handled the datagram. The ability to identify the respective routers can be helpful in instances where the content of the datagram received at the cloud-based system 205 is not the same as the content of the datagram as originally communicated by the surgical hub 206. For aspects where the communication path was pre-specified and included in the header of the communicated datagram, the ability to identify the respective

routers can allow for path validation and provide additional confidence of the authenticity of the received generator data.

[0673] Furthermore, according to various aspects, after authenticating the received generator data, the cloud-based system 205 can communicate a message (e.g., a handshake or similar message) to the surgical hub 206 via the Internet or another communication network, confirming/guaranteeing that the datagram communicated from the surgical hub 206 was received intact by the cloud-based system 205, thereby effectively closing the loop for that particular datagram.

[0674] Aspects of the above-described communication method, and/or variations thereof, can also be employed to communicate data other than generator data to the cloud-based system 205 and/or to communicate generator data and/or other data from the surgical hub 206 to systems and/or devices other than the cloud-based system 205. For example, according to various aspects, the generator data and/or other data can be communicated from the surgical hub 206 to a hand-held surgical device/instrument (e.g., wireless device/instrument 235), to a robotic interface of a surgical device/instrument (e.g., robot hub 222) and/or to other servers, including servers (e.g., similar to server 213) associated with other cloud-based systems (e.g., similar to cloud-based system 205) in accordance with the above-described communication method. For example, in certain instances, an EEPROM chip of a given surgical instrument can initially be provided with merely an electronic chip device ID. Upon connection of the given surgical instrument to the combination generator 3700, data can be downloaded from the cloud-based system 205 to the surgical hub 206 and subsequently to the EEPROM of the surgical instrument in accordance with the above-described communication method.

[0675] In addition to communicating generator data to the cloud-based system 205, the surgical hub 206 can also utilize the above-described method of communication, and/or variations thereof, to communicate data other than generator data to the cloud-based system 205. For example, the surgical hub 206 can also communicate other information associated with the surgical procedure to the cloud-based system 205. Such other information can include, for example, the type of surgical procedure being performed, the name of the facility where the surgical procedure is being performed, the location of the facility where the surgical procedure is being performed, an identification of the operating room within the facility where the surgical procedure is being performed, the name of the surgeon performing the surgical procedure, the age of the patient, and data associated with the condition of the patient (e.g., blood pressure, heart rate, current medications). According to various aspects, such other information may be stripped of all information which could identify the specific surgery, the patient, or the surgeon, so that the information is essentially anonymized for further processing and analysis by the cloud-based system 205. In other words, the stripped data is not correlated to a specific surgery, patient, or surgeon. The stripped information can be communicated to the cloud-based system 205 either together with or distinct from the communicated generator data.

[0676] For instances where the stripped/other data is to be communicated apart from the generator data, the stripped/other data can be time-stamped, compressed, and/or encrypted in a manner identical to or different from that



described above regarding the generator data, and the surgical hub **206** may be programmed/configured to generate a datagram which includes the encrypted stripped/other information in lieu of the encrypted generator data. The datagram can then be communicated from the surgical hub **206** through the Internet to the cloud-based system **205** following an IP which: (1) defines datagrams that encapsulate the encrypted stripped/other data to be delivered, and (2) defines addressing methods that are used to label the datagram with source and destination information.

[**0677**] For instances where the stripped/other information is to be communicated with the generator data, the stripped/other data can be time-stamped, compressed, and/or encrypted in a manner identical to or different from that described above regarding the generator data, and the surgical hub **206** may be programmed/configured to generate a datagram which includes both the encrypted generator data and the encrypted stripped/other information. An example of such a datagram is shown in FIG. **30**, where the payload **3804** of the datagram **3800** is divided into two or more distinct payload data portions (e.g., one for the encrypted generator data **3834**, one for the encrypted stripped/other information **3836**), with each portion having an identifying bit (e.g., generator data (GD) **3806**, other data (OD) **3812**), the associated encrypted data **3808**, **3814**, and the associated padding **3810**, **3816**, if needed, respectively. Further, as shown in FIG. **30**, the header **3802** may be the same as (e.g., IP address source **3818**, IP address destination **3820**, header length **3822**) or different from the header **3782** described with reference to the datagram **3780** shown in FIG. **29**. For example, the header **3802** may be different in that the header **3802** further includes a field designating the number of payload data portions **3824** (e.g., 2) included in the payload **3804** of the datagram **3800**. The header **3802** can also be different in that it can include fields designating the payload length **3826**, **3830** and the checksum value **3828**, **2832** for each payload data portion **3834**, **3836**, respectively. Although only two payload data portions are shown in FIG. **30**, it will be appreciated that the payload **3804** of the datagram **3800** may include any quantity/number of payload data portions (e.g., 1, 2, 3, 4, 5), where each payload data portion includes data associated with a different aspect of the surgical procedure. The datagram **3800** can then be communicated from the surgical hub **206** through the Internet to the cloud-based system **205** following an IP which: (1) defines datagrams that encapsulate the encrypted generator data and the encrypted stripped/other data to be delivered, and (2) defines addressing methods that are used to label the datagram with source and destination information.

[**0678**] As set forth above, it is an unfortunate reality that the outcomes of all surgical procedures are not always optimal and/or successful. For instances where a failure event is detected and/or identified, a variation of the above-described communication methods can be utilized to isolate surgical data which is associated with the failure event (e.g., failure event surgical data) from surgical data which is not associated with the failure event (e.g., non-failure event surgical data) and communicate the surgical data which is associated with the failure event (e.g., failure event data) from the surgical hub **206** to the cloud-based system **205** on a prioritized basis for analysis. According to one aspect of the present disclosure, failure event surgical data is com-

municated from the surgical hub **206** to the cloud-based system **205** on a prioritized basis relative to non-failure event surgical data.

[**0679**] FIG. **31** illustrates various aspects of a system-implemented method of identifying surgical data associated with a failure event (e.g., failure event surgical data) and communicating the identified surgical data to a cloud-based system **205** on a prioritized basis. The method comprises (1) receiving **3838** surgical data at a surgical hub **206**, wherein the surgical data is associated with a surgical procedure; (2) time-stamping **3840** the surgical data; (3) identifying **3842** a failure event associated with the surgical procedure; (4) determining **3844** which of the surgical data is associated with the failure event (e.g., failure event surgical data); (5) separating **3846** the surgical data associated with the failure event from all other surgical data (e.g., non-failure event surgical data) received at the surgical hub **206**; (6) chronologizing **3848** the surgical data associated with the failure event; (7) encrypting **3850** the surgical data associated with the failure event; and (8) communicating **3852** the encrypted surgical data to a cloud-based system **205** on a prioritized basis.

[**0680**] More specifically, various surgical data can be captured during a surgical procedure and the captured surgical data, as well as other surgical data associated with the surgical procedure, can be communicated to the surgical hub **206**. The surgical data can include, for example, data associated with a surgical device/instrument (e.g., FIG. **9**, surgical device/instrument **235**) utilized during the surgery, data associated with the patient, data associated with the facility where the surgical procedure was performed, and data associated with the surgeon. Either prior to or subsequent to the surgical data being communicated to and received by the surgical hub **206**, the surgical data can be time-stamped and/or stripped of all information which could identify the specific surgery, the patient, or the surgeon, so that the information is essentially anonymized for further processing and analysis by the cloud-based system **205**.

[**0681**] Once a failure event has been detected and/or identified (e.g., which can be either during or after the surgical procedure), the surgical hub **206** can determine which of the surgical data is associated with the failure event (e.g., failure event surgical data) and which of the surgical data is not associated with the surgical event (e.g., non-failure event surgical data). According to one aspect of the present disclosure, a failure event can include, for example, a detection of one or more misfired staples during a stapling portion of a surgical procedure. For example, in one aspect, referring to FIG. **9**, an endoscope **239** may take snapshots while a surgical device/instrument **235** comprising an end effector including a staple cartridge performs a stapling portion of a surgical procedure. In such an aspect, an imaging module **238** may compare the snapshots to stored images and/or images downloaded from the cloud-based system **205** that convey correctly fired staples to detect a misfired staple and/or evidence of a misfired staple (e.g., a leak). In another aspect, the imaging module **238** may analyze the snapshots themselves to detect a misfired staple and/or evidence of a misfired staple. In one alternative aspect, the surgical hub **206** may communicate the snapshots to the cloud-based system **205**, and a component of the cloud-based system **205** may perform the various imaging module functions described above to detect a misfired staple and/or evidence of a misfired staple and to report the

detection to the surgical hub 206. According to another aspect of the present disclosure, a failure event can include a detection of a tissue temperature which is below the expected temperature during a tissue-sealing portion of a surgical procedure and/or a visual indication of excessive bleeding or oozing following a surgical procedure (e.g., FIG. 9, via endoscope 239). For example, in one aspect, referring to FIG. 9, the surgical device/instrument 235 may comprise an end effector, including a temperature sensor and the surgical hub 206, and/or the cloud-based system may compare at least one temperature detected by the temperature sensor (e.g., during a tissue-sealing portion of a surgical procedure) to a stored temperature and/or a range of temperatures expected and/or associated with that surgical procedure to detect an inadequate/low sealing temperature. In another aspect, an endoscope 239 may take snapshots during a surgical procedure. In such an aspect, an imaging module 238 may compare the snapshots to stored images and/or images downloaded from the cloud-based system 205 that convey tissue correctly sealed at expected temperatures to detect evidence of an improper/insufficient sealing temperature (e.g., charring, oozing/bleeding). Further, in such an aspect, the imaging module 238 may analyze the snapshots themselves to detect evidence of an improper/insufficient sealing temperature (e.g., charring, oozing/bleeding). In one alternative aspect, the surgical hub 206 may communicate the snapshots to the cloud-based system 205, and a component of the cloud-based system 205 may perform the various imaging module functions described above to detect evidence of an improper/insufficient sealing temperature and to report the detection to the surgical hub 206. According to the various aspects described above, in response to the detected and/or identified failure event, the surgical hub 206 may download a program from the cloud-based system 205 for execution by the surgical device/instrument 235 that corrects the detected issue (i.e., program that alters surgical device/instrument parameters to prevent misfired staples, program that alters surgical device/instrument parameters to ensure correct sealing temperature).

**[0682]** In some aspects, a failure event is deemed to cover a certain time period, and all surgical data associated with that certain time period can be deemed to be associated with the failure event.

**[0683]** After the surgical data associated with the failure event has been identified, the identified surgical data (e.g., failure event surgical data) can be separated or isolated from all of the other surgical data associated with the surgical procedure (e.g., non-failure event surgical data). The separation can be realized, for example, by tagging or flagging the identified surgical data, by storing the identified surgical data apart from all of the other surgical data associated with the surgical procedure, or by storing only the other surgical data while continuing to process the identified surgical data for subsequent prioritized communication to the cloud-based system 205. According to various aspects, the tagging or flagging of the identified surgical data can occur during the communication process when the datagram is generated as described in more detail below.

**[0684]** The time-stamping of all of the surgical data (e.g., either before or after the surgical data is received at the surgical hub) can be utilized by a component of the surgical hub 206 to chronologize the identified surgical data associated with the failure event. The component of the surgical hub 206 which utilizes the time-stamping to chronologize

the identified surgical data can be, for example, the processor module 232, the processor 244 of the computer system 210, and/or combinations thereof. By chronologizing the identified surgical data, the cloud-based system 205 and/or other interested parties can subsequently better understand the conditions which were present leading up to the occurrence of the failure event and possibly pinpoint the exact cause of the failure event, thereby providing the knowledge to potentially mitigate a similar failure event from occurring during a similar surgical procedure performed at a future date.

**[0685]** Once the identified surgical data has been chronologized, the chronologized surgical data may be encrypted in a manner similar to that described above with respect to the encryption of the generator data. Thus, the identified surgical data can be encrypted to help ensure the confidentiality of the identified surgical data, either while it is being stored at the surgical hub 206 or while it is being transmitted to the cloud-based system 205 using the Internet or other computer networks. According to various aspects, a component of the surgical hub 206 utilizes an encryption algorithm to convert the identified surgical data from a readable version to an encoded version, thereby forming the encrypted surgical data associated with the failure event (e.g., FIGS. 25-27). The component of the surgical hub which utilizes the encryption algorithm can be, for example, the processor module 232, the processor 244 of the computer system 210, and/or combinations thereof. The utilized encryption algorithm can be a symmetric encryption algorithm or an asymmetric encryption algorithm.

**[0686]** After the identified surgical data has been encrypted, a component of the surgical hub can communicate the encrypted surgical data associated with the failure event (e.g., encrypted failure event surgical data) to the cloud-based system 205. The component of the surgical hub which communicates the encrypted surgical data to the cloud-based system 205 can be, for example, the processor module 232, a hub/switch 207/209 of the modular communication hub 203, the router 211 of the modular communication hub 203, or the communication module 247 of the computer system 210. According to various aspects, the communication of the encrypted surgical data (e.g., encrypted failure event surgical data) through the Internet can follow an IP which: (1) defines datagrams that encapsulate the encrypted surgical data to be delivered, and (2) defines addressing methods that are used to label the datagram with source and destination information. The datagram can be similar to the datagram shown in FIG. 29 or the datagram shown in FIG. 30, but can be different in that either the header or the payload of the datagram can include a field which includes a flag or a tag which identifies the encrypted surgical data (e.g., encrypted failure event surgical data) as being prioritized relative to other non-prioritized surgical data (e.g., encrypted non-failure event surgical data). An example of such a datagram is shown in FIG. 32, where the payload 3864 of the datagram 3860 includes a field which indicates (e.g., a prioritized designation 3834) that the payload 3864 includes prioritized surgical data (e.g., combination generator data 3868). According to various aspects, the payload 3864 of the datagram 3860 can also include non-flagged/non-tagged/non-prioritized surgical data 3836 (e.g., other surgical data 3874) as shown in FIG. 32.

**[0687]** According to various aspects, prior to the identified surgical data (e.g., failure event surgical data) being

encrypted, the identified surgical data can be compressed (if not already compressed by the source(s) of the relevant surgical data). The compression allows for a smaller representation of the surgical data associated with the failure event to be subsequently encrypted and communicated to the cloud-based system 205. For the compression, a component of the surgical hub 206 can utilize a compression algorithm to convert a representation of the identified surgical data to a smaller representation of the identified surgical data, thereby allowing for a more efficient and economical encryption of the identified surgical data (less data to encrypt utilizes less processing resources) and a more efficient and economical communication of the encrypted surgical data (smaller representations of the surgical data within the payload of the datagrams allow for more identified surgical data to be included in a given datagram, for more identified surgical data to be communicated within a given time period, and/or for identified surgical data to be communicated with fewer communication resources). The component of the surgical hub 206 which utilizes the compression algorithm can be, for example, the processor module 232, the processor 244 of the computer system 210, and/or combinations thereof. The utilized compression algorithm can be a lossless compression algorithm or a lossy compression algorithm.

[0688] In instances where other non-prioritized surgical data (e.g., non-failure event surgical data) is to be communicated with prioritized surgical data (e.g., failure event surgical data), the other non-prioritized surgical data can be time-stamped, compressed, and/or encrypted in a manner identical to or different from that described above regarding the surgical data identified as associated with a failure event (e.g., failure event surgical data), and the surgical hub 206 may be programmed/configured to generate a datagram which includes both the encrypted prioritized surgical data (e.g., encrypted failure event surgical data) and the encrypted other non-prioritized surgical data (e.g., encrypted non-failure event surgical data). For example, in light of FIG. 32, the payload 3864 of the datagram 3860 may be divided into two or more distinct payload data portions (e.g., one for the prioritized surgical data 3834, one for the non-prioritized surgical data 3836), with each portion having an identifying bit (e.g., generator data (GD) 3866, other data (OD) 3872), the associated encrypted data (e.g., encrypted prioritized surgical data 3868, encrypted non-prioritized surgical data 3874), and the associated padding 3870, 3876, if needed, respectively. Further, similar to FIG. 30, the header 3862 may be the same as (e.g., IP address source 3878, IP address destination 3880, header length 3882) or different from the header 3782 described with reference to the datagram 3780 shown in FIG. 29. For example, the header 3862 may be different in that the header 3862 further includes a field designating the number of payload data portions 3884 (e.g., 2) included in the payload 3864 of the datagram 3860. The header 3862 can also be different in that it can include fields designating the payload length 3886, 3890 and the checksum value 3888, 2892 for each payload data portion 3834, 3836, respectively. Although only two payload data portions are shown in FIG. 32, it will be appreciated that the payload 3864 of the datagram 3860 may include any quantity/number of payload data portions (e.g., 1, 2, 3, 4, 5), where each payload data portion includes data associated with a different aspect of the surgical procedure. The datagram 3860 can then be com-

municated from the surgical hub 206 through the Internet to the cloud-based system 205 following an IP which: (1) defines datagrams that encapsulate the encrypted generator data and the encrypted stripped/other data to be delivered, and (2) defines addressing methods that are used to label the datagram with source and destination information.

[0689] In some aspects, once a failure event associated with a surgical procedure has been identified, the surgical hub 206 and/or the cloud-based system 205 can subsequently flag or tag a surgical device/instrument 235 which was utilized during the surgical procedure for inoperability and/or removal. For example, in one aspect, information (e.g., serial number, ID) associated with the surgical device/instrument 235 and stored at the surgical hub 206 and/or the cloud-based system 205 can be utilized to effectively block the surgical device/instrument 235 from being used again (e.g., blacklisted). In another aspect, information (e.g., serial number, ID) associated with the surgical device/instrument can initiate the printing of a shipping slip and shipping instructions for returning the surgical device/instrument 235 back to a manufacturer or other designated party so that a thorough analysis/inspection of the surgical device/instrument 235 can be performed (e.g., to determine the cause of the failure). According to various aspects described herein, once the cause of a failure is determined (e.g., via the surgical hub 206 and/or the cloud-based system 205), the surgical hub 206 may download a program from the cloud-based system 205 for execution by the surgical device/instrument 235 that corrects the determined cause of the failure (i.e., program that alters surgical device/instrument parameters to prevent the failure from occurring again).

[0690] According to some aspects, the surgical hub 206 and/or the cloud-based system 205 can also provide/display a reminder (e.g., via hub display 215 and/or surgical device/instrument display 237) to administrators, staff, and/or other personnel to physically remove the surgical device/instrument 235 from the operating room (e.g., if detected as still present in the operating room) and/or to send the surgical device/instrument 235 to the manufacturer or the other designated party. In one aspect, the reminder may be set up to be provided/displayed periodically until an administrator can remove the flag or tag of the surgical device/instrument 235 from the surgical hub 206 and/or the cloud-based system 205. According to various aspects, an administrator may remove the flag or tag once the administrator can confirm (e.g., system tracking of the surgical device/instrument 235 via its serial number/ID) that the surgical device/instrument 235 has been received by the manufacturer or the other designated party. By using the above-described method to flag and/or track surgical data associated with a failure event, a closed loop control of the surgical data associated with the failure event and/or with a surgical device/instrument 235 can be realized. Additionally, in view of the above, it will be appreciated that the surgical hub 206 can be utilized to effectively manage the utilization (or non-utilization) of surgical devices/instruments 235 which have or potentially could be utilized during a surgical procedure.

[0691] In various aspects of the present disclosure, the surgical hub 206 and/or cloud-based system 205 may want to control which components (e.g., surgical device/instrument 235, energy device 241) are being utilized in its interactive surgical system 100/200 to perform surgical procedures (e.g., to minimize future failure events, to avoid the use of unauthorized or knock-off components).

[0692] As such, in various aspects of the present disclosure, since an interactive surgical system 100 may comprise a plurality of surgical hubs 106, a cloud-based system 105 and/or each surgical hub 106 of the interactive surgical system 100 may want to track component-surgical hub combinations utilized over time. In one aspect, upon/after a component (See FIG. 9, e.g., surgical device/instrument 235, energy device 241) is connected to/used with a particular surgical hub 106 (e.g., surgical device/instrument 235 wired/wirelessly connected to the particular surgical hub 106, energy device 241 connected to the particular surgical hub 106 via generator module 240), the particular surgical hub 106 may communicate a record/block of that connection/use (e.g., linking respective unique identifiers of the connected devices) to the cloud-based system 105 and/or to the other surgical hubs 106 in the interactive surgical system 100. For example, upon/after the connection/use of an energy device 241, a particular surgical hub 106 may communicate a record/block (e.g., linking a unique identifier of the energy device 241 to a unique identifier of a generator module 240 to a unique identifier of the particular surgical hub 106) to the cloud-based system 105 and/or other surgical hubs 106 in the interactive surgical system 100. In such an aspect, if this is the first time the component (e.g., energy device) is connected to/used with a surgical hub 106 in the interactive surgical system 100, the cloud-based system 105 and/or each surgical hub 106 of the interactive surgical system 100 may store the record/block as a genesis record/block. In such an aspect, the genesis record/block stored at the cloud-based system 105 and/or each surgical hub 106 may comprise a time stamp. However, in such an aspect, if this is not the first time the component (e.g., energy device 241) has been connected to/used with a surgical hub 106 in the interactive surgical system 100, the cloud-based system 105 and/or each surgical hub 106 of the interactive surgical system may store the record/block as a new record/block in a chain of record/blocks associated with the component. In such an aspect, the new record/block may comprise a cryptographic hash of the most recently communicated record/block stored at the cloud-based system 105 and/or each surgical hub 106, the communicated linkage data, and a time stamp. In such an aspect, each cryptographic hash links each new record/block (e.g., each use of the component) to its prior record/block to form a chain confirming the integrity of each prior record/block(s) back to an original genesis record/block (e.g., first use of the component). According to such an aspect, this blockchain of records/blocks may be developed at the cloud-based system 105 and/or each surgical hub 106 of the interactive surgical system 100 to permanently and verifiably tie usage of a particular component to one or more than one surgical hub 106 in the interactive surgical system 100 over time. Here, according to another aspect, this approach may be similarly applied to sub-components (e.g., handle, shaft, end effector, cartridge) of a component when/after the component is connected to/used with a particular surgical hub 106 of an interactive surgical system 100.

[0693] According to various aspects of the present disclosure, the cloud-based system 105 and/or each surgical hub 106 may utilize such records/blocks to trace usage of a particular component and/or a sub-component back to its initial usage in the interactive surgical system 100. For example, if a particular component (e.g., surgical device/instrument 235) is flagged/tagged as related to a failure event, the cloud-based system 105 and/or a surgical hub 106

may analyze such records/blocks to determine whether past usage of that component and/or a sub-component of that component contributed to or caused the failure event (e.g., overused). In one example, the cloud-based system 105 may determine that a sub-component (e.g., end effector) of that component may actually be contributing/causing the failure event and then tag/flag that component for inoperability and/or removal based on the determination.

[0694] According to another aspect, the cloud-based system 205 and/or surgical hub 206 may control which components (e.g., surgical device/instrument 235, energy device 241) are being utilized in an interactive surgical system 200 to perform surgical procedures by authenticating the component and/or its supplier/manufacture. In one aspect, the supplier/manufacture of a component may associate a serial number and a source ID with the component. In such an aspect, the supplier/manufacture may create/generate a private key for the serial number, encrypt the serial number with the private key, and store the encrypted serial number and the source ID on an electronic chip (e.g., memory) in the component prior to shipment to a surgical site. Here, upon/after connection of the component to a surgical hub 206, the surgical hub 206 may read the encrypted serial number and the source ID from the electronic chip. In response, the surgical hub 206 may send a message (i.e., comprising the encrypted serial number) to a server of the supplier/manufacture associated with the source ID (e.g., directly or via the cloud-based system 205). In such an aspect, the surgical hub 206 may encrypt the message using a public key associated with that supplier/manufacture. In response, the surgical hub 206 may receive a message (i.e., comprising the private key the supplier/manufacture generated for/associated with that encrypted serial number) from the supplier/manufacture server (e.g., directly or via the cloud-based system 205). In such an aspect, the supplier/manufacture server may encrypt the message using a public key associated with the surgical hub 206. Further, in such an aspect, the surgical hub 206 may then decrypt the message (e.g., using a private key paired to the public key used to encrypt the message) to reveal the private key associated with the encrypted serial number. The surgical hub 206 may then decrypt the encrypted serial number, using that private key, to reveal the serial number. Further, in such an aspect, the surgical hub 206 may then compare the decrypted serial number to a comprehensive list of authorized serial numbers (e.g., stored at the surgical hub 206 and/or the cloud-based system and/or downloaded from the cloud-based system, e.g., received separately from the supplier/manufacture) and permit use of the connected component if the decrypted serial number matches an authorized serial number. Initially, such a process permits the surgical hub 206 to authenticate the supplier/manufacture. In particular, the surgical hub 206 encrypted the message comprising the encrypted serial number using a public key associated with the supplier/manufacture. As such, receiving a response message (i.e., comprising the private key) authenticates the supplier/manufacture to the surgical hub 206 (i.e., otherwise the supplier/manufacture would not have access to the private key paired to the public key used by the surgical hub 206 to encrypt the message, and the supplier/manufacture would not have been able to associate the encrypted serial number received in the message to its already generated private key). Furthermore, such a process permits the surgical hub 206 to authenticate the connected component/device itself. In par-

ticular, the supplier/manufacture (e.g., just authenticated) encrypted the serial number of the component using the delivered private key. Upon secure receipt of the private key, the surgical hub 206 is able to decrypt the encrypted serial number (i.e., read from the connected component), which authenticates the component and/or its association with the supplier/manufacture (i.e., only that private key as received from that supplier/manufacture would decrypt the encrypted serial number). Nonetheless, the surgical hub 206 further verifies the component as authentic (e.g., compares the decrypted serial number to a comprehensive list of authorized serial numbers received separately from the supplier/manufacture). Notably, such aspects as described above can alternatively be performed by the cloud-based system 205 and/or a combination of the cloud-based system 205 and the surgical hub 206 to control which components (e.g., surgical device/instrument 235, energy device 241) are being utilized in an interactive surgical system 200 (e.g., to perform surgical procedures) by authenticating the component and/or its supplier/manufacture. In one aspect, such described approaches may prevent the use of knock-off component(s) within the interactive surgical system 200 and ensure the safety and well-being of surgical patients.

**[0695]** According to another aspect, the electronic chip of a component (e.g., surgical device/instrument 235, energy device 241) may store (e.g., in memory) data associated with usage of that component (i.e., usage data, e.g., number of uses with a limited use device, number of uses remaining, firing algorithms executed, designation as a single-use component). In such an aspect, the surgical hub 206 and/or the cloud-based system 205, upon/after connection of the component to the interactive surgical system, may read such usage data from the memory of a component and write back at least a portion of that usage data for storage (e.g., in memory 249) at the surgical hub 206 and/or for storage at the cloud-based system 205 (e.g., individually and/or under a blockchain approach discussed herein). According to such an aspect, the surgical hub 206 and/or the cloud-based system 205, upon/after a subsequent connection of that component to the interactive surgical system, may again read such usage data and compare that usage to previously stored usage data. Here, if a discrepancy exists or if a predetermined/authorized usage has been met, the surgical hub 206 and/or the cloud-based system 205 may prevent use of that component (e.g., blacklisted, rendered inoperable, flagged for removal) on the interactive surgical system 200. In various aspects, such an approach prevents bypass of the encryption chip systems. If the component's electronic chip/memory has been tampered with (e.g., memory reset, number of uses altered, firing algorithms altered, single-use device designated as a multi-use device), a discrepancy will exist, and the component's use will be controlled/prevented.

**[0696]** Additional details are disclosed in U.S. Patent Application Publication No. 2017/0086914, titled TECHNIQUES FOR OPERATING GENERATOR FOR DIGITALLY GENERATING ELECTRICAL SIGNAL WAVEFORMS AND SURGICAL INSTRUMENTS, which published on Mar. 30, 2017, which is incorporated herein by reference in its entirety.

#### Surgical Hub Coordination of Device Pairing in an Operating Room

**[0697]** One of the functions of the surgical hub 106 is to pair (also referred to herein as "connect" or "couple") with

other components of the surgical system 102 to control, gather information from, or coordinate interactions between the components of the surgical system 102. Since the operating rooms of a hospital are likely in close physical proximity to one another, a surgical hub 106 of a surgical system 102 may unknowingly pair with components of a surgical system 102 in a neighboring operating room, which would significantly interfere with the functions of the surgical hub 106. For example, the surgical hub 106 may unintentionally activate a surgical instrument in a different operating room or record information from a different ongoing surgical procedure in a neighboring operating room.

**[0698]** Aspects of the present disclosure present a solution, wherein a surgical hub 106 only pairs with detected devices of the surgical system 102 that are located within the bounds of its operating room.

**[0699]** Furthermore, the surgical hub 106 relies on its knowledge of the location of other components of the surgical system 102 within its operating room in making decisions about, for example, which surgical instruments should be paired with one another or activated. A change in the position of the surgical hub 106 or another component of the surgical system 102 can be problematic.

**[0700]** Aspects of the present disclosure further present a solution wherein the surgical hub 106 is configured to reevaluate or redetermine the bounds of its operating room upon detecting that the surgical hub 106 has been moved. Aspects of the present disclosure further present a solution wherein the surgical hub 106 is configured to redetermine the bounds of its operating room upon detection of a potential device of the surgical system 102, which can be an indication that the surgical hub 106 has been moved.

**[0701]** In various aspects, a surgical hub 106 is used with a surgical system 102 in a surgical procedure performed in an operating room. The surgical hub 106 comprises a control circuit configured to determine the bounds of the operating room, determine devices of the surgical system 102 located within the bounds of the operating room, and pair the surgical hub 106 with the devices of the surgical system 102 located within the bounds of the operating room.

**[0702]** In one aspect, the control circuit is configured to determine the bounds of the operating room after activation of the surgical hub 106. In one aspect, the surgical hub 106 includes a communication circuit configured to detect and pair with the devices of the surgical system located within the bounds of the operating room. In one aspect, the control circuit is configured to redetermine the bounds of the operating room after a potential device of the surgical system 102 is detected. In one aspect, the control circuit is configured to periodically determine the bounds of the operating room.

**[0703]** In one aspect, the surgical hub 106 comprises an operating room mapping circuit that includes a plurality of non-contact sensors configured to measure the bounds of the operating room.

**[0704]** In various aspects, the surgical hub 106 includes a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to pair the surgical hub with devices of the surgical system 102 located within the bounds of the operating room, as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer-readable instructions which, when executed, cause a machine to pair the surgical hub 106 with devices of

the surgical system **102** located within the bounds of the operating room, as described above.

**[0705]** FIGS. **35** and **36** are logic flow diagrams of processes depicting control programs or logic configurations for pairing the surgical hub **106** with devices of the surgical system **102** located within the bounds of the operating room, as described above.

**[0706]** The surgical hub **106** performs a wide range of functions that requires short- and long-range communication, such as assisting in a surgical procedure, coordinating between devices of the surgical system **102**, and gathering and transmitting data to the cloud **104**. To properly perform its functions, the surgical hub **106** is equipped with a communication module **130** capable of short-range communication with other devices of the surgical system **102**. The communication module **130** is also capable of long-range communication with the cloud **104**.

**[0707]** The surgical hub **106** is also equipped with an operating-room mapping module **133** which is capable of identifying the bounds of an operating room, and identifying devices of the surgical system **102** within the operating room. The surgical hub **106** is configured to identify the bounds of an operating room, and only pair with or connect to potential devices of the surgical system **102** that are detected within the operating room.

**[0708]** In one aspect, the pairing comprises establishing a communication link or pathway. In another aspect, the pairing comprises establishing a control link or pathway.

**[0709]** An initial mapping or evaluation of the bounds of the operating room takes place during an initial activation of the surgical hub **106**. Furthermore, the surgical hub **106** is configured to maintain spatial awareness during operation by periodically mapping its operating room, which can be helpful in determining if the surgical hub **106** has been moved. The reevaluation **3017** can be performed periodically or it can be triggered by an event such as observing a change in the devices of the surgical system **102** that are deemed within the operating room. In one aspect, the change is detection **3010** of a new device that was not previously deemed as within the bounds of the operating room, as illustrated in FIG. **37**. In another aspect, the change is a disappearance, disconnection, or un-pairing of a paired device that was previously deemed as residing within the operating room, as illustrated in FIG. **38**. The surgical hub **106** may continuously monitor **3035** the connection with paired devices to detect **3034** the disappearance, disconnection, or un-pairing of a paired device.

**[0710]** In other aspects, reevaluation triggering events can be, for example, changes in surgeons' positions, instrument exchanges, or sensing of a new set of tasks being performed by the surgical hub **106**.

**[0711]** In one aspect, the evaluation of the bounds of the room by the surgical hub **106** is accomplished by activation of a sensor array of the operating-room mapping module **133** within the surgical hub **106** which enables it to detect the walls of the operating room.

**[0712]** Other components of the surgical system **102** can be made to be spatially aware in the same, or a similar, manner as the surgical hub **106**. For example, a robotic hub **122** may also be equipped with an operating-room mapping module **133**.

**[0713]** The spatial awareness of the surgical hub **106** and its ability to map an operating room for potential components of the surgical system **102** allows the surgical hub **106**

to make autonomous decisions about whether to include or exclude such potential components as part of the surgical system **102**, which relieves the surgical staff from dealing with such tasks. Furthermore, the surgical hub **106** is configured to make inferences about, for example, the type of surgical procedure to be performed in the operating room based on information gathered prior to, during, and/or after the performance of the surgical procedure. Examples of gathered information include the types of devices that are brought into the operating room, time of introduction of such devices into the operating room, and/or the devices sequence of activation.

**[0714]** In one aspect, the surgical hub **106** employs the operating-room mapping module **133** to determine the bounds of the surgical theater (e.g., a fixed, mobile, or temporary operating room or space) using either ultrasonic or laser non-contact measurement devices.

**[0715]** Referring to FIG. **34**, ultrasound based non-contact sensors **3002** can be employed to scan the operating theater by transmitting a burst of ultrasound and receiving the echo when it bounces off a perimeter wall **3006** of an operating theater to determine the size of the operating theater and to adjust Bluetooth pairing distance limits. In one example, the non-contact sensors **3002** can be Ping ultrasonic distance sensors, as illustrated in FIG. **34**.

**[0716]** FIG. **34** shows how an ultrasonic sensor **3002** sends a brief chirp with its ultrasonic speaker **3003** and makes it possible for a micro-controller **3004** of the operating-room mapping module **133** to measure how long the echo takes to return to the ultrasonic sensor's ultrasonic microphone **3005**. The micro-controller **3004** has to send the ultrasonic sensor **3002** a pulse to begin the measurement. The ultrasonic sensor **3002** then waits long enough for the micro-controller program to start a pulse input command. Then, at about the same time the ultrasonic sensor **3002** chirps a 40 kHz tone, it sends a high signal to the micro-controller **3004**. When the ultrasonic sensor **3002** detects the echo with its ultrasonic microphone **3005**, it changes that high signal back to low. The micro-controller's pulse input command measures the time between the high and low changes and stores its measurement in a variable. This value can be used along with the speed of sound in air to calculate the distance between the surgical hub **106** and the operating-room wall **3006**.

**[0717]** In one example, as illustrated in FIG. **33**, a surgical hub **106** can be equipped with four ultrasonic sensors **3002**, wherein each of the four ultrasonic sensors is configured to assess the distance between the surgical hub **106** and a wall of the operating room **3000**. A surgical hub **106** can be equipped with more or less than four ultrasonic sensors **3002** to determine the bounds of an operating room.

**[0718]** Other distance sensors can be employed by the operating-room mapping module **133** to determine the bounds of an operating room. In one example, the operating-room mapping module **133** can be equipped with one or more photoelectric sensors that can be employed to assess the bounds of an operating room. In one example, suitable laser distance sensors can also be employed to assess the bounds of an operating room. Laser-based non-contact sensors may scan the operating theater by transmitting laser light pulses, receiving laser light pulses that bounce off the perimeter walls of the operating theater, and comparing the phase of the transmitted pulse to the received pulse to

determine the size of the operating theater and to adjust Bluetooth pairing distance limits.

[0719] Referring to the top left corner of FIG. 33, a surgical hub 106 is brought into an operating room 3000. The surgical hub 106 is activated at the beginning of the set-up that occurs prior to the surgical procedure. In the example of FIG. 33, the set-up starts at an actual time of 11:31:14 (EST) based on a real-time clock. However, at the stated procedure set-up start time, the surgical hub 106 starts 3001 an artificial randomized real-time clock timing scheme at artificial real time 07:36:00 to protect private patient information.

[0720] At artificial real time 07:36:01, the operating-room mapping module 133 employs the ultrasonic distance sensors to ultrasonically ping the room (e.g., sends out a burst of ultrasound and listens for the echo when it bounces off the perimeter walls of the operating room as described above) to verify the size of the operating room and to adjust pairing distance limits.

[0721] At artificial real time 07:36:03, the data is stripped and time-stamped. At artificial real time 07:36:05, the surgical hub 106 begins pairing devices located only within the operating room 3000 as verified using ultrasonic distance sensors 3002 of the operating-room mapping module 133. The top right corner of FIG. 33 illustrates several example devices that are within the bounds of the operating room 3000 and are paired with the surgical hub 106, including a secondary display device 3020, a secondary hub 3021, a common interface device 3022, a powered stapler 3023, a video tower module 3024, and a powered handheld dissector 3025. On the other hand, secondary hub 3021', secondary display device 3020', and powered stapler 3026 are all outside the bounds of the operating room 3000 and, accordingly, are not paired with the surgical hub 106.

[0722] In addition to establishing a communication link with the devices of the surgical system 102 that are within the operating room, the surgical hub 106 also assigns a unique identification and communication sequence or number to each of the devices. The unique sequence may include the device's name and a time stamp of when the communication was first established. Other suitable device information may also be incorporated into the unique sequence of the device.

[0723] As illustrated in the top left corner of FIG. 33, the surgical hub 106 has determined that the operating room 3000 bounds are at distances  $a$ ,  $-a$ ,  $b$ , and  $-b$  from the surgical hub 106. Since Device "D" is outside the determined bounds of its operating room 3000, the surgical hub 106 will not pair with the Device "D." FIG. 35 is an example algorithm illustrating how the surgical hub 106 only pairs with devices within the bounds of its operating room. After activation, the surgical hub 106 determines 3007 bounds of the operating room using the operating-room mapping module 133, as described above. After the initial determination, the surgical hub 106 continuously searches for or detects 3008 devices within a pairing range. If a device is detected 3010, the surgical hub 106 then determines 3011 whether the detected device is within the bounds of the operating room. The surgical hub 106 pairs 3012 with the device if it is determined that the device is within the bounds of the operating room. In certain instances, the surgical hub 106 will also assign 3013 an identifier to the device. If, however, the surgical hub 106 determines that the detected device is

outside the bounds of the operating room, the surgical hub 106 will ignore 3014 the device.

[0724] Referring to FIG. 36, after an initial determination of the bounds of the room, and after an initial pairing of devices located within such bounds, the surgical hub 106 continues to detect 3015 new devices that become available for pairing. If a new device is detected 3016, the surgical hub 106 is configured to reevaluate 3017 the bounds of the operating room prior to pairing with the new device. If the new device is determined 3018 to be within the newly determined bounds of the operating room, then the surgical hub 106 pairs with the device 3019 and assigns 3030 a unique identifier to the new device. If, however, the surgical hub 106 determines that the new device is outside the newly determined bounds of the operating room, the surgical hub 106 will ignore 3031 the device.

[0725] For pairing, the operating-room mapping module 133 contains a compass and integrated Bluetooth transceiver. Other communication mechanisms, which are not significantly affected by the hospital environment or geographical location, can be employed. Bluetooth Low Energy (BLE) beacon technology can currently achieve indoor distance measurements with accuracy of about 1-2 meters, with improved accuracy in closer proximities (within 0-6 meters). To improve the accuracy of the distance measurements, a compass is used with the BLE. The operating-room mapping module 133 utilizes the BLE and the compass to determine where modules are located in relation to the patient. For example, two modules facing each other (detected by compass) with greater than one meter distance between them may clearly indicate that the modules are on opposite sides of the patient. The more "Hub"-enabled modules that reside in the operating room, the greater the achievable accuracy becomes due to triangulation techniques.

[0726] In the situations where multiple surgical hubs 106, modules, and/or other peripherals are present in the same operating room, as illustrated in the top right corner of FIG. 33, the operating-room mapping module 133 is configured to map the physical location of each module that resides within the operating room. This information could be used by the user interface to display a virtual map of the room, enabling the user to more easily identify which modules are present and enabled, as well as their current status. In one aspect, the mapping data collected by surgical hubs 106 are uploaded to the cloud 104, where the data are analyzed for identifying how an operating room is physically setup, for example.

[0727] The surgical hub 106 is configured to determine a device's location by assessing transmission radio signal strength and direction. For Bluetooth protocols, the Received Signal Strength Indication (RSSI) is a measurement of the received radio signal strength. In one aspect, the devices of the surgical system 102 can be equipped with USB Bluetooth dongles. The surgical hub 106 may scan the USB Bluetooth beacons to get distance information. In another aspect, multiple high-gain antennas on a Bluetooth access point with variable attenuators can produce more accurate results than RSSI measurements. In one aspect, the hub is configured to determine the location of a device by measuring the signal strength from multiple antennas. Alternatively, in some examples, the surgical hub 106 can be

equipped with one or more motion sensor devices configured to detect a change in the position of the surgical hub 106.

[0728] Referring to the bottom left corner of FIG. 33, the surgical hub 106 has been moved from its original position, which is depicted in dashed lines, to a new position closer to the device “D,” which is still outside the bounds of the operating room 3000. The surgical hub 106 in its new position, and based on the previously determined bounds of the operating room, would naturally conclude that the device “D” is a potential component of the surgical system 102. However, the introduction of a new device is a triggering event for reevaluation 3017 of the bounds of the operating room, as illustrated in the example algorithm of FIGS. 35, 37. After performing the reevaluation, the surgical hub 106 determines that the operating room bounds have changed. Based on the new bounds, at distances  $a_{new}$ ,  $-a_{new}$ ,  $b_{new}$ , and  $-b_{new}$ , the surgical hub 106 concludes that it has been moved and that the Device “D” is outside the newly determined bounds of its operating room. Accordingly, the surgical hub 106 will still not pair with the Device “D.”

[0729] In one aspect, one or more of the processes depicted in FIGS. 35-39 can be executed by a control circuit of a surgical hub 106, as depicted in FIG. 10 (processor 244). In another aspect, one or more of the processes depicted in FIGS. 35-39 can be executed by a cloud computing system 104, as depicted in FIG. 1. In yet another aspect, one or more of the processes depicted in FIGS. 35-39 can be executed by at least one of the aforementioned cloud computing systems 104 and/or a control circuit of a surgical hub 106 in combination with a control circuit of a modular device, such as the microcontroller 461 of the surgical instrument depicted in FIG. 12, the microcontroller 620 of the surgical instrument depicted in FIG. 16, the control circuit 710 of the robotic surgical instrument 700 depicted in FIG. 17, the control circuit 760 of the surgical instruments 750, 790 depicted in FIGS. 18-19, or the controller 838 of the generator 800 depicted in FIG. 20.

#### Spatial Awareness of Surgical Hubs in Operating Rooms

[0730] During a surgical procedure, a surgical instrument such as an ultrasonic or an RF surgical instrument can be coupled to a generator module 140 of the surgical hub 106. In addition, a separate surgical instrument controller such as a foot, or hand, switch or activation device can be used by an operator of the surgical instrument to activate the energy flow from the generator to the surgical instrument. Multiple surgical instrument controllers and multiple surgical instruments can be used concurrently in an operating room. Pressing or activating the wrong surgical instrument controller can lead to undesirable consequences. Aspects of the present disclosure present a solution in which the surgical hub 106 coordinates the pairing of surgical instrument controllers and surgical instruments to ensure patient and operator safety.

[0731] Aspects of the present disclosure are presented for a surgical hub 106 configured to establish and sever pairings between components of the surgical system 102 within the bounds of the operating room to coordinate flow of information and control actions between such components. The surgical hub 106 can be configured to establish a pairing

between a surgical instrument controller and a surgical instrument that reside within the bounds of an operating room of surgical hub 106.

[0732] In various aspects, the surgical hub 106 can be configured to establish and sever pairings between components of the surgical system 102 based on operator request or situational and/or spatial awareness. The hub situational awareness is described in greater detail below in connection with FIG. 86.

[0733] Aspects of the present disclosure are presented for a surgical hub for use with a surgical system in a surgical procedure performed in an operating room. The surgical hub includes a control circuit that selectively forms and severs pairings between devices of the surgical system. In one aspect, the hub includes a control circuit is configured to pair the hub with a first device of the surgical system, assign a first identifier to the first device, pair the hub with a second device of the surgical system, assign a second identifier to the second device, and selectively pair the first device with the second device. In one aspect, the surgical hub includes a storage medium, wherein the control circuit is configured to store a record indicative of the pairing between the first device and the second device in the storage medium. In one aspect, the pairing between the first device and the second device defines a communication pathway therebetween. In one aspect, the pairing between the first device and the second device defines a control pathway for transmitting control actions from the second device to the first device.

[0734] Further to the above, in one aspect, the control circuit is further configured to pair the hub with a third device of the surgical system, assign a third identifier to the third device, sever the pairing between the first device and the second device, and selectively pair the first device with the third device. In one aspect, the control circuit is further configured to store a record indicative of the pairing between the first device and the third device in the storage medium. In one aspect, the pairing between the first device and the third device defines a communication pathway therebetween. In one aspect, the pairing between the first device and the third device defines a control pathway for transmitting control actions from the third device to the first device.

[0735] In various aspects, the surgical hub includes a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to selectively form and sever pairings between the devices of the surgical system, as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer-readable instructions which, when executed, cause a machine to selectively form and sever pairings between the devices of the surgical system, as described above. FIGS. 40 and 41 are logic flow diagrams of processes depicting control programs or logic configurations for selectively forming and severing pairings between the devices of the surgical system, as described above.

[0736] In one aspect, the surgical hub 106 establishes a first pairing with a surgical instrument and a second pairing with the surgical instrument controller. The surgical hub 106 then links the pairings together allowing the surgical instrument and the surgical instrument controller to operate with one another. In another aspect, the surgical hub 106 may sever an existing communication link between a surgical instrument and a surgical instrument controller, then link the



surgical instrument to another surgical instrument controller that is linked to the surgical hub 106.

[0737] In one aspect, the surgical instrument controller is paired to two sources. First, the surgical instrument controller is paired to the surgical hub 106, which includes the generator module 140, for control of its activation. Second, the surgical instrument controller is also paired to a specific surgical instrument to prevent inadvertent activation of the wrong surgical instrument.

[0738] Referring to FIGS. 40 and 42, the surgical hub 106 may cause the communication module 130 to pair 3100 or establish a first communication link 3101 with a first device 3102 of the surgical system 102, which can be a first surgical instrument. Then, the hub may assign 3104 a first identification number to the first device 3102. This is a unique identification and communication sequence or number that may include the device's name and a time stamp of when the communication was first established.

[0739] In addition, the surgical hub 106 may then cause the communication module 130 to pair 3106 or establish a second communication link 3107 with a second device 3108 of the surgical system 102, which can be a surgical instrument controller. The surgical hub 106 then assigns 3110 a second identification number to the second device 3108.

[0740] In various aspects, the steps of pairing a surgical hub 106 with a device may include detecting the presence of a new device, determining that the new device is within bounds of the operating room, as described above in greater detail, and only pairing with the new device if the new device is located within the bounds of the operating room.

[0741] The surgical hub 106 may then pair 3112 or authorize a communication link 3114 to be established between the first device 3102 and the second device 3108, as illustrated in FIG. 42. A record indicative of the communication link 3114 is stored by the surgical hub 106 in the storage array 134. In one aspect, the communication link 3114 is established through the surgical hub 106. In another aspect, as illustrated in FIG. 42, the communication link 3114 is a direct link between the first device 3102 and the second device 3108.

[0742] Referring to FIGS. 41 and 43, the surgical hub 106 may then detect and pair 3120 or establish a third communication link 3124 with a third device 3116 of the surgical system 102, which can be another surgical instrument controller, for example. The surgical hub 106 may then assign 3126 a third identification number to the third device 3116.

[0743] In certain aspects, as illustrated in FIG. 43, the surgical hub 106 may then pair 3130 or authorize a communication link 3118 to be established between the first device 3102 and the third device 3116, while causing the communication link 3114 to be severed 3128, as illustrated in FIG. 43. A record indicative of the formation of the communication link 3118 and severing of the communication link 3114 is stored by the surgical hub 106 in the storage array 134. In one aspect, the communication link 3118 is established through the surgical hub 106. In another aspect, as illustrated in FIG. 43, the communication link 3118 is a direct link between the first device 3102 and the third device 3116.

[0744] As described above, the surgical hub 106 can manage an indirect communication between devices of the surgical system 102. For example, in situations where the first device 3102 is a surgical instrument and the second device 3108 is a surgical instrument controller, an output of

the surgical instrument controller can be transmitted through the communication link 3107 to the surgical hub 106, which may then transmit the output to the surgical instrument through the communication link 3101.

[0745] In making a decision to connect or sever a connection between devices of the surgical system 102, the surgical hub 106 may rely on perioperative data received or generated by the surgical hub 106. Perioperative data includes operator input, hub-situational awareness, hub-spatial awareness, and/or cloud data. For example, a request can be transmitted to the surgical hub 106 from an operator user-interface to assign a surgical instrument controller to a surgical instrument. If the surgical hub 106 determines that the surgical instrument controller is already connected to another surgical instrument, the surgical hub 106 may sever the connection and establish a new connection per the operator's request.

[0746] In certain examples, the surgical hub 106 may establish a first communication link between the visualization system 108 and the primary display 119 to transmit an image, or other information, from the visualization system 108, which resides outside the sterile field, to the primary display 119, which is located within the sterile field. The surgical hub 106 may then sever the first communication link and establish a second communication link between a robotic hub 122 and the primary display 119 to transmit another image, or other information, from the robotic hub 122 to the primary display 119, for example. The ability of the surgical hub 106 to assign and reassign the primary display 119 to different components of the surgical system 102 allows the surgical hub 106 to manage the information flow within the operating room, particularly between components inside the sterile field and outside the sterile field, without physically moving these components.

[0747] In another example that involves the hub-situational awareness, the surgical hub 106 may selectively connect or disconnect devices of the surgical system 102 within an operating room based on the type of surgical procedure being performed or based on a determination of an upcoming step of the surgical procedure that requires the devices to be connected or disconnected. The hub situational awareness is described in greater detail below in connection with FIG. 86.

[0748] Referring to FIG. 44, the surgical hub 106 may track 3140 the progression of surgical steps in a surgical procedure and may coordinate pairing and unpairing of the devices of the surgical system 102 based upon such progression. For example, the surgical hub 106 may determine that a first surgical step requires use of a first surgical instrument, while a second surgical step, occurring after completion of the first surgical step, requires use of a second surgical instrument. Accordingly, the surgical hub 106 may assign a surgical instrument controller to the first surgical instrument for the duration of the first surgical step. After detecting completion 3142 of the first surgical step, the surgical hub 106 may cause the communication link between the first surgical instrument and the surgical instrument controller to be severed 3144. The surgical hub 106 may then assign the surgical instrument controller to the second surgical instrument by pairing 3146 or authorizing the establishment of a communication link between the surgical instrument controller and the second surgical instrument.

[0749] Various other examples of the hub-situational awareness, which can influence the decision to connect or disconnect devices of the surgical system 102, are described in greater detail below in connection with FIG. 86.

[0750] In certain aspects, the surgical hub 106 may utilize its spatial awareness capabilities, as described in greater detail elsewhere herein, to track progression of the surgical steps of a surgical procedure and autonomously reassign a surgical instrument controller from one surgical instrument to another surgical instrument within the operating room of the surgical hub 106. In one aspect, the surgical hub 106 uses Bluetooth pairing and compass information to determine the physical position of the components of the surgical system 102.

[0751] In the example illustrated in FIG. 2, the surgical hub 106 is paired with a first surgical instrument held by a surgical operator at the operating table and a second surgical instrument positioned on a side tray. A surgical instrument controller can be selectively paired with either the first surgical instrument or the second surgical instrument. Utilizing the Bluetooth pairing and compass information, the surgical hub 106 autonomously assigns the surgical instrument controller to the first surgical instrument because of its proximity to the patient.

[0752] After completion of the surgical step that involved using the first surgical instrument, the first surgical instrument may be returned to the side tray or otherwise moved away from the patient. Detecting a change in the position of the first surgical instrument, the surgical hub 106 may sever the communication link between the first surgical instrument and the surgical instrument controller to protect against unintended activation of the first surgical instrument by the surgical instrument controller. The surgical hub 106 may also reassign the surgical instrument controller to another surgical instrument if the surgical hub 106 detects that it has been moved to a new position at the operating table.

[0753] In various aspects, devices of the surgical system 102 are equipped with an easy hand-off operation mode that would allow one user to give activation control of a device they currently control to another surgical instrument controller within reach of another operator. In one aspect, the devices are equipped to accomplish the hand-off through a predetermined activation sequence of the devices that causes the devices that are activated in the predetermined activation sequence to pair with one another.

[0754] In one aspect, the activation sequence is accomplished by powering on the devices to be paired with one another in a particular order. In another aspect, the activation sequence is accomplished by powering on the devices to be paired with one another within a predetermined time period. In one aspect, the activation sequence is accomplished by activating communication components, such as Bluetooth, of the devices to be paired with one another in a particular order. In another aspect, the activation sequence is accomplished by activating communication components, such as Bluetooth, of the devices to be paired within one another within a predetermined time period.

[0755] Alternatively, the hand-off can also be accomplished by a selection of a device through one of the surgical-operator input devices. After the selection is completed, the next activation by another controller would allow the new controller to take control.

[0756] In various aspects, the surgical hub 106 can be configured to directly identify components of the surgical

system 102 as they are brought into an operating room. In one aspect, the devices of the surgical system 102 can be equipped with an identifier recognizable by the surgical hub 106, such as, for example, a bar code or an RFID tag. NFC can also be employed. The surgical hub 106 can be equipped with a suitable reader or scanner for detecting the devices brought into the operating room.

[0757] The surgical hub 106 can also be configured to check and/or update various control programs of the devices of the surgical system 102. Upon detecting and establishing a communication link of a device of the surgical system 102, the surgical hub 106 may check if its control program is up to date. If the surgical hub 106 determines that a later version of the control program is available, the surgical hub 106 may download the latest version from the cloud 104 and may update the device to the latest version. The surgical hub 106 may issue a sequential identification and communication number to each paired or connected device.

#### Cooperative Utilization of Data Derived from Secondary Sources by Intelligent Surgical Hubs

[0758] In a surgical procedure, the attention of a surgical operator must be focused on the tasks at hand. Receiving information from multiple sources, such as, for example, multiple displays, although helpful, can also be distracting. The imaging module 138 of the surgical hub 106 is configured to intelligently gather, analyze, organize/package, and disseminate relevant information to the surgical operator in a manner that minimizes distractions.

[0759] Aspects of the present disclosure are presented for cooperative utilization of data derived from multiple sources, such as, for example, an imaging module 138 of the surgical hub 106. In one aspect, the imaging module 138 is configured to overlay data derived from one or more sources onto a livestream destined for the primary display 119, for example. In one aspect, the overlaid data can be derived from one or more frames acquired by the imaging module 138. The imaging module 138 may commandeer image frames on their way for display on a local display such as, for example, the primary display 119. The imaging module 138 also comprises an image processor that may preform an array of local image processing on the commandeered images.

[0760] Furthermore, a surgical procedure generally includes a number of surgical tasks which can be performed by one or more surgical instruments guided by a surgical operator or a surgical robot, for example. Success or failure of a surgical procedure depends on the success or failure of each of the surgical tasks. Without relevant data on the individual surgical tasks, determining the reason for a failed surgical procedure is a question of probability.

[0761] Aspects of the present disclosure are presented for capturing one or more frames of a livestream of a surgical procedure for further processing and/or pairing with other data. The frames may be captured at the completion of a surgical task (also referred to elsewhere herein as "surgical step") to assess whether the surgical task was completed successfully. Furthermore, the frames, and the paired data, can be uploaded to the cloud for further analysis.

[0762] In one aspect, one or more captured images are used to identify at least one previously completed surgical task to evaluate the outcome of the surgical task. In one aspect, the surgical task is a tissue-stapling task. In another aspect, the surgical task is an advanced energy transection.

[0763] FIG. 45 is a logic flow diagram of a process 3210 depicting a control program or a logic configuration for overlaying information derived from one or more still frames of a livestream of a remote surgical site onto the livestream. The process 3210 includes receiving 3212 a livestream of a remote surgical site from a medical imaging device 124, for example, capturing 3214 at least one image frame of a surgical step of the surgical procedure from the livestream, deriving 3216 information relevant to the surgical step from data extracted from the at least one image frame, and overlaying 3218 the information onto the livestream.

[0764] In one aspect, the still frames can be of a surgical step performed at the remote surgical site. The still frames can be analyzed for information regarding completion of the surgical step. In one aspect, the surgical step comprises stapling tissue at the surgical site. In another aspect, the surgical task comprises applying energy to tissue at the surgical site.

[0765] FIG. 46 is a logic flow diagram of a process 3220 depicting a control program or a logic configuration for differentiating among surgical steps of a surgical procedure. The process 3220 includes receiving 3222 a livestream of a surgical site from a medical imaging device 124, for example, capturing 3224 at least one first image frame of a first surgical step of the surgical procedure from the livestream, deriving 3226 information relevant to the first surgical step from data extracted from the at least one image frame, capturing 3228 at least one second image frame of a second surgical step of the surgical procedure from the livestream, and differentiating 3229 among the first surgical step and the second surgical step based on the at least one first image frame and the at least one second image frame.

[0766] FIG. 47 is a logic flow diagram of a process 3230 depicting a control program or a logic configuration for differentiating among surgical steps of a surgical procedure. The process 3232 includes receiving 3232 a livestream of the surgical site from a medical imaging device 124, for example, capturing 3234 image frames of the surgical steps of the surgical procedure from the livestream and differentiating 3236 among the surgical steps based on data extracted from the image frames.

[0767] FIG. 48 is a logic flow diagram of a process 3240 depicting a control program or a logic configuration for identifying a staple cartridge from information derived from one or more still frames of staples deployed from the staple cartridge into tissue. The process 3240 includes receiving 3242 a livestream of the surgical site from medical imaging device 124, for example, capturing 3244 an image frame from the livestream, detecting 3246 a staple pattern in the image frame, wherein the staple pattern is defined by staples deployed from a staple cartridge into tissue at the surgical site. The process 3240 further includes identifying 3248 the staple cartridge based on the staple pattern.

[0768] In various aspects, one or more of the steps of the processes 3210, 3220, 3230, 3240 can be executed by a control circuit of an imaging module of a surgical hub, as depicted in FIGS. 3, 9, 10. In certain examples, the control circuit may include a processor and a memory coupled to the processor, wherein the memory stores instructions executable by the processor to perform one or more of the steps of the processes 3210, 3220, 3230, 3240. In certain examples, a non-transitory computer-readable medium stores computer-readable instructions which, when executed, cause a

machine to perform one or more of the steps of the processes 3210, 3220, 3230, 3240. For economy, the following description of the processes 3210, 3220, 3230, 3240 will be described as being executed by the control circuit of an imaging module of a surgical hub; however, it should be understood that the execution of the processes 3210, 3220, 3230, 3240 can be accomplished by any of the aforementioned examples.

[0769] Referring to FIGS. 34 and 49, a surgical hub 106 is in communication with a medical imaging device 124 located at a remote surgical site during a surgical procedure. The imaging module 138 receives a livestream of the remote surgical site transmitted by the imaging device 124 to a primary display 119, for example, in accordance with steps 3212, 3222, 3232, 3242.

[0770] Further to the above, the imaging module 138 of the surgical hub 106 includes a frame grabber 3200. The frame grabber 3200 is configured to capture (i.e., “grabs”) individual, digital still frames from the livestream transmitted by the imaging device 124, for example, to a primary display 119, for example, during a surgical procedure, in accordance with steps 3214, 3224, 3234, 3244. The captured still frames are stored and processed by a computer platform 3203 (FIG. 49) of the imaging module 138 to derive information about the surgical procedure. Processing of the captured frames may include performance of simple operations, such as histogram calculations, 2D filtering, and arithmetic operations on arrays of pixels to the performance of more complex tasks, such as object detection, 3D filtering, and the like.

[0771] In one aspect, the derived information can be overlaid onto the livestream. In one aspect, the still frames and/or the information resulting from processing the still frames can be communicated to a cloud 104 for data aggregation and further analysis.

[0772] In various aspects, the frame grabber 3200 may include a digital video decoder and a memory for storing the acquired still frames, such as, for example, a frame buffer. The frame grabber 3200 may also include a bus interface through which a processor can control the acquisition and access the data and a general purpose I/O for triggering image acquisition or controlling external equipment.

[0773] As described above, the imaging device 124 can be in the form of an endoscope, including a camera and a light source positioned at a remote surgical site, and configured to provide a livestream of the remote surgical site at the primary display 119, for example.

[0774] In various aspects, image recognition algorithms can be implemented to identify features or objects in still frames of a surgical site that are captured by the frame grabber 3200. Useful information pertaining to the surgical steps associated with the captured frames can be derived from the identified features. For example, identification of staples in the captured frames indicates that a tissue-stapling surgical step has been performed at the surgical site. The type, color, arrangement, and size of the identified staples can also be used to derive useful information regarding the staple cartridge and the surgical instrument employed to deploy the staples. As described above, such information can be overlaid on a livestream directed to a primary display 119 in the operating room.

[0775] The image recognition algorithms can be performed at least in part locally by the computer platform 3203 (FIG. 49) of the imaging module 138. In certain instances,

the image recognition algorithms can be performed at least in part by the processor module 132 of the surgical hub 106. An image database can be utilized in performance of the image recognition algorithms and can be stored in a memory 3202 of the computer platform 3203. Alternatively, the imaging database can be stored in the storage array 134 (FIG. 3) of the surgical hub 106. The image database can be updated from the cloud 104.

[0776] An example image recognition algorithm that can be executed by the computer platform 3203 may include a key points-based comparison and a region-based color comparison. The algorithm includes: receiving an input at a processing device, such as, for example, the computer platform 3203; the input, including data related to a still frame of a remote surgical site; performing a retrieving step, including retrieving an image from an image database and, until the image is either accepted or rejected, designating the image as a candidate image; performing an image recognition step, including using the processing device to perform an image recognition algorithm on the still frame and candidate images in order to obtain an image recognition algorithm output; and performing a comparison step, including: if the image recognition algorithm output is within a pre-selected range, accepting the candidate image as the still frame and if the image recognition algorithm output is not within the pre-selected range, rejecting the candidate image and repeating the retrieving, image recognition, and comparison steps.

[0777] Referring to FIGS. 50-52, in one example, a surgical step involves stapling and cutting tissue. FIG. 50 depicts a still frame 3250 of a stapled and cut tissue T. A staple deployment 3252 includes staples 3252', 3252" from a first staple cartridge. A second staple deployment 3254 includes staples 3254', 3254" from a second staple cartridge. A proximal portion 3253 of the staple deployment 3252 overlaps with a distal portion 3255 of the staple deployment 3254. Six rows of staples were deployed in each deployment. Tissue T was cut between the third and fourth rows of each deployment, but only one side of the stapled tissue T is fully shown.

[0778] In various aspects, the imaging module 138 identifies one or more of the staples 3252', 3252", 3254', 3254" in the still frame 3250, which were absent in a previous still frame captured by the frame grabber 3200. The imaging module 138 then concludes that a surgical stapling and cutting instrument has been used at the surgical site.

[0779] In the example of FIG. 50, the staple deployment 3252 includes two different staples 3252', 3252". Likewise, the staple deployment 3254 includes two different staples 3254', 3254". For brevity, the following description focuses on the staples 3252', 3252", but is equally applicable to the staples 3254', 3254". The staples 3252', 3252" are arranged in a predetermined pattern or sequence that forms a unique identifier corresponding to the staple cartridge that housed the staples 3252', 3252". The unique pattern can be in a single row or multiple rows of the staples 3250. In one example, the unique pattern can be achieved by alternating the staples 3252', 3252" at a predetermined arrangement.

[0780] In one aspect, multiple patterns can be detected in a firing of staples. Each pattern can be associated with a unique characteristic of the staples, the staple cartridge that housed the staples, and/or the surgical instrument that was

employed to fire the staple. For example, a firing of staples may include patterns that represent staple form, staple size, and/or location of the firing.

[0781] In the example, of FIG. 50, the imaging module 138 may identify a unique pattern of the staples 3252 from the still frame 3250. A database storing staple patterns and corresponding identification numbers of staple cartridges can then be explored to determine an identification number of a staple cartridge that housed the staples 3252.

[0782] The patterns of the example of FIG. 50 are based on only two different staples; however, other aspects may include three or more different staples. The different staples can be coated with different coatings, which can be applied to the staples by one or more of the following methods: anodizing, dyeing, electro-coating, photoluminescent coating, application of nitrides, methyl methacrylate, painting, powder coating, coating with paraffins, oil stains or phosphor coatings, the use of hydroxyapatite, polymers, titanium oxinitrides, zinc sulfides, carbides, etc. It should be noted that, while the listed coatings are fairly specific as disclosed herein, other coatings known in the art to distinguish the staple are within the contemplated scope of the present disclosure.

[0783] In the example of FIGS. 50-52, the staples 3252' are anodized staples, while the staples 3252" are non-anodized staples. In one aspect, the different staples may comprise two or more different colors. Different metal staples may comprise magnetic or radioactive staple markers that differentiate them from unmarked staples.

[0784] FIG. 51 illustrates a staple deployment 3272 deployed into tissue from a staple cartridge via a surgical instrument. Only three staple rows 3272a, 3272b, 3272c are depicted in FIG. 51. The rows 3272a, 3272b, 3272c are arranged between a medial line, where the tissue was cut, and a lateral line at the tissue edge. For clarity, the inner row 3272a of staples is redrawn separately to the left and the outer two rows 3272b, 3272c are redrawn separately to the right. A proximal end 3273 and a distal end portion of the staple deployment 3272 are also redrawn in FIG. 51 for clarity.

[0785] The staple deployment 3272 includes two different staples 3272', 3272" that are arranged in predetermined patterns that serve various functions. For example, the inner row 3272a comprises a pattern of alternating staples 3272', 3272", which defines a metric for distance measurements in the surgical field. In other words, the pattern of the inner row 3272a acts as a ruler for measuring distances, which can be helpful in accurately determining the position of a leak, for example. The outer rows 3272b, 3272c define a pattern that represents an identification number of the staple cartridge that housed the staples 3272', 3272".

[0786] Furthermore, unique patterns at the ends of the staple deployment 3272 identify the proximal end portion 3273 and distal end portion 3275. In the example of FIG. 51, a unique arrangement of three staples 3272" identifies the distal end 3275, while a unique arrangement of four staples 3272" identifies the proximal end 3273. Identification of the proximal and distal ends of a staple deployment allows the imaging module 128 to distinguish between different staple deployments within a captured frame, which can be useful in pointing the source of a leak, for example.

[0787] In various aspects, the imaging module 138 may detect a sealed tissue in a still frame of a remote surgical site captured by the frame grabber 3200. Detection of the sealed

tissue can be indicative of a surgical step that involves applying therapeutic energy to tissue.

**[0788]** Sealing tissue can be accomplished by the application of energy, such as electrical energy, for example, to tissue captured or clamped within an end effector of a surgical instrument in order to cause thermal effects within the tissue. Various mono-polar and bi-polar RF surgical instruments and harmonic surgical instruments have been developed for such purposes. In general, the delivery of energy to captured tissue can elevate the temperature of the tissue and, as a result, the energy can at least partially denature proteins within the tissue. Such proteins, like collagen, for example, can be denatured into a proteinaceous amalgam that intermixes and fuses, or seals, together as the proteins renature.

**[0789]** Accordingly, sealed tissue has a distinct color and/or shape that can be detected by the imaging module **138** using image recognition algorithms, for example. In addition, smoke detection at the surgical site can indicate that therapeutic energy application to the tissue is in progress.

**[0790]** Further to the above, the imaging module **138** of the surgical hub **106** is capable of differentiating between surgical steps of a surgical procedure based on the captured frames. As described above, a still frame that comprises fired staples is indicative of a surgical step involving tissue stapling, while a still frame that comprises a sealed tissue is indicative of a surgical step involving energy application to tissue.

**[0791]** In one aspect, the surgical hub **106** may selectively overlay information relevant to a previously completed surgical task onto the livestream. For example, the overlaid information may comprise image data from a still frame of the surgical site captured during the previously completed surgical task. Furthermore, guided by common landmark locations at the surgical site, the imaging module **138** can interlace one image frame to another to establish and detect surgical locations and relationship data of a previously completed surgical task.

**[0792]** In one example, the surgical hub **106** is configured to overlay information regarding a potential leak in a tissue treated by stapling or application of therapeutic energy in a previously completed surgical task. The potential leak can be spotted by the imaging module **138** during the processing of a still frame of the tissue. The surgical operator can be alerted about the leak by overlaying information about the potential leak onto the livestream.

**[0793]** In various aspects, still frames of an end effector of a surgical instrument at a surgical site can be used to identify the surgical instrument. For example, the end effector may include an identification number that can be recognized by the imaging module **138** during image processing of the still frame. Accordingly, the still frames captured by the imaging module **138** may be used to identify a surgical instrument utilized in a surgical step of a surgical procedure. The still frames may also include useful information regarding the performance of the surgical instrument. All such information can be uploaded to the cloud **104** for data aggregation and further analysis.

**[0794]** In various examples, the surgical hub **106** may also selectively overlay information relevant to a current or upcoming surgical task, such as an anatomical location or a surgical instrument suitable for the surgical task.

**[0795]** The imaging module **138** may employ various images and edge detection techniques to track a surgical site

where a surgical instrument was used to complete a surgical task. Success or failure of the surgical task can then be assessed. For example, a surgical instrument can be employed to seal and/or cut tissue at the surgical site. A still frame of the surgical site can be stored in the memory **3202** or the storage array **134** of the surgical hub **106**, for example, upon completion of the surgical task.

**[0796]** In the following surgical step, the quality of the seal can be tested via different mechanisms. To ensure that the testing is accurately applied to the treated tissue, the stored still frame of the surgical site is overlaid onto the livestream in search of a match. Once a match is found, the testing can take place. One or more additional still frames can be taken during the testing, which can be later analyzed by the imaging module **138** of the surgical hub **106**. The testing mechanisms include bubble detection, bleeding detection, dye detection (where a dye is employed at the surgical site), and/or burst stretch detection (where a localized strain is applied adjacent to an anastomosis site), for example.

**[0797]** The imaging module **138** may capture still frames of the response of the treated tissue to these tests, which can be stored in the memory **3202** or the storage array **134** of the surgical hub **106**, for example. The still frames can be stored alone or in combination with other data, such as, for example, data from the surgical instrument that performed the tissue treatment. The paired data can also be uploaded to the cloud **104** for additional analysis and/or pairing.

**[0798]** In various aspects, the still frames captured by the frame grabber **3200** can be processed locally, paired with other data, and can also be transmitted to the cloud **104**. The size of the processed and/or transmitted data will depend on the number of captured frames. In various aspects, the rate at which the frame grabber **3200** captures the still frames from the livestream can be varied in an effort to reduce the size of the data without sacrificing quality.

**[0799]** In one aspect, the frame-capturing rate may depend on the type of surgical task being performed. Certain surgical tasks may need a higher number of still frames than others for an evaluation of success or failure. The frame-capturing rate can be scaled to accommodate such needs.

**[0800]** In one aspect, the frame-capturing rate is dependent upon the detected motion of the imaging device **124**. In use, an imaging device **124** may target one surgical site for a period of time. Observing no or minor changes in the still frames captured while the imaging device **124** is not being moved, the imaging module **138** may reduce the frame-capturing rate of the frame grabber **3200**. If the situation changes, however, where frequent motion is detected, the imaging module **138** may respond by increasing the frame-capturing rate of the frame grabber **3200**. In other words, the imaging module **138** may be configured to correlate the frame-capturing rate of the frame grabber **3200** with the detected degree of motion of the imaging device **124**.

**[0801]** For additional efficiency, only portions of the still frames, where motion is detected, need to be stored, processed, and/or transmitted to the cloud **104**. The imaging module **138** can be configured to select the portions of the still frames where motion is detected. In one example, motion detection can be achieved by comparing a still frame to a previously captured still frame. If movement is detected, the imaging module **138** may cause the frame grabber **3200**

to increase the frame-capturing rate, but only the portions where motion is detected are stored, processed, and/or transmitted to the cloud **104**.

**[0802]** In another aspect, the data size can be managed by scaling the resolution of the captured information based on the area of the screen where the focal point is or where end effectors are located, for example. The remainder of the screen could be captured at a lower resolution.

**[0803]** In one aspect, the corners of the screen and the edges could generally be captured at a lower resolution. The resolution, however, can be scaled up if an event of significance is observed.

**[0804]** During a surgical procedure, the surgical hub **106** can be connected to various operating-room monitoring devices, such as, for example, heart rate monitors and insufflation pumps. Data collected from these devices can improve the situational awareness of the surgical hub **106**. The hub situational awareness is described in greater detail below in connection with FIG. **86**.

**[0805]** In one example, the surgical hub **106** can be configured to utilize patient data received from a heart rate monitor connected along with data regarding the location of the surgical site to assess proximity of the surgical site to sensory nerves. An increase in the patient's heart rate, when combined with anatomical data indicating that the surgical site is in a region high in sensory nerves, can be construed as an indication of sensory nerve proximity. Anatomical data can be available to the surgical hub **106** through accessing patient records (e.g., an EMR database containing patient records).

**[0806]** The surgical hub **106** may be configured to determine the type of surgical procedure being performed on a patient from data received from one or more of the operating-room monitoring devices, such as, for example, heart rate monitors and insufflation pumps. Abdominal surgical procedures generally require insufflation of the abdomen, while insufflation is not required in theoretic surgery. The surgical hub **106** can be configured to determine whether a surgical procedure is an abdominal or a thoracic surgical procedure by detecting whether the insufflation pump is active. In one aspect, the surgical hub **106** may be configured to monitor insufflation pressure on the output side of the insufflation pump in order to determine whether the surgical procedure being performed is one that requires insufflation.

**[0807]** The surgical hub **106** may also gather information from other secondary devices in the operating room to assess, for example, whether the surgical procedure is a vascular or avascular procedure.

**[0808]** The surgical hub **106** may also monitor AC current supply to one or more of its components to assess whether a component is active. In one example, the surgical hub **106** is configured to monitor AC current supply to the generator module to assess whether the generator is active, which can be an indication that the surgical procedure being performed is one that requires application of energy to seal tissue.

**[0809]** In various aspects, secondary devices in the operating room that are incapable of communication with the surgical hub **106** can be equipped with communication interface devices (communication modules) that can facilitate pairing of these devices with the surgical hub **106**. In one aspect, the communication interface devices may be configured to be bridging elements, which would allow them two-way communication between the surgical hub **106** and such devices.

**[0810]** In one aspect, the surgical hub **106** can be configured to control one or more operational parameters of a secondary device through a communication interface device. For example, the surgical hub **106** can be configured to increase or decrease the insufflation pressure through a communication interface device coupled to an insufflation device.

**[0811]** In one aspect, the communication interface device can be configured to engage with an interface port of the device. In another aspect, the communication interface device may comprise an overlay or other interface that directly interacts with a control panel of the secondary device. In other aspects, the secondary devices, such as, for example, the heart rate monitor and/or the insufflation devices, can be equipped with integrated communication modules that allow them to pair with the hub for two-way communication therewith.

**[0812]** In one aspect, the surgical hub **106** can also be connected through a communication interface device, for example, to muscle pads that are connected to the neuro-stim detection devices to improve resolution of a nerve-sensing device.

**[0813]** Furthermore, the surgical hub **106** can also be configured to manage operating room supplies. Different surgical procedures require different supplies. For example, two different surgical procedures may require different sets of surgical instruments. Certain surgical procedures may involve using a robotic system, while others may not. Furthermore, two different surgical procedures may require staple cartridges that are different in number, type, and/or size. Accordingly, the supplies brought into the operating room can provide clues as to the nature of the surgical procedure that will be performed.

**[0814]** In various aspects, the surgical hub **106** can be integrated with an operating room supplies scanner to identify items pulled into the operating room and introduced into the sterile field. The surgical hub **106** may utilize data from the operating room supplies scanner, along with data from the devices of the surgical system **102** that are paired with the surgical hub **106**, to autonomously determine the type of surgical procedure that will be performed. In one example, the surgical hub **106** may record a list of serial numbers of the smart cartridge that are going to be used in the surgical procedure. During the surgical procedure, the surgical hub **106** may gradually remove the staples that have been fired, based on information collected from the staple cartridge chips. In one aspect, the surgical hub **106** is configured to make sure that all the items are accounted for at the end of the procedure.

#### Surgical Hub Control Arrangements

**[0815]** In a surgical procedure, a second surgical hub may be brought into an operating room already under the control of a first surgical hub. The second surgical hub can be, for example, a surgical robotic hub brought into the operating room as a part of a robotic system. Without coordination between the first and second surgical hubs, the robotic surgical hub will attempt to pair with all the other components of the surgical system **102** that are within the operating room. The confusion arising from the competition between two hubs in a single operating room can lead to undesirable consequences. Also, sorting out the instrument distribution between the hubs during the surgical procedure can be time consuming.

[0816] Aspects of the present disclosure are presented for a surgical hub for use with a surgical system in a surgical procedure performed in an operating room. A control circuit of the surgical hub is configured to determine the bounds of the operating room and establish a control arrangement with a detected surgical hub located within the bounds of the operating room.

[0817] In one aspect, the control arrangement is a peer-to-peer arrangement. In another aspect, the control arrangement is a master-slave arrangement. In one aspect, the control circuit is configured to select one of a master mode of operation or a slave mode of operation in the master-slave arrangement. In one aspect, the control circuit is configured to surrender control of at least one surgical instrument to the detected surgical hub in the slave mode of operation.

[0818] In one aspect, the surgical hub includes an operating room mapping circuit that includes a plurality of non-contact sensors configured to measure the bounds of the operating room.

[0819] In various aspects, the surgical hub includes a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to coordinate a control arrangement between surgical hubs, as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer-readable instructions which, when executed, cause a machine to coordinate a control arrangement between surgical hubs, as described above.

[0820] Aspects of the present disclosure are presented for a surgical system comprising two independent surgical hubs that are configured to interact with one another. Each of the hubs has their own linked surgical devices and the control designation of and distribution of where data is recorded and processed. This interaction causes one or both hubs to change how they were behaving before the interaction. In one aspect, the change involves a redistribution of devices previously assigned to each of the hubs. In another aspect, the change involves establishing a master-slave arrangement between the hubs. In yet another aspect, the change can be a change in the location of the processing shared between the hubs.

[0821] FIG. 53 is a logic flow diagram of a process depicting a control program or a logic configuration for coordinating a control arrangement between surgical hubs. The process of FIG. 53 is similar in many respects to the process of FIG. 35 except that the process of FIG. 53 addresses detection of a surgical hub by another surgical hub. As illustrated in FIG. 53, the surgical hub 106 determines 3007 the bounds of the operating room. After the initial determination, the surgical hub 106 continuously searches for or detects 3008 devices within a pairing range. If a device is detected 3010, and if the detected device is located 3011 within the bounds of the operating room, the surgical hub 106 pairs 3012 with the device and assigns 3013 an identifier to the device. If through an initial interaction, as described below in greater detail, the surgical hub 106 determines 3039 that the device is another surgical hub, a control arrangement is established 3040 therebetween.

[0822] Referring to FIG. 54, a robotic surgical hub 3300 enters an operating room already occupied by a surgical hub 3300. The robotic surgical hub 3310 and the surgical hub 3300 are similar in many respects to other surgical hubs described in greater detail elsewhere herein, such as, for example, the surgical hubs 106. For example, the robotic

surgical hub 3310 includes non-contact sensors configured to measure the bounds of the operating room, as described in greater detail elsewhere herein in connection with FIGS. 33, 34.

[0823] As the robotic surgical hub 3310 is powered up, it determines the bounds of the operating room and begins to pair with other components of the surgical system 102 that are located within the bounds of the operating room. The robotic surgical hub 3310 pairs with a robotic advanced energy tool 3311, a robotic stapler 3312, a monopolar energy tool 3313, and a robotic visualization tower 3314, which are all located within the bounds of the operating room. The surgical hub 3300 is already paired with a handheld stapler 3301, a handheld powered dissector 3302, a secondary display 3303, a surgeon interface 3304, and a visualization tower 3305. Since the handheld stapler 3301, the handheld powered dissector 3302, the secondary display 3303, the surgeon interface 3304, and the visualization tower 3305 are already paired with the surgical hub 3300, such devices cannot pair with another surgical hub without permission from the surgical hub 3300.

[0824] Further to the above, the robotic surgical hub 3310 detects and/or is detected by the surgical hub 3300. A communication link is established between the communication modules of the surgical hubs 3300, 3310. The surgical hubs 3300, 3310 then determine the nature of their interaction by determining a control arrangement therebetween. In one aspect, the control arrangement can be a master-slave arrangement. In another aspect, the control arrangement can be a peer-to-peer arrangement.

[0825] In the example of FIG. 54, a master-slave arrangement is established. The surgical hubs 3300, 3310 request permission from a surgical operator for the robotic surgical hub 3310 to take control of the operating room from the surgical hub 3300. The permission can be requested through a surgeon interface or console 3304. Once permission is granted, the robotic surgical hub 3310 requests the surgical hub 3300 to transfer control to the robotic surgical hub 3310.

[0826] Alternatively, the surgical hubs 3300, 3310 can negotiate the nature of their interaction without external input based on previously gathered data. For example, the surgical hubs 3300, 3310 may collectively determine that the next surgical task requires use of a robotic system. Such determination may cause the surgical hub 3300 to autonomously surrender control of the operating room to the robotic surgical hub 3310. Upon completion of the surgical task, the robotic surgical hub 3310 may then autonomously return the control of the operating room to surgical hub 3300.

[0827] The outcome of the interaction between the surgical hubs 3300, 3310 is illustrated on the right of FIG. 54. The surgical hub 3300 has transferred control to the robotic surgical hub 3310, which has also taken control of the surgeon interface 3304 and the secondary display 3303 from the surgical hub 3300. The robotic surgical hub 3310 assigns new identification numbers to the newly transferred devices. The surgical hub 3300 retains control the handheld stapler 3301, the handheld powered dissector 3302, and visualization tower 3305. In addition, the surgical hub 3300 performs a supporting role, wherein the processing and storage capabilities of the surgical hub 3300 are now available to the robotic surgical hub 3310.

[0828] FIG. 55 is a logic flow diagram of a process depicting a control program or a logic configuration for

coordinating a control arrangement between surgical hubs. In various aspects, two independent surgical hubs will interact with one another in a predetermined manner to assess the nature of their relationship. In one example, after establishing **3321** a communication link, the surgical hubs exchange **3322** data packets. A data packet may include type, identification number, and/or status of a surgical hub. A data packet may further include a record of devices under control of the surgical hub and/or any limited communication connections, such as data ports for other secondary operating room devices.

**[0829]** The control arrangement between the surgical hubs is then determined **3323** based on input from a surgical operator or autonomously between the surgical hubs. The surgical hubs may store instructions as to how to determine a control arrangement therebetween. The control arrangement between two surgical hubs may depend on the type of surgical procedure being performed. The control arrangement between two surgical hubs may depend on their types, identification information, and/or status. The control arrangement between two surgical hubs may depend on the devices paired with the surgical hubs. The surgical hubs then redistribute **3324** the devices of the surgical system **102** therebetween based upon the determined control arrangement.

**[0830]** In the master-slave arrangement, the record communication can be unidirectional from the slave hub to the master hub. The master hub may also require the slave hub to hand-off some of its wireless devices to consolidate communication pathways. In one aspect, the slave hub can be relegated to a relay configuration with the master hub originating all commands and recording all data. The slave hub can remain linked to the master hub for a distributed sub-processing of the master hub commands, records, and/or controls. Such interaction expands the processing capacity of the dual linked hubs beyond the capabilities of the master hub by itself.

**[0831]** In a peer-to-peer arrangement, each surgical hub may retain control of its devices. In one aspect, the surgical hubs may cooperate in controlling a surgical instrument. In one aspect, an operator of the surgical instrument may designate the surgical hub that will control the surgical instrument at the time of its use.

**[0832]** Referring generally to FIGS. **56-61**, the interaction between surgical hubs can be extended beyond the bounds of the operating room. In various aspects, surgical hubs in separate operating rooms may interact with one another within predefined limits. Depending on their relative proximity, surgical hubs in separate operating rooms may interact through any suitable wired or wireless data communication network such as Bluetooth and WiFi. As used here, a “data communication network” represents any number of physical, virtual, or logical components, including hardware, software, firmware, and/or processing logic configured to support data communication between an originating component and a destination component, where data communication is carried out in accordance with one or more designated communication protocols over one or more designated communication media.

**[0833]** In various aspects, a first surgical operator in a first operating room may wish to consult a second surgical operator in a second operating room, such as in case of an emergency. A temporary communication link may be established between the surgical hubs of the first and second

operating room to facilitate the consult while the first and second surgical operators remain in their respective operating rooms.

**[0834]** The surgical operator being consulted can be presented with a consult request through the surgical hub in his/her operating room. If the surgical operator accepts, he/she will have access to all the data compiled by the surgical hub requesting the consult. The surgical operator may access all previously stored data, including a full history of the procedure. In addition, a livestream of the surgical site at the requesting operating room can be transmitted through the surgical hubs to a display at the receiving operating room.

**[0835]** When a consult request begins, the receiving surgical hub begins to record all received information in a temporarily storage location, which can be a dedicated portion of the storage array of the surgical hub. At the end of the consult, the temporary storage location is purged from all the information. In one aspect, during a consult, the surgical hub records all accessible data, including blood pressure, ventilation data, oxygen stats, generator settings and uses, and all patient electronic data. The recorded data will likely be more than the data stored by the surgical hub during normal operation, which is helpful in providing the surgical operator being consulted with as much information as possible for the consult.

**[0836]** Referring to FIG. **56**, a non-limiting example of an interaction between surgical hubs in different operating rooms is depicted. FIG. **56** depicts an operating room OR 1 that includes a surgical system **3400** supporting a thoracic segmentectomy and a second operating room OR 3 that includes a surgical system **3410** supporting a colorectal procedure. The surgical system **3400** includes surgical hub **3401**, surgical hub **3402**, and robotic surgical hub **3403**. The surgical system **3400** further includes a personal interface **3406**, a primary display **3408**, and secondary displays **3404**, **3405**. The surgical system **3410** includes a surgical hub **3411** and a secondary display **3412**. For clarity, several components of the surgical systems **3400**, **3410** are removed.

**[0837]** In the example of FIG. **56**, the surgical operator of OR 3 is requesting a consult from the surgical operator of OR 1. A surgical hub **3411** of the OR 3 transmits the consult request to one of the surgical hubs of the OR 1, such as the surgical hub **3401**. In OR 1, the surgical hub **3401** presents the request at a personal interface **3406** held by the surgical operator. The consult is regarding selecting an optimal location of a colon transection. The surgical operator of OR 1, through a personal interface **3406**, recommends an optimal location for the transection site that avoids a highly vascular section of the colon. The recommendation is transmitted in real time through the surgical hubs **3401**, **3411**. Accordingly, the surgical operator is able to respond to the consult request in real time without having to leave the sterile field of his own operating room. The surgical operator requesting the consult also did not have to leave the sterile field of OR 3.

**[0838]** If the surgical hub **3401** is not in communication with the personal interface **3406**, it may relay the message to another surgical hub such as, for example, the surgical hub **3402** or the robotic surgical hub **3403**. Alternatively, the surgical hub **3401** may request control of the personal interface **3406** from another surgical hub.

**[0839]** In any event, if the surgical operator of OR 1 decides to accept the consult request, a livestream, or



frames, of a surgical site **3413** of the colorectal procedure of OR 3 is transmitted to OR 1 through a connection established between the surgical hubs **3401**, **3411**, for example. FIG. **57** illustrates a livestream of the surgical site **3413** displayed on a secondary display of OR 3. The surgical hubs **3401**, **3411** cooperate to transmit the livestream of the surgical site of OR 3 to the personal interface **3406** of the OR 1, as illustrated in FIG. **58**.

[**0840**] Referring to FIGS. **59-61**, the surgical operator may expand the laparoscopic livestream from OR 3 onto the primary display **3405** in OR 1, for example, through the controls of the personal interface **3406**. The personal interface **3406** allows the surgical operator to select a destination for the livestream by presenting the surgical operator with icons that represent the displays that are available in OR 1, as illustrated in FIG. **60**. Other navigation controls **3407** are available to the surgical operator through the personal interface **3406**, as illustrated in FIG. **61**. For example, the personal interface **3406** includes navigation controls for adjusting the livestream of the surgical site of OR 3 in OR 1 by the surgical operator moving his or her fingers on the livestream displayed on the personal interface **3406**. To visualize the high vasculature regions, the consulted surgical operator may change the view of the livestream from OR 3 through the personal interface **3406** to an advanced imaging screen. The surgical operator may then manipulate the image in multiple planes to see the vascularization using a wide-angle multi-spectral view, for example.

[**0841**] As illustrated in FIG. **61**, the surgical operator also has access to an array of relevant information **3420**, such as, for example, heart rate, blood pressure, ventilation data, oxygen stats, generator settings and uses, and all patient electronic data of the patient in OR 3.

#### Data Management and Collection

[**0842**] In one aspect the surgical hub provides data storage capabilities. The data storage includes creation and use of self-describing data including identification features, management of redundant data sets, and storage of the data in a manner of paired data sets which can be grouped by surgery but not necessarily keyed to actual surgical dates and surgeons to maintain data anonymity. The following description incorporates by reference all of the “hub” and “cloud” analytics system hardware and software processing techniques to implement the specific data management and collection techniques described hereinbelow, as incorporated by reference herein. FIGS. **62-80** will be described in the context of the interactive surgical system **100** environment including a surgical hub **106**, **206** described in connection FIGS. **1-11** and intelligent instruments and generators described in connection with FIGS. **12-21**.

#### Electronic Medical Record (EMR) Interaction

[**0843**] FIG. **62** is a diagram **4000** illustrating a technique for interacting with a patient Electronic Medical Record (EMR) database **4002**, according to one aspect of the present disclosure. In one aspect, the present disclosure provides a method of embedding a key **4004** within the EMR database **4002** located within the hospital or medical facility. A data barrier **4006** is provided to preserve patient data privacy and allows the reintegration of stripped and isolated data pairs, as described hereinbelow, from the surgical hub **106**, **206** or the cloud **104**, **204**, to be reassembled. A schematic diagram

of the surgical hub **206** is described generally in FIGS. **1-11** and in particular in FIGS. **9-10**. Therefore, in the description of FIG. **62**, the reader is guided to FIG. **1-11** and in particular FIGS. **9-10** for any implementation details of the surgical hub **206** that may be omitted here for conciseness and clarity of disclosure. Returning to FIG. **62**, the method allows the users full access to all the data collected during a surgical procedure and patient information stored in the form of electronic medical records **4012**. The reassembled data can be displayed on a monitor **4010** coupled to the surgical hub **206** or secondary monitors but is not permanently stored on any surgical hub storage device **248**. The reassembled data is temporarily stored in a storage device **248** located either in the surgical hub **206** or the cloud **204** and is deleted at the end of its use and overwritten to insure it cannot be recovered. The key **4004** in the EMR database **4002** is used to reintegrate anonymized hub data back into full integrated patient electronic medical records **4012** data.

[**0844**] As shown in FIG. **62**, the EMR database **4002** is located within the hospital data barrier **4006**. The EMR database **4002** may be configured for storing, retrieving, and managing associative arrays, or other data structures known today as a dictionary or hash. Dictionaries contain a collection of objects, or records, which in turn have many different fields within them, each containing data. The patient electronic medical records **4012** may be stored and retrieved using a key **4004** that uniquely identifies the patient electronic medical record **4012**, and is used to quickly find the data within the EMR database **4002**. The key-value EMR database **4002** system treats the data as a single opaque collection which may have different fields for every record.

[**0845**] Information from the EMR database **4002** may be transmitted to the surgical hub **206** and the patient electronic medical records **4012** data is redacted and stripped before it is sent to an analytics system based either on the hub **206** or the cloud **204**. An anonymous data file **4016** is created by redacting personal patient data and stripping relevant patient data **4018** from the patient electronic medical record **4012**. As used herein, the redaction process includes deleting or removing personal patient information from the patient electronic medical record **4012** to create a redacted record that includes only anonymous patient data. A redacted record is a record from which sensitive patient information has been expunged. Un-redacted data may be deleted **4019**. The relevant patient data **4018** may be referred to herein as stripped/extracted data **4018**. The relevant patient data **4018** is used by the surgical hub **206** or cloud **204** processing engines for analytic purposes and may be stored on the storage device **248** of the surgical hub **206** or may be stored on the cloud **204** based analytics system storage device **205**. The surgical hub anonymous data file **4016** can be rebuilt using a key **4004** stored in the EMR database **4002** to reintegrate the surgical hub anonymous data file **4016** back into a fully integrated patient electronic medical record **4012**. The relevant patient data **4018** that is used in analytic processes may include information such as the patient’s diagnoses of emphysema, pre-operative treatment (e.g., chemotherapy, radiation, blood thinner, blood pressure medication, etc.), typical blood pressures, or any data that alone cannot be used to ascertain the identity of the patient. Data **4020** to be redacted includes personal information removed from the patient electronic medical record **4012**, may include age, employer, body mass index (BMI), or any data that can be used to ascertain the identity of the patient. The

surgical hub **206** creates a unique anonymous procedure ID number (e.g., 380i4z), for example, as described in FIG. **63**. Within the EMR database **4002** located in the hospital data barrier **4006**, the surgical hub **206** can reunite the data in the anonymous data file **4016** stored on the surgical hub **206** storage device **248** with the data in the patient electronic medical record **4012** stored on the EMR database **4002** for surgeon review. The surgical hub **206** displays the combined patient electronic medical record **4012** on a display or monitor **4010** coupled to the surgical hub **206**. Ultimately, un-redacted data is deleted **4019** from the surgical hub **206** storage **248**.

Creation of a Hospital Data Barrier, Inside which  
the Data from Hubs can be Compared Using  
Non-Anonymized Data and Outside of which the  
Data has to be Stripped

**[0846]** In one aspect, the present disclosure provides a surgical hub **206** as described in FIGS. **9** and **10**, for example, where the surgical hub **206** comprises a processor **244**; and a memory **249** coupled to the processor **244**. The memory **249** stores instructions executable by the processor **244** to interrogate a surgical instrument **235**, retrieve a first data set from the surgical instrument **235**, interrogate a medical imaging device **238**, retrieve a second data set from the medical imaging device **238**, associate the first and second data sets by a key, and transmit the associated first and second data sets to a remote network, e.g., the cloud **204**, outside of the surgical hub **206**. The surgical instrument **235** is a first source of patient data and the first data set is associated with a surgical procedure. The medical imaging device **238** is a second source of patient data and the second data set is associated with an outcome of the surgical procedure. The first and second data records are uniquely identified by the key.

**[0847]** In another aspect, the surgical hub **206** provides a memory **249** storing instructions executable by the processor **244** to retrieve the first data set using the key, anonymize the first data set, retrieve the second data set using the key, anonymize the second data set, pair the anonymized first and second data sets, and determine success rate of surgical procedures grouped by the surgical procedure based on the anonymized paired first and second data sets.

**[0848]** In another aspect, the surgical hub **206** provides a memory **249** storing instructions executable by the processor **244** to retrieve the anonymized first data set, retrieve the anonymized second data set, and reintegrate the anonymized first and second data sets using the key.

**[0849]** In another aspect, the first and second data sets define first and second data payloads in respective first and second data packets.

**[0850]** In various aspects, the present disclosure provides a control circuit to associate the first and second data sets by a key as described above. In various aspects, the present disclosure provides a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to associate the first and second data sets by a key as described above.

**[0851]** During a surgical procedure it would be desirable to monitor data associated with the surgical procedure to enable configuration and operation of instruments used during the procedure to improve surgical outcomes. The technical challenge is to retrieve the data in a manner that maintains the anonymity of the patient to maintain privacy

of the data associated with the patient. The data may be used for conglomeration with other data without individualizing the data.

**[0852]** One solution provides a surgical hub **206** to interrogate an electronic medical records database **4002** for patient electronic medical records **4012** data, strip out desirable or relevant patient data **4018** from the patient electronic medical record **4012**, and redact any personal information that could be used to identify the patient. The redaction technique removes any information that could be used to correlate the stripped relevant patient data **4018** to a specific patient, surgery, or time. The surgical hub **206** and the instruments **235** coupled to the surgical hub **206** can then be configured and operated based on the stripped relevant patient data **4018**.

**[0853]** As disclosed in connection with FIG. **62**, extracting (or stripping) relevant patient data **4018** from a patient electronic medical record **4012** while redacting any information that can be used to correlate the patient with the surgery or a scheduled time of the surgery enables the relevant patient data **4018** to be anonymized. The anonymous data file **4016** can then be sent to the cloud **204** for aggregation, processing, and manipulation. The anonymous data file **4016** can be used to configure the surgical instrument **235**, or any of the modules shown in FIGS. **9** and **10** or the surgical hub **206** during the surgery based on the extracted anonymous data file **4016**.

**[0854]** In one aspect, a hospital data barrier **4006** is created such that inside the data barrier **4006** data from various surgical hubs **206** can be compared using non-anonymized un-redacted data and outside the data barrier **4006** data from various surgical hubs **206** are stripped to maintain anonymity and protect the privacy of the patient and the surgeon. This aspect is discussed further in connection with FIG. **66**.

**[0855]** In one aspect, the data from a surgical hub **206** can be exchanged between surgical hubs **206** (e.g., hub-to-hub, switch-to-switch, or router-to-router) to provide in-hospital analysis and display of the data. FIG. **1** shows an example of multiple hubs **106** in communication with each other and with the cloud **104**. This aspect also is discussed further in connection with FIG. **66**.

**[0856]** In another aspect, an artificial time measure is substituted for a real time clock for all information stored internally within an instrument **235**, a robot located in a robot hub **222**, a surgical hub **206**, and/or hospital computer equipment. The anonymized data, which may include anonymized patient and surgeon data, is transmitted to the server **213** in the cloud **204** and it is stored in the cloud storage device **205** coupled to the server **213**. The substitution of an artificial real time clock enables anonymizing the patient data and surgeon data while maintaining data continuity. In one aspect, the instrument **235**, robot hub **222**, surgical hub **206**, and/or the cloud **204** are configured to obscure patient identification (ID) while maintaining data continuity. This aspect is discussed further in connection with FIG. **63**.

**[0857]** Within the surgical hub **206**, a local decipher key **4004** allows information retrieved from the surgical hub **206** itself to reinstate the real-time information from the anonymized data set located in the anonymous data file **4016**. The data stored on the hub **206** or the cloud **204**, however, cannot be reinstated to real-time information from the anonymized data set in the anonymous data file **4016**. The key **4004** is held locally in the surgical hub **206** computer/storage device

**248** in an encrypted format. The surgical hub **206** network processor ID is part of the decryption mechanism such that if the key **4004** and data is removed, the anonymized data set in the anonymous data file **4016** cannot be restored without being on the original surgical hub **206** computer/storage device **248**.

Substituting Artificial Time Measure for Real Time  
Clock for all Internally Stored Information and  
Sent to the Cloud as a Means to Anonymizing the  
Patient and Surgeon Data

[**0858**] FIG. **63** illustrates a process **4030** of anonymizing a surgical procedure by substituting an artificial time measure for a real time clock for all information stored internally within the instrument, robot, surgical hub, and/or hospital computer equipment, according to one aspect of the present disclosure. As shown in FIG. **63**, the surgical procedure set-up start time **4032** was scheduled to begin at an actual time of 11:31:14 (EST) based on a real time clock. At the stated procedure set-up start time **4032**, the surgical hub **206** starts **4034** an artificial randomized real time clock timing scheme at artificial real time at 07:36:00. The surgical hub **206** then ultrasonically pings **4036** the operating theater (e.g., sends out a burst of ultrasound and listens for the echo when it bounces off the perimeter walls of an operating theater (e.g., a fixed, mobile, temporary, or field the operating room) as described in connection with FIG. **64** to verify the size of the operating theater and to adjust short range wireless, e.g., Bluetooth, pairing distance limits at artificial real time 07:36:01. At artificial real time 07:36:03, the surgical hub **206** strips **4038** the relevant data and applies a time stamp to the stripped data. At artificial real time 07:36:05, the surgical hub **206** wakes up and begins pairing **4040** only devices located within the operating theater as verified using the ultrasonic pinging **4036** process.

[**0859**] FIG. **64** illustrates ultrasonic pinging of an operating room wall to determine a distance between a surgical hub and the operating room wall, in accordance with at least one aspect of the present disclosure. With reference also to FIG. **2**, the spatial awareness of the surgical hub **206** and its ability to map an operating room for potential components of the surgical system allows the surgical hub **206** to make autonomous decisions about whether to include or exclude such potential components as part of the surgical system, which relieves the surgical staff from dealing with such tasks. Furthermore, the surgical hub **206** is configured to make inferences about, for example, the type of surgical procedure to be performed in the operating room based on information gathered prior to, during, and/or after the performance of the surgical procedure. Examples of gathered information include the types of devices that are brought into the operating room, time of introduction of such devices into the operating room, and/or the devices sequence of activation.

[**0860**] In one aspect, the surgical hub **206** employs the operating-room mapping module, such as, for example, the non-contact sensor module **242** to determine the bounds of the surgical theater (e.g., a fixed, mobile, or temporary operating room or space) using either ultrasonic or laser non-contact measurement devices.

[**0861**] Referring now to FIG. **64**, ultrasound based non-contact sensors **3002** can be employed to scan the operating theater by transmitting a burst of ultrasound and receiving the echo when it bounces off a perimeter wall **3006** of an

operating theater to determine the size of the operating theater and to adjust short range wireless, e.g., Bluetooth, pairing distance limits. In one example, the non-contact sensors **3002** can be Ping ultrasonic distance sensors, as illustrated in FIG. **64**.

[**0862**] FIG. **64** shows how an ultrasonic sensor **3002** sends a brief chirp with its ultrasonic speaker **3003** and makes it possible for a micro-controller **3004** of the operating-room mapping module to measure how long the echo takes to return to the ultrasonic sensor's ultrasonic microphone **3005**. The micro-controller **3004** has to send the ultrasonic sensor **3002** a pulse to begin the measurement. The ultrasonic sensor **3002** then waits long enough for the micro-controller program to start a pulse input command. Then, at about the same time the ultrasonic sensor **3002** chirps a 40 kHz tone, it sends a high signal to the micro-controller **3004**. When the ultrasonic sensor **3002** detects the echo with its ultrasonic microphone **3005**, it changes that high signal back to low. The micro-controller's pulse input command measures the time between the high and low changes, and stores it measurement in a variable. This value can be used along with the speed of sound in air to calculate the distance between the surgical hub **106** and the operating-room wall **3006**.

[**0863**] In one example, a surgical hub **206** can be equipped with four ultrasonic sensors **3002**, wherein each of the four ultrasonic sensors is configured to assess the distance between the surgical hub **206** and a wall of the operating room **3000**. A surgical hub **206** can be equipped with more or less than four ultrasonic sensors **3002** to determine the bounds of an operating room.

[**0864**] Other distance sensors can be employed by the operating-room mapping module to determine the bounds of an operating room. In one example, the operating-room mapping module can be equipped with one or more photoelectric sensors that can be employed to assess the bounds of an operating room. In one example, suitable laser distance sensors can also be employed to assess the bounds of an operating room. Laser based non-contact sensors may scan the operating theater by transmitting laser light pulses, receiving laser light pulses that bounce off the perimeter walls of the operating theater, and comparing the phase of the transmitted pulse to the received pulse to determine the size of the operating theater and to adjust short range wireless, e.g., Bluetooth, pairing distance limits.

Stripping Out Data from Images and Connected  
Smart Instrument Data to Allow Conglomeration  
but not Individualization

[**0865**] In one aspect, the present disclosure provides a data stripping method which interrogates the electronic patient records provided, extracts the relevant portions to configure and operate the surgical hub and instruments coupled to the surgical hub, while anonymizing the surgery, patient, and all identifying parameters to maintain patient privacy.

[**0866**] With reference now back to FIG. **63** and also to FIGS. **1-11** to show interaction with an interactive surgical system **100** environment including a surgical hub **106**, **206**, once the size of the operating theater has been verified and Bluetooth pairing is complete, based on artificial real time, the computer processor **244** of the surgical hub **206** begins stripping **4038** data received from the modules coupled to the surgical hub **206**. In one example, the processor **244**

begins stripping 4083 images received from the imaging module 238 and connected smart instruments 235, for example. Stripping 4038 the data allows conglomeration of the data but not individualization of the data. This enables stripping 4038 the data identifier, linking the data, and monitoring an event while maintaining patient privacy by anonymizing the data.

[0867] With reference to FIGS. 1-64, in one aspect, a data stripping 4038 method is provided. In accordance with the data stripping 4038 method, the surgical hub 206 processor 244 interrogates the patient records stored in the surgical hub database 238 and extracts the relevant portions of the patient records to configure and operate the surgical hub 206 and its instruments 235, robots, and other modular devices, e.g., modules. The data stripping 4038 method anonymizes the surgical procedure, patient, and all identifying parameters associated with the surgical procedure. Stripping 4038 the data on the fly ensures that at no time the data is correlated to a specific patient, surgical procedure, surgeon, time or other possible identifier that can be used to correlate the data.

[0868] The data may be stripped 4038 for compilation of the base information at a remote cloud 204 database storage device 205 coupled to the remote server 213. The data stored in the database storage device 248 can be used in advanced cloud based analytics, as described in U.S. Provisional Patent Application Ser. No. 62/611,340, filed Dec. 28, 2017, titled CLOUD-BASED MEDICAL ANALYTICS, which is incorporated herein by reference in its entirety. A copy of the information with data links intact also can be stored into the patient EMR database 4002 (FIG. 62). For example, the surgical hub 206 may import patient tissue irregularities or co-morbidities to add to an existing data set stored in the database 248. The data may be stripped 4038 before the surgery and/or may be stripped 4038 as the data is transmitted to the cloud 204 database storage device 205 coupled to the remote server 213.

[0869] With continued reference to FIGS. 1-11 and 62-64, FIG. 65 is a diagram 4050 depicting the process of importing patient electronic medical records 4012 containing surgical procedure and relevant patient data 4018 stored in the EMR database 4002, stripping 4038 the relevant patient data 4018 from the imported medical records 4012, and identifying 4060 smart device implications 4062, or inferences, according to one aspect of the present disclosure. As shown in FIG. 65, the patient electronic medical records 4012, containing information stored in the patient EMR database 4002, are retrieved from the EMR database 4002, imported into the surgical hub 206, and stored in the surgical hub 206 storage device 248. Un-redacted data is removed or deleted 4019 from the patient electronic medical records 4012 before they are stored in the surgical hub 206 storage device 248 as an anonymous data file 4016 (FIG. 62). The relevant patient data 4018 is then stripped 4038 from the medical records 4012 to remove the desired relevant patient data 4018 and delete 4019 un-redacted data to maintain patient anonymity. In the illustrated example, the stripped data 4058 includes emphysema, high blood pressure, small lung cancer, warfarin/blood thinner, and/or radiation pretreatment. The stripped data 4058 is employed to identify 4060 smart device implications while maintaining patient anonymity as described hereinbelow.

[0870] Although the surgical procedure data and relevant patient data 4018 is described as being imported from

patient electronic medical records 4012 stored in the EMR database 4002, in various aspects, the surgical procedure data and relevant patient data 4018 may be retrieved from a modular device coupled to the surgical hub 206 before being stored in the EMR database 4002. For example, the surgical hub 206 may interrogate the module to retrieve the surgical procedure data and relevant patient data 4018 from the module. As described herein, a module includes an imaging module 238 that is coupled to an endoscope 239, a generator module 240 that is coupled to an energy device 241, a smoke evacuator module 226, a suction/irrigation module 228, a communication module 230, a processor module 232, a storage array 234, a smart device/instrument 235 optionally coupled to a display 237, and a non-contact sensor module 242, among other modules as illustrated in FIGS. 3 and 8-10.

[0871] For example, the anonymized stripped data 4058 may be employed to identify 4060 catastrophic failures of instruments, and other smart devices, and may initiate an automatic archive process and submission of data for further implications analysis. For example, the implication of detecting a counterfeit component or adapter on an original equipment manufacturer (OEM) device would be to initiate documentation of the component and recording of the results and outcome of its use. For example, the surgical hub 206 may execute situational awareness algorithms as described in connection FIG. 86. In one aspect, the surgical hub 206 may initially receive or identify a variety of implications 4062 that are derived from anonymized stripped data 4058. The surgical hub 206 is configured to control the instruments 235, or other modules, so that they operate correspondingly to the derived implications 4062. In one example, the surgical hub 206 control logic identifies that (i) lung tissue may be more fragile than normal (e.g., due to emphysema), (ii) hemostasis issues are more likely (e.g., due to high blood pressure and/or the patient being on a blood thinner, such as warfarin), (iii) cancer may be more aggressive (e.g., due to the target of the procedure being a small cell lung cancer), and (iv) lung tissue may be stiffer and more prone to fracture (e.g., due to the patient having received a radiation pretreatment). The control logic or processor 244 of the surgical hub 206 then interprets how this data impacts the instruments 235, or other modules, so that the instruments 235 are operated consistently with the data and then communicates the corresponding adjustments to each of the instruments 235.

[0872] In one example relating to a stapler type of surgical instrument 235, based on the implications 4062 identified 4060 from the anonymized stripped data 4058, the control logic or processor 244 of the surgical hub 206 may (i) notify the stapler to adjust the compression rate threshold parameter, (ii) adjust the surgical hub 206 visualization threshold value to quantify the bleeding and internal parameters, (iii) notify the combo generator module 240 of the lung tissue and vessel tissue types so that the power and generator module 240 control algorithms are adjusted accordingly, (iv) notify the imaging module 238 of the aggressive cancer tag to adjust the margin ranges accordingly, (v) notify the stapler of the margin parameter adjustment needed (the margin parameter corresponds to the distance or amount of tissue around the cancer that will be excised), and (vi) notify the stapler that the tissue is potentially fragile. Furthermore, the anonymized stripped data 4058, upon which the implications 4060 are based, is identified by the surgical hub 206 and is fed into the situational awareness algorithm (see FIG.

86). Examples include, without limitations, thoracic lung resection, e.g., segmentectomy, among others.

[0873] FIG. 66 is a diagram 4070 illustrating the application of cloud based analytics to un-redacted data, stripped relevant patient data 4018, and independent data pairs, according to one aspect of the present disclosure. As shown, multiple surgical hubs Hub #1 4072, Hub #3 4074, and Hub #4 4076 are located within the hospital data barrier 4006 (see also FIG. 62). The un-redacted patient electronic medical record 4012 including patient data and surgery related data may be used and exchanged between the surgical hubs: Hub #1 4072, Hub #3 4074, and Hub #4 4076 located within the hospital data barrier 4006. Prior to transmitting the un-redacted patient electronic medical record 4012 containing patient data and surgery related data outside the hospital data barrier 4006, however, the patient electronic medical record 4012 patient data is redacted and stripped to create an anonymous data file 4016 containing anonymized information for further analysis and processing of the redacted/stripped data by a cloud based analytic processes in the cloud 204.

[0874] FIG. 67 is a logic flow diagram 4080 of a process depicting a control program or a logic configuration for associating patient data sets from first and second sources of data, according to one aspect of the present disclosure. With reference to FIG. 67 and with reference also to FIGS. 1-11 to show interaction with an interactive surgical system 100 environment including a surgical hub 106, 206, in one aspect, the present disclosure provides a surgical hub 206, comprising a processor 244; and a memory 249 coupled to the processor 244. The memory 249 stores instructions executable by the processor 244 to interrogate 4082 a surgical instrument 235, retrieve 4084 a first data set from the surgical instrument 235, interrogate 4086 a medical imaging device 238, retrieve 4088 a second data set from the medical imaging device 238, associate 4090 the first and second data sets by a key, and transmit the associated first and second data sets to a remote network outside of the surgical hub 206. The surgical instrument 235 is a first source of patient data and the first data set is associated with a surgical procedure. The medical imaging device 238 is a second source of patient data and the second data set is associated with an outcome of the surgical procedure. The first and second data records are uniquely identified by the key.

[0875] In another aspect, the surgical hub 206 provides a memory 249 storing instructions executable by the processor 244 to retrieve the first data set using the key, anonymize the first data set, retrieve the second data set using the key, anonymize the second data set, pair the anonymized first and second data sets, and determine success rate of surgical procedures grouped by the surgical procedure based on the anonymized paired first and second data sets.

[0876] In another aspect, the surgical hub 206 provides a memory 249 storing instructions executable by the processor 244 to retrieve the anonymized first data set, retrieve the anonymized second data set, and reintegrate the anonymized first and second data sets using the key.

[0877] FIG. 68 is a logic flow diagram of a process 4400 depicting a control program or a logic configuration for stripping data to extract relevant portions of the data to configure and operate the surgical hub 206 and modules (e.g., instruments 235) coupled to the surgical hub 206, according to one aspect of the present disclosure. With

reference to FIG. 68 and with reference also to FIGS. 1-11 to show interaction with an interactive surgical system 100 environment including a surgical hub 106, 206, in one aspect, the surgical hub 206 may be configured to interrogate a module coupled to surgical hub 206 for data, and strip the data to extract relevant portions of the data to configure and operate the surgical hub 206 and modules (e.g., instruments 235) coupled to the surgical hub 206 and anonymize the surgery, patient, and other parameters that can be used to identify the patient to maintain patient privacy. According to the process 4400, in one aspect the present disclosure provides a surgical hub 206 including a processor 244, a modular communication hub 203 coupled to the processor 244, where the modular communication hub 203 is configured to connect modular devices located in one or more operating theaters to the surgical hub 206. The processor 244 is coupled to a memory 249, where the memory 249 stores instructions executable by the processor 244 to cause the processor to interrogate 4402 a modular device coupled to the processor 244 via the modular communication hub 203. The modular device is a source of data sets that include patient identity data and surgical procedure data. The processor 244 receives 4404 a data set from the modular device. The processor 244 discards 4406 the patient identity data and any portion of the surgical procedure data that identifies the patient from the data set. The processor 244 extracts 4408 anonymous data from the data set and creates 4410 an anonymized data set. The processor 244 configures 4412 the operation of the surgical hub 206 or the modular device based on the anonymized data set.

[0878] In another aspect, where the anonymized data set includes catastrophic failure of a modular device, the memory 249 stores instructions executable by the processor 244 to initiate automatic archiving and submission of data for implications analysis based on the catastrophic failure of the modular device. In another aspect, the memory 249 stores instructions executable by the processor 244 to detect counterfeit component information from the anonymized data set. In another aspect, the memory 249 stores instructions executable by the processor 244 to derive implications of the modular device from the anonymized data set and the memory 249 stores instructions executable by the processor 244 to configure the modular device to operate based on the derived implications or to configure the surgical hub based on the derived implications. In another aspect, the memory 249 stores instructions executable by the processor 244 to conglomerate the anonymized data. In another aspect, the memory 249 stores instructions executable by the processor 244 to extract the anonymized data prior to storing the received data in a storage device coupled to the surgical hub. In another aspect, the memory 249 stores instructions executable by the processor to transmit the anonymized data to a remote network outside of the surgical hub, compile the anonymized data at the remote network, and store a copy of the data set from the modular device in a patient electronic medical records database.

#### Storage of Data Creation and Use of Self-Describing Data Including Identification Features

[0879] In one aspect, the present disclosure provides self-describing data packets generated at the issuing instrument and including identifiers for all devices that handled the packet. The self description allows the processor to interpret

the data in the self-describing packet without knowing the data type in advance prior to receipt of the self-describing packet. The data applies to every data point or data string and includes the type of data, the source of the self-describing packet, the device identification that generated the packet, the units, the time of generation of the packet, and an authentication that the data contained in the packet is unaltered. When the processor (in the device or the surgical hub) receives an unexpected packet and verifies the source of the packet, the processor alters the collection techniques to be ready for any subsequent packets from that source.

[0880] With reference also to FIGS. 1-11 to show interaction with an interactive surgical system 100 environment including a surgical hub 106, 206, during a surgical procedure being performed in a surgical hub 206 environment, the size and quantity of data being generated by surgical devices 235 coupled to the surgical hub 206 can become quite large. Also, data exchanged between the surgical devices 235 and/or the surgical hub 206 can become quite large.

[0881] One solution provides a techniques for minimizing the size of the data and handling the data within a surgical hub 206 by generating a self-describing packet. The self-describing packet is initially assembled by the instrument 235 that generated it. The packet is then ordered and encrypted by generating an encryption certificate which is unique for each data packet. The data is then communicated from the instrument 235 via encrypted wired or wireless protocols and stored on the surgical hub 206 for processing and transmission to the cloud 204 analytics engine. Each self-describing data packet includes an identifier to identify the specific instrument that generated it and the time it was generated. A surgical hub 206 identifier is added to the packet when the packet is received by the surgical hub 206.

[0882] In one aspect, the present disclosure provides a surgical hub 206 comprising a processor 244 and a memory 249 coupled to the processor 244. The memory 249 storing instructions executable by the processor 244 to receive a first data packet from a first source, receive a second data packet from a second source, associate the first and second data packets, and generate a third data packet comprising the first and second data payloads. The first data packet comprises a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate. The first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet. The second data packet comprises a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate. The second preamble defines the second data payload and the second encryption certificate verifies the authenticity of the second data packet.

[0883] In another aspect, the memory 249 stores instructions executable by the processor 244 to determine that a data payload is from a new source, verify the new source of the data payload, and alter a data collection process at the surgical hub to receive subsequent data packets from the new source.

[0884] In another aspect, the memory 249 stores instructions executable by the processor 244 to associate the first and second data packets based on a key. In another aspect, the memory 249 stores instructions executable by the processor 244 to anonymize the data payload of the third data packet. In another aspect, the memory 249 stores instructions executable by the processor 244 to receive an anonymized

third data packet and reintegrate the anonymized third data packet into the first and second data packets using the key.

[0885] In various aspects, the present disclosure provides a control circuit to receive and process data packets as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer readable instructions, which when executed, causes a machine to receive and process data packets as described above.

[0886] In other aspects, the present disclosure a method of generating a data packet comprising self-describing data. In one aspect, a surgical instrument includes a processor and a memory coupled to the processor, a control circuit, and/or a computer-readable medium configured to generate a data packet comprising a preamble, a data payload, a source of the data payload, and an encryption certificate. The preamble defines the data payload and the encryption certificate verifies the authenticity of the data packet. In various aspects, the data packet may be generated by any module coupled to the surgical hub. Self-describing data packets minimize data size and data handling in the surgical hub.

[0887] In one aspect, the present disclosure provides a self-describing data packet generated at an issuing device (e.g., instrument, tool, robot). The self-describing data packet comprises identifiers for all devices that handle the data packet along a communication path; a self description to enable a processor to interpret that data contained in the data packet without having been told in advance of receipt of the data packet along a path; data for every data point or data string; and type of data, source of data, device IDs that generated the data, units of the data, time of generation, and authentication that the data packet is unaltered. In another aspect, when a processor receives a data packet from an unexpected source and verifies the source of the data, the processor alters the data collection technique to prepare for any subsequent data packets from the source.

[0888] In the creation and use of a data packet comprising self-describing data, the surgical hub includes identification features. The hub and intelligent devices use self-describing data packets to minimize data size and data handling. In a surgical hub that generates large volumes of data, the self-describing data packets minimize data size and data handling, thus saving time and enabling the operating theater to run more efficiently.

[0889] FIG. 69 illustrates a self-describing data packet 4100 comprising self-describing data, according to one aspect of the present disclosure. With reference also to FIGS. 1-11 to show interaction with an interactive surgical system 100 environment including a surgical hub 106, 206, in one aspect, self-describing data packets 4100 as shown in FIG. 69 are generated at an issuing instrument 235, or device or module located in or in communication with the operating theater, and include identifiers for all devices 235 that handle the packet along a communication path. The self description allows a processor 244 to interpret the data payload of the packet 4100 without having advance knowledge of the definition of the data payload prior to receiving the self-describing data packet 4100. The processor 244 can interpret the data payload by parsing an incoming self-describing packet 4100 as it is received and identifying the data payload without being notified in advance that the self-describing packet 4100 was received. The data is for every data point or data string. The data payload includes type of data, source

of data, device IDs that generated the data, data units, time when data was generated, and an authentication that the self-describing data packet 4100 is unaltered. Once the processor 244, which may be located either in the device or the surgical hub 206, receives an unexpected self-describing data packet 4100 and verifies the source of the self-describing data packet 4100, the processor 244 alters the data collection means to be ready for any subsequent self-describing data packets 4100 from that source. In one example, the information contained in a self-describing packet 4100 may be recorded during the first firing 4172 in the lung tumor resection surgical procedure described in connection with FIGS. 71-75.

[0890] The self-describing data packet 4100 includes not only the data but a preamble which defines what the data is and where the data came from as well as an encryption certificate verifying the authenticity of each data packet 4100. As shown in FIG. 69, the data packet 4100 may comprise a self-describing data header 4102 (e.g., force-to-fire [FTF], force-to-close [FTC], energy amplitude, energy frequency, energy pulse width, speed of firing, and the like), a device ID 4104 (e.g., 002), a shaft ID 4106 (e.g., W30), a cartridge ID 4108 (e.g., ESN736), a unique time stamp 4110 (e.g., 09:35:15), a force-to-fire value 4112 (e.g., 85) when the self-describing data header 4102 includes FTF (force-to-fire), otherwise, this position in the data packet 4100 includes the value of force-to-close, energy amplitude, energy frequency, energy pulse width, speed of firing, and the like. The data packet 4100, further includes tissue thickness value 4114 (e.g., 1.1 mm), and an identification certificate of data value 4116 (e.g., 01101010001001) that is unique for each data packet 4100. Once the self-describing data packet 4100 is received by another instrument 235, surgical hub 206, cloud 204, etc., the receiver parses the self-describing data header 4102 and based on its value knows what data type is contained in the self-describing data packet 4100. TABLE 1 below lists the value of the self-describing data header 4102 and the corresponding data value.

TABLE 1

Self-Describing Data Header (4102)	Data Type
FTF	Force To Fire (N)
FTC	Force To Close (N)
EA	Energy Amplitude (J)
EF	Energy Frequency (Hz)
EPW	Energy Pulse Width (Sec)
SOF	Speed Of Firing (mm/sec)

[0891] Each self-describing data packet 4100 comprising self-describing data is initially assembled by the instrument 235, device, or module that generated the self-describing data packet 4100. Subsequently, the self-describing data packet 4100 comprising self-describing data is ordered and encrypted to generate an encryption certificate. The encryption certificate is unique for each self-describing data packet 4100. That data is then communicated via encrypted wired or wireless protocols and stored on the surgical hub 206 for processing and transmission to the cloud 204 analytics engine.

[0892] Each self-describing data packet 4100 comprising self-describing data includes a device ID 4104 to identify the specific instrument 235 that generated the self-describing

data packet 4100, a time stamp 4110 to indicate the time that the data packet 4100 was generated, and when the self-describing data packet 4100 is received by the surgical hub 206. The surgical hub 206 ID also may be added to the self-describing data packet 4100.

[0893] Each of the self-describing data packets 4100 comprising self-describing data may include a packet wrapper that defines the beginning of the data packet 4100 and the end of the data packet 4100 including any identifiers necessary to forecast the number and order of the bits in the self-describing data packet.

[0894] The surgical hub 206 also manages redundant data sets. As the device 235 functions and interconnects with other surgical hubs 206, multiple sets of the same data may be created and stored on various devices 235. Accordingly, the surgical hub 206 manages multiple images of redundant data as well as anonymization and security of data. The surgical hub 206 also provides temporary visualization and communication, incident management, peer-to-peer processing or distributed processing, and storage backup and protection of data.

[0895] FIG. 70 is a logic flow diagram 4120 of a process depicting a control program or a logic configuration for using data packets comprising self-describing data, according to one aspect of the present disclosure. With reference to FIGS. 1-69, in one aspect, the present disclosure provides a surgical hub 206 comprising a processor 244 and a memory 249 coupled to the processor 244. The memory 249 storing instructions executable by the processor 244 to receive a first data packet from a first source, receive a second data packet from a second source, associate the first and second data packets, and generate a third data packet comprising the first and second data payloads. The first data packet comprises a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate. The first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet. The second data packet comprises a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate. The second preamble defines the second data payload and the second encryption certificate verifies the authenticity of the second data packet.

[0896] In another aspect, the memory 249 stores instructions executable by the processor 244 to determine that a data payload is from a new source, verify the new source of the data payload, and alter a data collection process at the surgical hub to receive subsequent data packets from the new source.

[0897] In another aspect, the memory 249 stores instructions executable by the processor 244 to associate the first and second data packets based on a key. In another aspect, the memory 249 stores instructions executable by the processor 244 to anonymize the data payload of the third data packet. In another aspect, the memory 244 stores instructions executable by the processor 244 to receive an anonymized third data packet and reintegrate the anonymized third data packet into the first and second data packets using the key.

[0898] FIG. 71 is a logic flow diagram 4130 of a process depicting a control program or a logic configuration for using data packets comprising self-describing data, according to one aspect of the present disclosure. With reference to FIG. 71 and with reference also to FIGS. 1-11 to show interaction with an interactive surgical system 100 environ-

ment including a surgical hub **106, 206**, in one aspect, the present disclosure provides a surgical hub **206** comprising a processor **244** and a memory **249** coupled to the processor **244**. The memory **249** storing instructions executable by the processor **244** to receive **4132** a first self-describing data packet from a first data source, the first self-describing data packet comprising a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate. The first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet. The memory **249** storing instructions executable by the processor **244** to parse **4134** the received first preamble and interpret **4136** the first data payload based on the first preamble.

**[0899]** In various aspects, the memory **249** stores instructions executable by the processor **244** to receive a second self-describing data packet from a second data source, the second self-describing data packet comprising a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate. The second preamble defines the second data payload and the second encryption certificate verifies the authenticity of the second data packet. The memory **249** storing instructions executable by the processor **244** to parse the received second preamble, interpret the second data payload based on the second preamble, associate the first and second self-describing data packets, and generate a third self-describing data packet comprising the first and second data payloads. In one aspect, the memory stores instructions executable by the processor to anonymize the data payload of the third self-describing data packet.

**[0900]** In various aspects, the memory stores instructions executable by the processor to determine that a data payload was generated by a new data source, verify the new data source of the data payload, and alter a data collection process at the surgical hub to receive subsequent data packets from the new data source. In one aspect, the memory stores instructions executable by the processor to associate the first and second self-describing data packets based on a key. In another aspect, the memory stores instructions executable by the processor to receive an anonymized third self-describing data packet and reintegrate the anonymized third self-describing data packet into the first and second self-describing data packets using the key.

Storage of the Data in a Manner of Paired Data Sets which can be Grouped by Surgery but not Necessarily Keyed to Actual Surgical Dates and Surgeons

**[0901]** In one aspect, the present disclosure provides a data pairing method that allows a surgical hub to interconnect a device measured parameter with a surgical outcome. The data pair includes all the relevant surgical data or patient qualifiers without any patient identifier data. The data pair is generated at two separate and distinct times. The disclosure further provides configuring and storing the data in such a manner as to be able to rebuild a chronological series of events or merely a series of coupled but unconstrained data sets. The disclosure further provides storing data in an encrypted form and having predefined backup and mirroring to the cloud.

**[0902]** To determine the success or failure of a surgical procedure, data stored in a surgical instrument should be correlated with the outcome of the surgical procedure while

simultaneously anonymizing the data to protect the privacy of the patient. One solution is to pair data associated with a surgical procedure, as recorded by the surgical instrument during the surgical procedure, with data assessing the efficacy of the procedure. The data is paired without identifiers associated with surgery, patient, or time to preserve anonymity. The paired data is generated at two separate and distinct times.

**[0903]** In one aspect, the present disclosure provides a surgical hub configured to communicate with a surgical instrument. The surgical hub comprises a processor and a memory coupled to the processor. The memory storing instructions executable by the processor to receive a first data set associated with a surgical procedure, receive a second data set associated with the efficacy of the surgical procedure, anonymize the first and second data sets by removing information that identifies a patient, a surgery, or a scheduled time of the surgery, and store the first and second anonymized data sets to generate a data pair grouped by surgery. The first data set is generated at a first time, the second data set is generated at a second time, and the second time is separate and distinct from the first time.

**[0904]** In another aspect, the memory stores instructions executable by the processor to reconstruct a series of chronological events based on the data pair. In another aspect, the memory stores instructions executable by the processor to reconstruct a series of coupled but unconstrained data sets based on the data pair. In another aspect, the memory stores instructions executable by the processor to encrypt the data pair, define a backup format for the data pair, and mirror the data pair to a cloud storage device.

**[0905]** In various aspects, the present disclosure provides a control circuit to receive and process data sets as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer readable instructions, which when executed, causes a machine to receive and process data sets as described above.

**[0906]** Storage of paired anonymous data enables the hospital or surgeon to use the data pairs locally to link to specific surgeries or to store the data pairs to analyze overall trends without extracting specific events in chronological manner.

**[0907]** In one aspect, the surgical hub provides user defined storage and configuration of data. Storage of the data may be made in a manner of paired data sets which can be grouped by surgery, but not necessarily keyed to actual surgical dates and surgeons. This technique provides data anonymity with regard to the patient and surgeon.

**[0908]** In one aspect, the present disclosure provides a data pairing method. The data pairing method comprises enabling a surgical hub to interconnect a device measured parameter with an outcome, wherein a data pair includes all the relevant tissue or patient qualifiers without any of the identifiers, wherein the data pair is generated at two distinct and separate times. In another aspect, the present disclosure provides a data configuration that includes whether the data is stored in such a manner as to enable rebuilding a chronological series of events or merely a series of coupled but unconstrained data sets. In another aspect, the data may be stored in an encrypted form. The stored data may comprise a predefined backup and mirroring to the cloud.

**[0909]** The data may be encrypted locally to the device. The data backup may be automatic to an integrated load secondary storage device. The device and/or the surgical hub



may be configured to maintain the time of storage of the data and compile and transmit the data to another location for storage, e.g., another surgical hub or a cloud storage device. The data may be grouped together and keyed for transmission to the cloud analytics location. A cloud based analytics system is described in commonly-owned U.S. Provisional Patent Application Ser. No. 62/611,340, filed Dec. 28, 2017, titled CLOUD-BASED MEDICAL ANALYTICS, which is incorporated herein by reference in its entirety.

[0910] In another aspect, the hub provides user selectable options for storing the data. In one technique, the hub enables the hospital or the surgeon to select if the data should be stored in such a manner that it could be used locally in a surgical hub to link to specific surgeries. In another technique, the surgical hub enables the data to be stored as data pairs so that overall trends can be analyzed without specific events extracted in a chronological manner.

[0911] FIG. 72 is a diagram 4150 of a tumor 4152 embedded in the right superior posterior lobe 4154 of the right lung 4156, according to one aspect of the present disclosure. To remove the tumor 4152, the surgeon cuts around the tumor 4152 along the perimeter generally designated as a margin 4158. A fissure 4160 separates the upper lobe 4162 and the middle lobe 4164 of the right lung 4156. In order to cut out the tumor 4152 about the margin 4158, the surgeon must cut the bronchial vessels 4166 leading to and from the middle lobe 4164 and the upper lobe 4162 of the right lung 4156. The bronchial vessels 4166 must be sealed and cut using a device such as a surgical stapler, electro-surgical instrument, ultrasonic instrument, a combo electro-surgical/ultrasonic instrument, and/or a combo stapler/electro-surgical device generally represented herein as the instrument/device 235 coupled to the surgical hub 206. The device 235 is configured to record data as described above, which is formed as a data packet, encrypted, stored, and/or transmitted to a remote data storage device 105 and processed by the server 113 in the cloud 104. FIGS. 77 and 78 are diagrams that illustrate the right lung 4156 and the bronchial tree 4250 embedded within the parenchyma tissue of the lung.

[0912] In one aspect, the data packet may be in the form of the self-describing data 4100 described in connection with FIGS. 69-71. The self-describing data packet 4100 will contain the information recorded by the device 235 during the procedure. Such information may include, for example, a self-describing data header 4102 (e.g., force-to-fire [FTF], force-to-close [FTC], energy amplitude, energy frequency, energy pulse width, speed of firing, and the like) based on the particular variable. The device ID 4104 (e.g., 002) of the instrument/device 235 used in the procedure including components of the instrument/device 235 such as the shaft ID 4106 (e.g., W30) and the cartridge ID 4108 (e.g., ESN736). The self-describing packet 4100 also records a unique time stamp 4110 (e.g., 09:35:15) and procedural variables such as a force-to-fire value 4112 (e.g., 85) when the self-describing data header 4102 includes FTF (force-to-fire), otherwise, this position in the data packet 4100 includes the value of force-to-close (FTC), energy amplitude, energy frequency, energy pulse width, speed of firing, and the like, as shown in TABLE 1, for example. The data packet 4100, further may include tissue thickness value 4114 (e.g., 1.1 mm), which in this example refers to the thickness of the bronchial vessel 4166 exposed in the fissure 4160 that were sealed and cut. Finally, each self-describing packet 4100 includes an identification certificate of data value 4116 (e.g.,

01101010001001) that uniquely identifies each data packet 4100 transmitted by the device/instrument 235 to the surgical hub 206, further transmitted from the surgical hub 206 to the cloud 204 and stored on the storage device 205 coupled to the server 213, and/or further transmitted to the robot hub 222 and stored.

[0913] The data transmitted by way of a self-describing data packet 4100 is sampled by the instrument device 235 at a predetermined sample rate. Each sample is formed into a self-describing data packet 4100 which is transmitted to the surgical hub 206 and eventually is transmitted from the surgical hub 206 to the cloud 204. The samples may be stored locally in the instrument device 235 prior to packetizing or may be transmitted on the fly. The predetermined sampling rate and transmission rate are dictated by communication traffic in the surgical hub 206 and may be adjusted dynamically to accommodate current bandwidth limitations. Accordingly, in one aspect, the instrument device 235 may record all the samples taken during surgery and at the end of the procedure packetize each sample into a self-describing packet 4100 and transmit the self-describing packet 4100 to the surgical hub 206. In another aspect, the sampled data may be packetized as it is recorded and transmitted to the surgical hub 206 on the fly.

[0914] FIG. 73 is a diagram 4170 of a lung tumor resection surgical procedure including four separate firings of a surgical stapler device 235 to seal and cut bronchial vessels 4166 exposed in the fissure 4160 leading to and from the upper and lower lobes 4162, 4164 of the right lung 4156 shown in FIG. 72, according to one aspect of the present disclosure. The surgical stapler device 235 is identified by a Device ID "002". The data from each firing of the surgical stapler device 235 is recorded and formed into a data packet 4100 comprising self-describing data as shown in FIG. 70. The self-describing data packet 4100 shown in FIG. 70 is representative of the first firing of device "002" having a staple cartridge serial number of ESN736, for example. In the following description, reference also is made to FIGS. 12-19 for descriptions of various architectures of instruments/devices 235 that include a processor or a control circuit coupled to a memory for recording (e.g., saving or storing) data collected during a surgical procedure.

[0915] The first firing 4172 is recorded at anonymous time 09:35:15. The first firing 4172 seals and severs a first bronchial vessel 4166 leading to and from the middle lobe 4164 and the upper lobe 4162 of the right lung 4156 into a first portion 4166a and a second portion 4166b, where each portion 4166a, 4166b is sealed by respective first and second staple lines 4180a, 4180b. Information associated with the first firing 4172, for example the information described in connection with FIG. 70, is recorded in the surgical stapler device 235 memory and is used to build a first self-describing data packet 4100 described in connection with FIGS. 69-71. The first self-describing packet 4100 may be transmitted upon completion of the first firing 4172 or may be kept stored in the surgical stapler device 235 memory until the surgical procedure is completed. Once transmitted by the surgical stapler device 235, the first self-describing data packet 4100 is received by the surgical hub 206. The first self-describing data packet 4100 is anonymized by stripping and time stamping 4038 the data, as discussed, for example, in connection with FIG. 63. After the lung resection surgical is completed, the integrity of the seals of the first and second staple lines 4182a, 4182b will be evaluated as shown in FIG.

74, for example, and the results of the evaluation will be paired with information associated with the first firing 4172.

[0916] The second firing 4174 seals and severs a second bronchial vessel of the bronchial vessels 4166 leading to and from the middle lobe 4164 and the upper lobe 4162 of the right lung 4156 into a first portion 4166c and a second portion 4166d, where each portion 4166c, 4166d is sealed by first and second staple lines 4180c, 4180d. Information associated with the second firing 4174, for example the information described in connection with FIGS. 69-71, is recorded in the surgical stapler device 235 memory and is used to build a second self-describing data packet 4100 described in connection with FIGS. 69-71. The second self-describing data packet 4100 may be transmitted upon completion of the second firing 4174 or may be kept stored in the surgical stapler device 235 memory until the surgical procedure is completed. Once transmitted by the surgical stapler device 235, the second self-describing data packet 4100 is received by the surgical hub 206. The second self-describing data packet 4100 is anonymized by stripping and time stamping 4038 the data as discussed, for example, in connection with FIG. 63. After the lung resection surgical is completed, the integrity of the seals of the first and second staple lines 4182c, 4182d will be evaluated as shown in FIG. 74, for example, and the results of the evaluation will be paired with information associated with the second firing 4174.

[0917] The third firing 4176 is recorded at anonymous time 09:42:12. The third firing 4176 seals and severs an outer portion of the upper and middle lobes 4162, 4164 of the right lung 4156. First and second staple lines 4182a, 4182b are used to seal the outer portion of the upper and middle lobes 4162, 4162. Information associated with the third firing 4176, for example the information described in connection with FIGS. 69-71, is recorded in the surgical stapler device 235 memory and is used to build a third self-describing data packet 4100 described in connection with FIGS. 69-71. The third self-describing packet 4100 may be transmitted upon completion of the third firing 4176 or may be kept stored in the surgical stapler device 235 memory until the surgical procedure is completed. Once transmitted by the surgical stapler device 235, the third self-describing data packet 4100 is received by the surgical hub 206. The third self-describing data packet 4100 is anonymized by stripping and time stamping 4038 the data, as discussed, for example, in connection with FIG. 63. After the lung resection surgical is completed, the integrity of the seals of the first and second staple lines 4180a, 4180b will be evaluated as shown in FIG. 74, for example, and the results of the evaluation will be paired with information associated with the third firing 4172.

[0918] The fourth firing 4178 seals and severs an inner portion of the upper and middle lobes 4162, 4162 of the right lung 4156. First and second staple lines 4182c, 4182d are used to seal the inner portions of the upper and middle lobes 4162, 4164. Information associated with the fourth firing 4178, for example the information described in connection with FIG. 70, is recorded in the surgical stapler device 235 memory and is used to build a fourth self-describing data packet 4100 described in connection with FIGS. 69-71. The fourth self-describing packet 4100 may be transmitted upon completion of the fourth firing 4178 or may be kept stored in the surgical stapler device 235 memory until the surgical procedure is completed. Once transmitted by the surgical

stapler device 235, the fourth self-describing data packet 4100 is received by the surgical hub 206. The fourth self-describing data packet 4100 is anonymized by stripping and time stamping 4038 the data, as discussed, for example, in connection with FIG. 63. After the lung resection surgical is completed, the integrity of the seals of the first and second staple lines 4182a, 4182b will be evaluated as shown in FIG. 74, for example, and the results of the evaluation will be paired with information associated with the fourth firing 4172.

[0919] FIG. 74 is a graphical illustration 4190 of a force-to-close (FTC) versus time curve 4192 and a force-to-fire (FTF) versus time curve 4194 characterizing the first firing 4172 of device 002 shown in FIG. 73, according to one aspect of the present disclosure. The surgical stapler device 235 is identified as 002 with a 30 mm staple cartridge S/N ESN736 with a PVS shaft S/N M3615N (Shaft ID W30). The surgical stapler device 235 was used for the first firing 4172 to complete the lung resection surgical procedure shown in FIG. 73. As shown in FIG. 74, the peak force-to-fire force of 85 N. is recorded at anonymous time 09:35:15. Algorithms in the surgical stapler device 235 determine a tissue thickness of about 1.1 mm. As described hereinbelow, the FTC versus time curve 4192 and the FTF versus time curve 4194 characterizing the first firing 4172 of the surgical device 235 identified by ID 002 will be paired with the outcome of the lung resection surgical procedure, transmitted to the surgical hub 206, anonymized, and either stored in the surgical hub 206 or transmitted to the cloud 204 for aggregation, further processing, analysis, etc.

[0920] FIG. 75 is a diagram 4200 illustrating a staple line visualization laser Doppler to evaluate the integrity of staple line seals by monitoring bleeding of a vessel after a firing of a surgical stapler, according to one aspect of the present disclosure. A laser Doppler technique is described in above under the heading "Advanced Imaging Acquisition Module," in U.S. Provisional Patent Application Ser. No. 62/611, 341, filed Dec. 28, 2017, and titled INTERACTIVE SURGICAL PLATFORM, which is hereby incorporated by reference herein in its entirety. The laser Doppler provides an image 4202 suitable for inspecting seals along the staple lines 4180a, 4180b, 4182a and for visualizing bleeding 4206 of any defective seals. Laser Doppler inspection of the first firing 4172 of device 002 shows a defective seal at the first staple line 4180a of the first portion 4166a of the bronchial vessel sealed during the first firing 4172. The staple line 4180a seal is bleeding 4206 out at a volume of 0.5 cc. The image 4202 is recorded at anonymous time 09:55:15 4204 and is paired with the force-to-close curve 4192 and force-to-fire curve 4194 shown in FIG. 74. The data pair set is grouped by surgery and is stored locally in the surgical hub 206 storage 248 and/or remotely to the cloud 204 storage 205 for aggregation, processing, and analysis, for example. For example, the cloud 204 analytics engine associates the information contained in the first self-describing packet 4100 associated with the first firing 4172 and indicate that a defective seal was produced at the staple line 4166a. Over time, this information can be aggregated, analyzed, and used to improve outcomes of the surgical procedure, such as, resection of a lung tumor, for example.

[0921] FIG. 76 illustrates two paired data sets 4210 grouped by surgery, according to one aspect of the present disclosure. The upper paired data set 4212 is grouped by one surgery and a lower paired data set 4214 grouped by another

surgery. The upper paired data set **4212**, for example, is grouped by the lung tumor resection surgery discussed in connection with FIGS. **73-76**. Accordingly, the rest of the description of FIG. **76** will reference information described in FIGS. **32-35** as well as FIGS. **1-21** to show interaction with an interactive surgical system **100** environment including a surgical hub **106, 206**. The lower paired data set **4214** is grouped by a liver tumor resection surgical procedure where the surgeon treated parenchyma tissue. The upper paired data set is associated with a failed staple line seal and the bottom paired data set is associated with a successful staple line seal. The upper and lower paired data sets **4212, 4214** are sampled by the instrument device **235** and each sample formed into a self-describing data packet **4100** which is transmitted to the surgical hub **206** and eventually is transmitted from the surgical hub **206** to the cloud **204**. The samples may be stored locally in the instrument device **235** prior to packetizing or may be transmitted on the fly. Sampling rate and transmission rate are dictated by communication traffic in the surgical hub **206** and may be adjusted dynamically to accommodate current bandwidth limitations.

[0922] The upper paired data set **4212** includes a left data set **4216** recorded by the instrument/device **235** during the first firing **4172** linked **4224** to a right data set **4218** recorded at the time the staple line seal **4180a** of the first bronchial vessel **4166a** was evaluated. The left data set **4216** indicates a “Vessel” tissue type **4236** having a thickness **4238** of 1.1 mm. Also included in the left data set **4216** is the force-to-close curve **4192** and force-to-fire curve **4194** versus time (anonymous real time) recorded during the first firing **4172** of the lung tumor resection surgical procedure. The left data set **4216** shows that the force-to-fire peaked at 85 Lbs. and recorded at anonymous real time **4240**  $t_{1a}$  (09:35:15). The right data set **4218** depicts the staple line visualization curve **4228** depicting leakage versus time. The right data set **4218** indicates that a “Vessel” tissue type **4244** having a thickness **4246** of 1.1 mm experienced a staple line **4180a** seal failure **4242**. The staple line visualization curve **4228** depicts leakage volume (cc) versus time of the staple line **4180a** seal. The staple line visualization curve **4228** shows that the leakage volume reached 0.5 cc, indicating a failed staple line **4180a** seal of the bronchial vessel **4166a**, recorded at anonymous time **4248** (09:55:15).

[0923] The lower paired data set **4214** includes a left data set **4220** recorded by the instrument/device **235** during a firing linked **4226** to a right data set **4222** recorded at the time the staple line seal of the parenchyma tissue was evaluated. The left data set **4220** indicates a “Parenchyma” tissue type **4236** having a thickness **4238** of 2.1 mm. Also included in the left data set **4220** is the force-to-close curve **4230** and force-to-fire curve **4232** versus time (anonymous real time) recorded during the first firing of the liver tumor resection surgical procedure. The left data set **4220** shows that the force-to-fire peaked at 100 Lbs. and recorded at anonymous real time **4240**  $t_{1b}$  (09:42:12). The right data set **4222** depicts the staple line visualization curve **4228** depicting leakage versus time. The right data set **4234** indicates that a “Parenchyma” tissue type **4244** having a thickness **4246** of 2.2 mm experienced a successful staple line seal. The staple line visualization curve **4234** depicts leakage volume (cc) versus time of the staple line seal. The staple line visualization curve **4234** shows that the leakage volume

was 0.0 cc, indicating a successful staple line seal of the parenchyma tissue, recorded at anonymous time **4248** (10:02:12).

[0924] The paired date sets **4212, 4214** grouped by surgery are collected for many procedures and the data contained in the paired date sets **4212, 4214** is recorded and stored in the cloud **204** storage **205** anonymously to protect patient privacy, as described in connection with FIGS. **62-69**. In one aspect, the paired date sets **4212, 4214** data are transmitted from the instrument/device **235**, or other modules coupled to the surgical hub **206**, to the surgical hub **206** and to the cloud **204** in the form of the self-describing packet **4100** as described in connection with FIGS. **71** and **72** and surgical procedure examples described in connection with FIGS. **72-76**. The paired date sets **4212, 4214** data stored in the cloud **204** storage **205** is analyzed in the cloud **204** to provide feedback to the instrument/device **235**, or other modules coupled to the surgical hub **206**, notifying a surgical robot coupled to the robot hub **222**, or the surgeon, that the conditions identified by the left data set ultimately lead to either a successful or failed seal. As described in connection with FIG. **76**, the upper left data set **4216** led to a failed seal and the bottom left data set **4220** led to a successful seal. This is advantageous because the information provided in a paired data set grouped by surgery can be used to improve resection, transection, and creation of anastomosis in a variety of tissue types. The information can be used to avoid pitfalls that may lead to a failed seal.

[0925] FIG. **77** is a diagram of the right lung **4156** and FIG. **78** is a diagram of the bronchial tree **4250** including the trachea **4252** and the bronchi **4254, 4256** of the lungs. As shown in FIG. **77**, the right lung **4156** is composed of three lobes divided into the upper lobe **4162**, the middle lobe **4160**, and the lower lobe **4165** separated by the oblique fissure **4167** and horizontal fissure **4160**. The left lung is composed of only two smaller lobes due to the position of heart. As shown in FIG. **78**, inside each lung, the right bronchus **4254** and the left bronchus **4256** divide into many smaller airways called bronchioles **4258**, greatly increasing surface area. Each bronchiole **4258** terminates with a cluster of air sacs called alveoli **4260**, where gas exchange with the bloodstream occurs.

[0926] FIG. **79** is a logic flow diagram **4300** of a process depicting a control program or a logic configuration for storing paired anonymous data sets grouped by surgery, according to one aspect of the present disclosure. With reference to FIGS. **1-79**, in one aspect, the present disclosure provides a surgical hub **206** configured to communicate with a surgical instrument **235**. The surgical hub **206** comprises a processor **244** and a memory **249** coupled to the processor **244**. The memory **249** storing instructions executable by the processor **244** to receive **4302** a first data set from a first source, the first data set associated with a surgical procedure, receive **4304** a second data set from a second source, the second data set associated with the efficacy of the surgical procedure, anonymize **4306** the first and second data sets by removing information that identifies a patient, a surgery, or a scheduled time of the surgery, and store **4308** the first and second anonymized data sets to generate a data pair grouped by surgery. The first data set is generated at a first time, the second data set is generated at a second time, and the second time is separate and distinct from the first time.

[0927] In another aspect, the memory **249** stores instructions executable by the processor **244** to reconstruct a series

of chronological events based on the data pair. In another aspect, the memory 249 stores instructions executable by the processor 244 to reconstruct a series of coupled but unconstrained data sets based on the data pair. In another aspect, the memory 249 stores instructions executable by the processor 244 to encrypt the data pair, define a backup format for the data pair, and mirror the data pair to a cloud 204 storage device 205.

#### Determination of Data to Transmit to Cloud Based Medical Analytics

[0928] In one aspect, the present disclosure provides a communication hub and storage device for storing parameters and status of a surgical device what has the ability to determine when, how often, transmission rate, and type of data to be shared with a cloud based analytics system. The disclosure further provides techniques to determine where the analytics system communicates new operational parameters for the hub and surgical devices.

[0929] In a surgical hub environment, large amounts of data can be generated rather quickly and may cause storage and communication bottlenecks in the surgical hub network. One solution may include local determination of when and what data is transmitted for to the cloud-based medical analytics system for further processing and manipulation of surgical hub data. The timing and rate at which the surgical hub data is exported can be determined based on available local data storage capacity. User defined inclusion or exclusion of specific users, patients, or procedures enable data sets to be included for analysis or automatically deleted. The time of uploads or communications to the cloud-based medical analytics system may be determined based on detected surgical hub network down time or available capacity.

[0930] With reference to FIGS. 1-79, in one aspect, the present disclosure provides a surgical hub 206 comprising a storage device 248, a processor 244 coupled to the storage device 248, and a memory 249 coupled to the processor 244. The memory 249 stores instructions executable by the processor 244 to receive data from a surgical instrument 235, determine a rate at which to transfer the data to a remote cloud-based medical analytics network 204 based on available storage capacity of the storage device 248, determine a frequency at which to transfer the data to the remote cloud-based medical analytics network 204 based on the available storage capacity of the storage device 248 or detected surgical hub network 206 down time, and determine a type of data to transfer the data to a remote cloud-based medical analytics network 204 based on inclusion or exclusion of data associated with a users, patient, or surgical procedure.

[0931] In another aspect, the memory 249 stores instructions executable by the processor 244 to receive new operational parameters for the surgical hub 206 or the surgical instrument 235.

[0932] In various aspects, the present disclosure provides a control circuit to determine, rate, frequency and type of data to transfer the data to the remote cloud-based medical analytics network as described above. In various aspects, the present disclosure provides a non-transitory computer-readable medium storing computer readable instructions which, when executed, causes a machine to determine, rate, frequency and type of data to transfer to the remote cloud-based medical analytics network.

[0933] In one aspect, the surgical hub 206 is configured to determine what data to transmit to the cloud based analytics system 204. For example, a surgical hub 206 modular device 235 that includes local processing capabilities may determine the rate, frequency, and type of data to be transmitted to the cloud based analytics system 204 for analysis and processing.

[0934] In one aspect, the surgical hub 206 comprises a modular communication hub 203 and storage device 248 for storing parameters and status of a device 235 that has the ability to determine when and how often data can be shared with a cloud based analytics system 204, the transmission rate and the type of data that can be shared with the cloud based analytics system 204. In another aspect, the cloud analytics system 204 communicates new operational parameters for the surgical hub 206 and surgical devices 235 coupled to the surgical hub 206. A cloud based analytics system 204 is described in commonly-owned U.S. Provisional Patent Application Ser. No. 62/611,340, filed Dec. 28, 2017, and titled CLOUD-BASED MEDICAL ANALYTICS, which is incorporated herein by reference in its entirety.

[0935] In one aspect, a device 235 coupled to a local surgical hub 206 determines when and what data is transmitted to the cloud analytics system 204 for company analytic improvements. In one example, the available local data storage capacity remaining in the storage device 248 controls the timing and rate at which the data is exported. In another example, user defined inclusion or exclusion of specific users, patients, or procedures allows data sets to be included for analysis or automatically deleted. In yet another example, detected network down time or available capacity determines the time of uploads or communications.

[0936] In another aspect, transmission of data for diagnosis of failure modes is keyed by specific incidents. For example, user defined failure of a device, instrument, or tool within a procedure initiates archiving and transmission of data recorded with respect to that instrument for failure modes analysis. Further, when a failure event is identified, all the data surrounding the event is archived and packaged for sending back for predictive informatics (PI) analytics. Data that is part of a PI failure is flagged for storage and maintenance until either the hospital or the cloud based analytics system releases the hold on the data.

[0937] Catastrophic failures of instruments may initiate an automatic archive and submission of data for implications analysis. Detection of a counterfeit component or adapter on an original equipment manufacturer (OEM) device initiates documentation of the component and recording of the results and outcome of its use.

[0938] FIG. 80 is a logic flow diagram 4320 of a process depicting a control program or a logic configuration for determining rate, frequency, and type of data to transfer to a remote cloud-based analytics network, according to one aspect of the present disclosure. With reference to FIGS. 1-80, in one aspect, the present disclosure provides a surgical hub 206 comprising a storage device 248, a processor 244 coupled to the storage device 248, and a memory 249 coupled to the processor 244. The memory 249 stores instructions executable by the processor 244 to receive 4322 data from a surgical instrument 235, determine 4324 a rate at which to transfer the data to a remote cloud-based medical analytics network 204 based on available storage capacity of the storage device 248. Optionally, the memory 249 stores instructions executable by the processor 244 to determine

**4326** a frequency at which to transfer the data to the remote cloud-based medical analytics network **204** based on the available storage capacity of the storage device **248**. Optionally, the memory **249** stores instructions executable by the processor **244** to detect surgical hub network downtime and to determine **4326** a frequency at which to transfer the data to the remote cloud-based medical analytics network **204** based on the detected surgical hub network **206** down time. Optionally, the memory **249** stores instructions executable by the processor **244** to determine **4328** a type of data to transfer the data to a remote cloud-based medical analytics network **204** based on inclusion or exclusion of data associated with a users, patient, or surgical procedure.

[**0939**] In another aspect, the memory **249** stores instructions executable by the processor **244** to receive new operational parameters for the surgical hub **206** or the surgical instrument **235**.

[**0940**] In one aspect, the present disclosure provides a surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: interrogate a surgical instrument, wherein the surgical instrument is a first source of patient data; retrieve a first data set from the surgical instrument, wherein the first data set is associated with a patient and a surgical procedure; interrogate a medical imaging device, wherein the medical imaging device is a second source of patient data; retrieve a second data set from the medical imaging device, wherein the second data set is associated with the patient and an outcome of the surgical procedure; associate the first and second data sets by a key; and transmit the associated first and second data sets to remote network outside of the surgical hub. The present disclosure further provides, a surgical hub wherein the memory stores instructions executable by the processor to: retrieve the first data set using the key; anonymize the first data set by removing its association with the patient; retrieve the second data set using the key; anonymize the second data set by removing its association with the patient; pair the anonymized first and second data sets; and determine success rates of surgical procedures grouped by the surgical procedure based on the anonymized paired first and second data sets. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to: retrieve the anonymized first data set; retrieve the anonymized second data set; and reintegrate the anonymized first and second data sets using the key. The present disclosure further provides a surgical hub, wherein the first and second data sets define first and second data payloads in respective first and second data packets. The present disclosure further provides a control circuit to perform any one of the above recited functions and/or a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to perform any one of the above recited functions.

[**0941**] In another aspect, the present disclosure provides a surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive a first data packet from a first source, the first data packet comprising a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate, wherein the first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet; receive a second data packet from a second source,

the second data packet comprising a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate, wherein the second preamble defines the second data payload and the second encryption certificate verifies the authenticity of the second data packet; associate the first and second data packets; and generate a third data packet comprising the first and second data payloads. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to: determine that a data payload is from a new source; verify the new source of the data payload; and alter a data collection process at the surgical hub to receive subsequent data packets from the new source. The present disclosure further provides a surgical, wherein the memory stores instructions executable by the processor to associate the first and second data packets based on a key. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to anonymize the data payload of the third data packet. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to receive an anonymized third data packet and reintegrate the anonymized third data packet into the first and second data packets using the key. The present disclosure further provides a control circuit to perform any one of the above recited functions and/or a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to perform any one of the above recited functions.

[**0942**] In another aspect, the present disclosure provides a surgical hub configured to communicate with a surgical instrument, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive a first data set associated with a surgical procedure, wherein the first data set is generated at a first time; receive a second data set associated with the efficacy of the surgical procedure, wherein the second data set is generated at a second time, wherein the second time is separate and distinct from the first time; anonymize the first and second data sets by removing information that identifies a patient, a surgery, or a scheduled time of the surgery; and store the first and second anonymized data sets to generate a data pair grouped by surgery. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to reconstruct a series of chronological events based on the data pair. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to reconstruct a series of coupled but unconstrained data sets based on the data pair. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to: encrypt the data pair; define a backup format for the data pair; and mirror the data pair to a cloud storage device. The present disclosure further provides a control circuit to perform any one of the above recited functions and/or a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to perform any one of the above recited functions.

[**0943**] In another aspect, the present disclosure provides a surgical hub comprising: a storage device; a processor coupled to the storage device; and a memory coupled to the processor, the memory storing instructions executable by the

processor to: receive data from a surgical instrument; determine a rate at which to transfer the data to a remote cloud-based medical analytics network based on available storage capacity of the storage device; determine a frequency at which to transfer the data to the remote cloud-based medical analytics network based on the available storage capacity of the storage device or detected surgical hub network down time; and determine a type of data to transfer the data to a remote cloud-based medical analytics network based on inclusion or exclusion of data associated with a users, patient, or surgical procedure. The present disclosure further provides a surgical hub, wherein the memory stores instructions executable by the processor to receive new operational parameters for the surgical hub or the surgical instrument. The present disclosure further provides a control circuit to perform any one of the above recited functions and/or a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to perform any one of the above recited functions.

**[0944]** In another aspect, the present disclosure provides a surgical hub comprising: a control configured to: receive data from a surgical instrument; determine a rate at which to transfer the data to a remote cloud-based medical analytics network based on available storage capacity of the storage device; determine a frequency at which to transfer the data to the remote cloud-based medical analytics network based on the available storage capacity of the storage device or detected surgical hub network down time; and determine a type of data to transfer the data to a remote cloud-based medical analytics network based on inclusion or exclusion of data associated with a users, patient, or surgical procedure.

#### Surgical Hub Situational Awareness

**[0945]** Although an “intelligent” device including control algorithms that respond to sensed data can be an improvement over a “dumb” device that operates without accounting for sensed data, some sensed data can be incomplete or inconclusive when considered in isolation, i.e., without the context of the type of surgical procedure being performed or the type of tissue that is being operated on. Without knowing the procedural context (e.g., knowing the type of tissue being operated on or the type of procedure being performed), the control algorithm may control the modular device incorrectly or suboptimally given the particular context-free sensed data. For example, the optimal manner for a control algorithm to control a surgical instrument in response to a particular sensed parameter can vary according to the particular tissue type being operated on. This is due to the fact that different tissue types have different properties (e.g., resistance to tearing) and thus respond differently to actions taken by surgical instruments. Therefore, it may be desirable for a surgical instrument to take different actions even when the same measurement for a particular parameter is sensed. As one specific example, the optimal manner in which to control a surgical stapling and cutting instrument in response to the instrument sensing an unexpectedly high force to close its end effector will vary depending upon whether the tissue type is susceptible or resistant to tearing. For tissues that are susceptible to tearing, such as lung tissue, the instrument’s control algorithm would optimally ramp down the motor in response to an unexpectedly high force to close to avoid tearing the tissue. For tissues that are resistant

to tearing, such as stomach tissue, the instrument’s control algorithm would optimally ramp up the motor in response to an unexpectedly high force to close to ensure that the end effector is clamped properly on the tissue. Without knowing whether lung or stomach tissue has been clamped, the control algorithm may make a suboptimal decision.

**[0946]** One solution utilizes a surgical hub including a system that is configured to derive information about the surgical procedure being performed based on data received from various data sources and then control the paired modular devices accordingly. In other words, the surgical hub is configured to infer information about the surgical procedure from received data and then control the modular devices paired to the surgical hub based upon the inferred context of the surgical procedure. FIG. 81 illustrates a diagram of a situationally aware surgical system 5100, in accordance with at least one aspect of the present disclosure. In some exemplifications, the data sources 5126 include, for example, the modular devices 5102 (which can include sensors configured to detect parameters associated with the patient and/or the modular device itself), databases 5122 (e.g., an EMR database containing patient records), and patient monitoring devices 5124 (e.g., a blood pressure (BP) monitor and an electrocardiography (EKG) monitor). The surgical hub 5104 can be configured to derive the contextual information pertaining to the surgical procedure from the data based upon, for example, the particular combination(s) of received data or the particular order in which the data is received from the data sources 5126. The contextual information inferred from the received data can include, for example, the type of surgical procedure being performed, the particular step of the surgical procedure that the surgeon is performing, the type of tissue being operated on, or the body cavity that is the subject of the procedure. This ability by some aspects of the surgical hub 5104 to derive or infer information related to the surgical procedure from received data can be referred to as “situational awareness.” In one exemplification, the surgical hub 5104 can incorporate a situational awareness system, which is the hardware and/or programming associated with the surgical hub 5104 that derives contextual information pertaining to the surgical procedure from the received data.

**[0947]** The situational awareness system of the surgical hub 5104 can be configured to derive the contextual information from the data received from the data sources 5126 in a variety of different ways. In one exemplification, the situational awareness system includes a pattern recognition system, or machine learning system (e.g., an artificial neural network), that has been trained on training data to correlate various inputs (e.g., data from databases 5122, patient monitoring devices 5124, and/or modular devices 5102) to corresponding contextual information regarding a surgical procedure. In other words, a machine learning system can be trained to accurately derive contextual information regarding a surgical procedure from the provided inputs. In another exemplification, the situational awareness system can include a lookup table storing pre-characterized contextual information regarding a surgical procedure in association with one or more inputs (or ranges of inputs) corresponding to the contextual information. In response to a query with one or more inputs, the lookup table can return the corresponding contextual information for the situational awareness system for controlling the modular devices 5102. In one exemplification, the contextual information received by the

situational awareness system of the surgical hub **5104** is associated with a particular control adjustment or set of control adjustments for one or more modular devices **5102**. In another exemplification, the situational awareness system includes a further machine learning system, lookup table, or other such system, which generates or retrieves one or more control adjustments for one or more modular devices **5102** when provided the contextual information as input.

**[0948]** A surgical hub **5104** incorporating a situational awareness system provides a number of benefits for the surgical system **5100**. One benefit includes improving the interpretation of sensed and collected data, which would in turn improve the processing accuracy and/or the usage of the data during the course of a surgical procedure. To return to a previous example, a situationally aware surgical hub **5104** could determine what type of tissue was being operated on; therefore, when an unexpectedly high force to close the surgical instrument's end effector is detected, the situationally aware surgical hub **5104** could correctly ramp up or ramp down the motor of the surgical instrument for the type of tissue.

**[0949]** As another example, the type of tissue being operated can affect the adjustments that are made to the compression rate and load thresholds of a surgical stapling and cutting instrument for a particular tissue gap measurement. A situationally aware surgical hub **5104** could infer whether a surgical procedure being performed is a thoracic or an abdominal procedure, allowing the surgical hub **5104** to determine whether the tissue clamped by an end effector of the surgical stapling and cutting instrument is lung (for a thoracic procedure) or stomach (for an abdominal procedure) tissue. The surgical hub **5104** could then adjust the compression rate and load thresholds of the surgical stapling and cutting instrument appropriately for the type of tissue.

**[0950]** As yet another example, the type of body cavity being operated in during an insufflation procedure can affect the function of a smoke evacuator. A situationally aware surgical hub **5104** could determine whether the surgical site is under pressure (by determining that the surgical procedure is utilizing insufflation) and determine the procedure type. As a procedure type is generally performed in a specific body cavity, the surgical hub **5104** could then control the motor rate of the smoke evacuator appropriately for the body cavity being operated in. Thus, a situationally aware surgical hub **5104** could provide a consistent amount of smoke evacuation for both thoracic and abdominal procedures.

**[0951]** As yet another example, the type of procedure being performed can affect the optimal energy level for an ultrasonic surgical instrument or radio frequency (RF) electro-surgical instrument to operate at. Arthroscopic procedures, for example, require higher energy levels because the end effector of the ultrasonic surgical instrument or RF electro-surgical instrument is immersed in fluid. A situationally aware surgical hub **5104** could determine whether the surgical procedure is an arthroscopic procedure. The surgical hub **5104** could then adjust the RF power level or the ultrasonic amplitude of the generator (i.e., "energy level") to compensate for the fluid filled environment. Relatedly, the type of tissue being operated on can affect the optimal energy level for an ultrasonic surgical instrument or RF electro-surgical instrument to operate at. A situationally aware surgical hub **5104** could determine what type of surgical procedure is being performed and then customize the energy level for the ultrasonic surgical instrument or RF

electrosurgical instrument, respectively, according to the expected tissue profile for the surgical procedure. Furthermore, a situationally aware surgical hub **5104** can be configured to adjust the energy level for the ultrasonic surgical instrument or RF electro-surgical instrument throughout the course of a surgical procedure, rather than just on a procedure-by-procedure basis. A situationally aware surgical hub **5104** could determine what step of the surgical procedure is being performed or will subsequently be performed and then update the control algorithms for the generator and/or ultrasonic surgical instrument or RF electro-surgical instrument to set the energy level at a value appropriate for the expected tissue type according to the surgical procedure step.

**[0952]** As yet another example, data can be drawn from additional data sources **5126** to improve the conclusions that the surgical hub **5104** draws from one data source **5126**. A situationally aware surgical hub **5104** could augment data that it receives from the modular devices **5102** with contextual information that it has built up regarding the surgical procedure from other data sources **5126**. For example, a situationally aware surgical hub **5104** can be configured to determine whether hemostasis has occurred (i.e., whether bleeding at a surgical site has stopped) according to video or image data received from a medical imaging device. However, in some cases the video or image data can be inconclusive. Therefore, in one exemplification, the surgical hub **5104** can be further configured to compare a physiologic measurement (e.g., blood pressure sensed by a BP monitor communicably connected to the surgical hub **5104**) with the visual or image data of hemostasis (e.g., from a medical imaging device **124** (FIG. 2) communicably coupled to the surgical hub **5104**) to make a determination on the integrity of the staple line or tissue weld. In other words, the situational awareness system of the surgical hub **5104** can consider the physiological measurement data to provide additional context in analyzing the visualization data. The additional context can be useful when the visualization data may be inconclusive or incomplete on its own.

**[0953]** Another benefit includes proactively and automatically controlling the paired modular devices **5102** according to the particular step of the surgical procedure that is being performed to reduce the number of times that medical personnel are required to interact with or control the surgical system **5100** during the course of a surgical procedure. For example, a situationally aware surgical hub **5104** could proactively activate the generator to which an RF electro-surgical instrument is connected if it determines that a subsequent step of the procedure requires the use of the instrument. Proactively activating the energy source allows the instrument to be ready for use as soon as the preceding step of the procedure is completed.

**[0954]** As another example, a situationally aware surgical hub **5104** could determine whether the current or subsequent step of the surgical procedure requires a different view or degree of magnification on the display according to the feature(s) at the surgical site that the surgeon is expected to need to view. The surgical hub **5104** could then proactively change the displayed view (supplied by, e.g., a medical imaging device for the visualization system **108**) accordingly so that the display automatically adjusts throughout the surgical procedure.

**[0955]** As yet another example, a situationally aware surgical hub **5104** could determine which step of the surgical procedure is being performed or will subsequently be per-

formed and whether particular data or comparisons between data will be required for that step of the surgical procedure. The surgical hub **5104** can be configured to automatically call up data screens based upon the step of the surgical procedure being performed, without waiting for the surgeon to ask for the particular information.

[0956] Another benefit includes checking for errors during the setup of the surgical procedure or during the course of the surgical procedure. For example, a situationally aware surgical hub **5104** could determine whether the operating theater is setup properly or optimally for the surgical procedure to be performed. The surgical hub **5104** can be configured to determine the type of surgical procedure being performed, retrieve the corresponding checklists, product location, or setup needs (e.g., from a memory), and then compare the current operating theater layout to the standard layout for the type of surgical procedure that the surgical hub **5104** determines is being performed. In one exemplification, the surgical hub **5104** can be configured to compare the list of items for the procedure (scanned by the scanner **5132** depicted in FIG. **85B**, for example) and/or a list of devices paired with the surgical hub **5104** to a recommended or anticipated manifest of items and/or devices for the given surgical procedure. If there are any discontinuities between the lists, the surgical hub **5104** can be configured to provide an alert indicating that a particular modular device **5102**, patient monitoring device **5124**, and/or other surgical item is missing. In one exemplification, the surgical hub **5104** can be configured to determine the relative distance or position of the modular devices **5102** and patient monitoring devices **5124** via proximity sensors, for example. The surgical hub **5104** can compare the relative positions of the devices to a recommended or anticipated layout for the particular surgical procedure. If there are any discontinuities between the layouts, the surgical hub **5104** can be configured to provide an alert indicating that the current layout for the surgical procedure deviates from the recommended layout.

[0957] As another example, a situationally aware surgical hub **5104** could determine whether the surgeon (or other medical personnel) was making an error or otherwise deviating from the expected course of action during the course of a surgical procedure. For example, the surgical hub **5104** can be configured to determine the type of surgical procedure being performed, retrieve the corresponding list of steps or order of equipment usage (e.g., from a memory), and then compare the steps being performed or the equipment being used during the course of the surgical procedure to the expected steps or equipment for the type of surgical procedure that the surgical hub **5104** determined is being performed. In one exemplification, the surgical hub **5104** can be configured to provide an alert indicating that an unexpected action is being performed or an unexpected device is being utilized at the particular step in the surgical procedure.

[0958] Overall, the situational awareness system for the surgical hub **5104** improves surgical procedure outcomes by adjusting the surgical instruments (and other modular devices **5102**) for the particular context of each surgical procedure (such as adjusting to different tissue types) and validating actions during a surgical procedure. The situational awareness system also improves surgeons' efficiency in performing surgical procedures by automatically suggesting next steps, providing data, and adjusting displays and

other modular devices **5102** in the surgical theater according to the specific context of the procedure.

[0959] FIG. **82A** illustrates a logic flow diagram of a process **5000a** for controlling a modular device **5102** according to contextual information derived from received data, in accordance with at least one aspect of the present disclosure. In other words, a situationally aware surgical hub **5104** can execute the process **5000a** to determine appropriate control adjustments for modular devices **5102** paired with the surgical hub **5104** before, during, or after a surgical procedure as dictated by the context of the surgical procedure. In the following description of the process **5000a**, reference should also be made to FIG. **81**. In one exemplification, the process **5000a** can be executed by a control circuit of a surgical hub **5104**, as depicted in FIG. **10** (processor **244**). In another exemplification, the process **5000a** can be executed by a cloud computing system **104**, as depicted in FIG. **1**. In yet another exemplification, the process **5000a** can be executed by a distributed computing system including at least one of the aforementioned cloud computing system **104** and/or a control circuit of a surgical hub **5104** in combination with a control circuit of a modular device, such as the microcontroller **461** of the surgical instrument depicted in FIG. **12**, the microcontroller **620** of the surgical instrument depicted in FIG. **16**, the control circuit **710** of the robotic surgical instrument **700** depicted in FIG. **17**, the control circuit **760** of the surgical instruments **750**, **790** depicted in FIGS. **18** and **19**, or the controller **838** of the generator **800** depicted in FIG. **20**. For economy, the following description of the process **5000a** will be described as being executed by the control circuit of a surgical hub **5104**; however, it should be understood that the description of the process **5000a** encompasses all of the aforementioned exemplifications.

[0960] The control circuit of the surgical hub **5104** executing the process **5000a** receives **5004a** data from one or more data sources **5126** to which the surgical hub **5104** is communicably connected. The data sources **5126** include, for example, databases **5122**, patient monitoring devices **5124**, and modular devices **5102**. In one exemplification, the databases **5122** can include a patient EMR database associated with the medical facility at which the surgical procedure is being performed. The data received **5004a** from the data sources **5126** can include perioperative data, which includes preoperative data, intraoperative data, and/or post-operative data associated with the given surgical procedure. The data received **5004a** from the databases **5122** can include the type of surgical procedure being performed or the patient's medical history (e.g., medical conditions that may or may not be the subject of the present surgical procedure). In one exemplification depicted in FIG. **83A**, the control circuit can receive **5004a** the patient or surgical procedure data by querying the patient EMR database with a unique identifier associated with the patient. The surgical hub **5104** can receive the unique identifier from, for example, a scanner **5128** for scanning the patient's wristband **5130** encoding the unique identifier associated with the patient when the patient enters the operating theater, as depicted in FIG. **85A**. In one exemplification, the patient monitoring devices **5124** include BP monitors, EKG monitors, and other such devices that are configured to monitor one or more parameters associated with a patient. As with the modular devices **5102**, the patient monitoring devices **5124** can be paired with the surgical hub **5104** such that the



surgical hub **5104** receives **5004a** data therefrom. In one exemplification, the data received **5004a** from the modular devices **5102** that are paired with (i.e., communicably coupled to) the surgical hub **5104** includes, for example, activation data (i.e., whether the device is powered on or in use), data of the internal state of the modular device **5102** (e.g., force to fire or force to close for a surgical cutting and stapling device, pressure differential for an insufflator or smoke evacuator, or energy level for an RF or ultrasonic surgical instrument), or patient data (e.g., tissue type, tissue thickness, tissue mechanical properties, respiration rate, or airway volume).

[0961] As the process **5000a** continues, the control circuit of the surgical hub **5104** can derive **5006a** contextual information from the data received **5004a** from the data sources **5126**. The contextual information can include, for example, the type of procedure being performed, the particular step being performed in the surgical procedure, the patient's state (e.g., whether the patient is under anesthesia or whether the patient is in the operating room), or the type of tissue being operated on. The control circuit can derive **5006a** contextual information according to data from either an individual data source **5126** or combinations of data sources **5126**. Further, the control circuit can derive **5006a** contextual information according to, for example, the type (s) of data that it receives, the order in which the data is received, or particular measurements or values associated with the data. For example, if the control circuit receives data from an RF generator indicating that the RF generator has been activated, the control circuit could thus infer that the RF electro-surgical instrument is now in use and that the surgeon is or will be performing a step of the surgical procedure utilizing the particular instrument. As another example, if the control circuit receives data indicating that a laparoscope imaging device has been activated and an ultrasonic generator is subsequently activated, the control circuit can infer that the surgeon is on a laparoscopic dissection step of the surgical procedure due to the order in which the events occurred. As yet another example, if the control circuit receives data from a ventilator indicating that the patient's respiration is below a particular rate, then the control circuit can determine that the patient is under anesthesia.

[0962] The control circuit can then determine **5008a** what control adjustments are necessary (if any) for one or more modular devices **5102** according to the derived **5006a** contextual information. After determining **5008a** the control adjustments, the control circuit of the surgical hub **5104** can then control **5010a** the modular devices according to the control adjustments (if the control circuit determined **5008a** that any were necessary). For example, if the control circuit determines that an arthroscopic procedure is being performed and that the next step in the procedure utilizes an RF or ultrasonic surgical instrument in a liquid environment, the control circuit can determine **5008a** that a control adjustment for the generator of the RF or ultrasonic surgical instrument is necessary to preemptively increase the energy output of the instrument (because such instruments require increased energy in liquid environments to maintain their effectiveness). The control circuit can then control **5010a** the generator and/or the RF or ultrasonic surgical instrument accordingly by causing the generator to increase its output and/or causing the RF or ultrasonic surgical instrument to increase the energy drawn from the generator. The control

circuit can control **5010a** the modular devices **5102** according to the determined **5008a** control adjustment by, for example, transmitting the control adjustments to the particular modular device to update the modular device's **5102** programming. In another exemplification wherein the modular device(s) **5102** and the surgical hub **5104** are executing a distributed computing architecture, the control circuit can control **5010a** the modular device **5102** according to the determined **5008a** control adjustments by updating the distributed program.

[0963] FIGS. **82B-D** illustrate representative implementations of the process **5000a** depicted in FIG. **82A**. As with the process **5000a** depicted in FIG. **82A**, the processes illustrated in FIGS. **82B-D** can, in one exemplification, be executed by a control circuit of the surgical hub **5104**. FIG. **82B** illustrates a logic flow diagram of a process **5000b** for controlling a second modular device according to contextual information derived from perioperative data received from a first modular device, in accordance with at least one aspect of the present disclosure. In the illustrated exemplification, the control circuit of the surgical hub **5104** receives **5004b** perioperative data from a first modular device. The perioperative data can include, for example, data regarding the modular device **5102** itself (e.g., pressure differential, motor current, internal forces, or motor torque) or data regarding the patient with which the modular device **5102** is being utilized (e.g., tissue properties, respiration rate, airway volume, or laparoscopic image data). After receiving **5004b** the perioperative data, the control circuit of the surgical hub **5104** derives **5006b** contextual information from the perioperative data. The contextual information can include, for example, the procedure type, the step of the procedure being performed, or the status of the patient. The control circuit of the surgical hub **5104** then determines **5008b** control adjustments for a second modular device based upon the derived **5006b** contextual information and then controls **5010b** the second modular device accordingly. For example, the surgical hub **5104** can receive **5004b** perioperative data from a ventilator indicating that the patient's lung has been deflated, derive **5006b** the contextual information therefrom that the subsequent step in the particular procedure type utilizes a medical imaging device (e.g., a scope), determine **5008b** that the medical imaging device should be activated and set to a particular magnification, and then control **5010b** the medical imaging device accordingly.

[0964] FIG. **82C** illustrates a logic flow diagram of a process **5000c** for controlling a second modular device according to contextual information derived from perioperative data received from a first modular device and the second modular device. In the illustrated exemplification, the control circuit of the surgical hub **5104** receives **5002c** perioperative data from a first modular device and receives **5004c** perioperative data from a second modular device. After receiving **5002c**, **5004c** the perioperative data, the control circuit of the surgical hub **5104** derives **5006c** contextual information from the perioperative data. The control circuit of the surgical hub **5104** then determines **5008c** control adjustments for the second modular device based upon the derived **5006c** contextual information and then controls **5010c** the second modular device accordingly. For example, the surgical hub **5104** can receive **5002c** perioperative data from a RF electro-surgical instrument indicating that the instrument has been fired, receive **5004c** perioperative data from a surgical stapling instrument indi-

cating that the instrument has been fired, derive **5006c** the contextual information therefrom that the subsequent step in the particular procedure type requires that the surgical stapling instrument be fired with a particular force (because the optimal force to fire can vary according to the tissue type being operated on), determine **5008c** the particular force thresholds that should be applied to the surgical stapling instrument, and then control **5010c** the surgical stapling instrument accordingly.

[0965] FIG. 82D illustrates a logic flow diagram of a process **5000d** for controlling a third modular device according to contextual information derived from perioperative data received from a first modular device and a second modular device. In the illustrated exemplification, the control circuit of the surgical hub **5104** receives **5002d** perioperative data from a first modular device and receives **5004d** perioperative data from a second modular device. After receiving **5002d**, **5004d** the perioperative data, the control circuit of the surgical hub **5104** derives **5006d** contextual information from the perioperative data. The control circuit of the surgical hub **5104** then determines **5008d** control adjustments for a third modular device based upon the derived **5006d** contextual information and then controls **5010d** the third modular device accordingly. For example, the surgical hub **5104** can receive **5002d**, **5004d** perioperative data from an insufflator and a medical imaging device indicating that both devices have been activated and paired to the surgical hub **5104**, derive **5006d** the contextual information therefrom that a video-assisted thoracoscopic surgery (VATS) procedure is being performed, determine **5008d** that the displays connected to the surgical hub **5104** should be set to display particular views or information associated with the procedure type, and then control **5010d** the displays accordingly.

[0966] FIG. 83A illustrates a diagram of a surgical system **5100** including a surgical hub **5104** communicably coupled to a particular set of data sources **5126**. A surgical hub **5104** including a situational awareness system can utilize the data received from the data sources **5126** to derive contextual information regarding the surgical procedure that the surgical hub **5104**, the modular devices **5102** paired to the surgical hub **5104**, and the patient monitoring devices **5124** paired to the surgical hub **5104** are being utilized in connection with. The inferences (i.e., contextual information) that one exemplification of the situational awareness system can derive from the particular set of data sources **5126** are depicted in dashed boxes extending from the data source(s) **5126** from which they are derived. The contextual information derived from the data sources **5126** can include, for example, what step of the surgical procedure is being performed, whether and how a particular modular device **5102** is being used, and the patient's condition.

[0967] In the example illustrated in FIG. 83A, the data sources **5126** include a database **5122**, a variety of modular devices **5102**, and a variety of patient monitoring devices **5124**. The surgical hub **5104** can be connected to various databases **5122** to retrieve therefrom data regarding the surgical procedure that is being performed or is to be performed. In one exemplification of the surgical system **5100**, the databases **5122** include an EMR database of a hospital. The data that can be received by the situational awareness system of the surgical hub **5104** from the databases **5122** can include, for example, start (or setup) time or operational information regarding the procedure (e.g., a

segmentectomy in the upper right portion of the thoracic cavity). The surgical hub **5104** can derive contextual information regarding the surgical procedure from this data alone or from the combination of this data and data from other data sources **5126**.

[0968] The surgical hub **5104** can also be connected to (i.e., paired with) a variety of patient monitoring devices **5124**. In one exemplification of the surgical system **5100**, the patient monitoring devices **5124** that can be paired with the surgical hub **5104** can include a pulse oximeter (SpO<sub>2</sub> monitor) **5114**, a BP monitor **5116**, and an EKG monitor **5120**. The perioperative data that can be received by the situational awareness system of the surgical hub **5104** from the patient monitoring devices **5124** can include, for example, the patient's oxygen saturation, blood pressure, heart rate, and other physiological parameters. The contextual information that can be derived by the surgical hub **5104** from the perioperative data transmitted by the patient monitoring devices **5124** can include, for example, whether the patient is located in the operating theater or under anesthesia. The surgical hub **5104** can derive these inferences from data from the patient monitoring devices **5124** alone or in combination with data from other data sources **5126** (e.g., the ventilator **5118**).

[0969] The surgical hub **5104** can also be connected to (i.e., paired with) a variety of modular devices **5102**. In one exemplification of the surgical system **5100**, the modular devices **5102** that can be paired with the surgical hub **5104** can include a smoke evacuator **5106**, a medical imaging device **5108**, an insufflator **5110**, a combined energy generator **5112** (for powering an ultrasonic surgical instrument and/or an RF electrosurgical instrument), and a ventilator **5118**.

[0970] The medical imaging device **5108** includes an optical component and an image sensor that generates image data. The optical component includes a lens or a light source, for example. The image sensor includes a charge-coupled device (CCD) or a complementary metal-oxide-semiconductor (CMOS), for example. In various exemplifications, the medical imaging device **5108** includes an endoscope, a laparoscope, a thoracoscope, and other such imaging devices. Various additional components of the medical imaging device **5108** are described above. The perioperative data that can be received by the surgical hub **5104** from the medical imaging device **5108** can include, for example, whether the medical imaging device **5108** is activated and a video or image feed. The contextual information that can be derived by the surgical hub **5104** from the perioperative data transmitted by the medical imaging device **5108** can include, for example, whether the procedure is a VATS procedure (based on whether the medical imaging device **5108** is activated or paired to the surgical hub **5104** at the beginning or during the course of the procedure). Furthermore, the image or video data from the medical imaging device **5108** (or the data stream representing the video for a digital medical imaging device **5108**) can be processed by a pattern recognition system or a machine learning system to recognize features (e.g., organs or tissue types) in the field of view (FOV) of the medical imaging device **5108**, for example. The contextual information that can be derived by the surgical hub **5104** from the recognized features can include, for example, what type of surgical procedure (or step thereof) is being performed, what organ is being operated on, or what body cavity is being operated in.

[0971] In one exemplification depicted in FIG. 83B, the smoke evacuator 5106 includes a first pressure sensor  $P_1$  configured to detect the ambient pressure in the operating theater, a second pressure sensor  $P_2$  configured to detect the internal downstream pressure (i.e., the pressure downstream from the inlet), and a third pressure sensor  $P_3$  configured to detect the internal upstream pressure. In one exemplification, the first pressure sensor  $P_1$  can be a separate component from the smoke evacuator 5106 or otherwise located externally to the smoke evacuator 5106. The perioperative data that can be received by the surgical hub 5104 from the smoke evacuator 5106 can include, for example, whether the smoke evacuator 5106 is activated, pressure readings from each of the sensors  $P_1$ ,  $P_2$ ,  $P_3$ , and pressure differentials between pairs of the sensors  $P_1$ ,  $P_2$ ,  $P_3$ . The perioperative data can also include, for example, the type of tissue being operated on (based upon the chemical composition of the smoke being evacuated) and the amount of tissue being cut. The contextual information that can be derived by the surgical hub 5104 from the perioperative data transmitted by the smoke evacuator 5106 can include, for example, whether the procedure being performed is utilizing insufflation. The smoke evacuator 5106 perioperative data can indicate whether the procedure is utilizing insufflation according to the pressure differential between  $P_3$  and  $P_1$ . If the pressure sensed by  $P_3$  is greater than the pressure sensed by  $P_1$  (i.e.,  $P_3 - P_1 > 0$ ), then the body cavity to which the smoke evacuator 5106 is connected is insufflated. If the pressure sensed by  $P_3$  is equal to the pressure sensed by  $P_1$  (i.e.,  $P_3 - P_1 = 0$ ), then the body cavity is not insufflated. When the body cavity is not insufflated, the procedure may be an open type of procedure.

[0972] The insufflator 5110 can include, for example, pressure sensors and current sensors configured to detect internal parameters of the insufflator 5110. The perioperative data that can be received by the surgical hub 5104 from the insufflator can include, for example, whether the insufflator 5110 is activated and the electrical current drawn by the insufflator's 5110 pump. The surgical hub 5104 can determine whether the insufflator 5110 is activated by, for example, directly detecting whether the device is powered on, detecting whether there is a pressure differential between an ambient pressure sensor and a pressure sensor internal to the surgical site, or detecting whether the pressure valves of the insufflator 5110 are pressurized (activated) or non-pressurized (deactivated). The contextual information that can be derived by the surgical hub 5104 from the perioperative data transmitted by the insufflator 5110 can include, for example, the type of procedure being performed (e.g., insufflation is utilized in laparoscopic procedures, but not arthroscopic procedures) and what body cavity is being operated in (e.g., insufflation is utilized in the abdominal cavity, but not in the thoracic cavity). In some exemplifications, the inferences derived from the perioperative data received from different modular devices 5102 can be utilized to confirm and/or increase the confidence of prior inferences. For example, if the surgical hub 5104 determines that the procedure is utilizing insufflation because the insufflator 5110 is activated, the surgical hub 5104 can then confirm that inference by detecting whether the perioperative data from the smoke evacuator 5106 likewise indicates that the body cavity is insufflated.

[0973] The combined energy generator 5112 supplies energy to one or more ultrasonic surgical instruments or RF

electrosurgical instruments connected thereto. The perioperative data that can be received by the surgical hub 5104 from the combined energy generator 5112 can include, for example, the mode that the combined energy generator 5112 is set to (e.g., a vessel sealing mode or a cutting/coagulation mode). The contextual information that can be derived by the surgical hub 5104 from the perioperative data transmitted by the combined energy generator 5112 can include, for example, the surgical procedural type (based on the number and types of surgical instruments that are connected to the energy generator 5112) and the procedural step that is being performed (because the particular surgical instrument being utilized or the particular order in which the surgical instruments are utilized corresponds to different steps of the surgical procedure). Further, the inferences derived by the surgical hub 5104 can depend upon inferences and/or perioperative data previously received by the surgical hub 5104. Once the surgical hub 5104 has determined the general category or specific type of surgical procedure being performed, the surgical hub 5104 can determine or retrieve an expected sequence of steps for the surgical procedure and then track the surgeon's progression through the surgical procedure by comparing the detected sequence in which the surgical instruments are utilized relative to the expected sequence.

[0974] The perioperative data that can be received by the surgical hub 5104 from the ventilator 5118 can include, for example, the respiration rate and airway volume of the patient. The contextual information that can be derived by the surgical hub 5104 from the perioperative data transmitted by the ventilator 5118 can include, for example, whether the patient is under anesthesia and whether the patient's lung is deflated. In some exemplifications, certain contextual information can be inferred by the surgical hub 5104 based on combinations of perioperative data from multiple data sources 5126. For example, the situational awareness system of the surgical hub 5104 can be configured to infer that the patient is under anesthesia when the respiration rate detected by the ventilator 5118, the blood pressure detected by the BP monitor 5116, and the heart rate detected by the EKG monitor 5120 fall below particular thresholds. For certain contextual information, the surgical hub 5104 can be configured to only derive a particular inference when the perioperative data from a certain number or all of the relevant data sources 5126 satisfy the conditions for the inference.

[0975] As can be seen from the particular exemplified surgical system 5100, the situational awareness system of a surgical hub 5104 can derive a variety of contextual information regarding the surgical procedure being performed from the data sources 5126. The surgical hub 5104 can utilize the derived contextual information to control the modular devices 5102 and make further inferences about the surgical procedure in combination with data from other data sources 5126. It should be noted that the inferences depicted in FIG. 83A and described in connection with the depicted surgical system 5100 are merely exemplary and should not be interpreted as limiting in any way. Furthermore, the surgical hub 5104 can be configured to derive a variety of other inferences from the same (or different) modular devices 5102 and/or patient monitoring devices 5124. In other exemplifications, a variety of other modular devices 5102 and/or patient monitoring devices 5124 can be paired to the surgical hub 5104 in the operating theater and data

received from those additional modular devices **5102** and/or patient monitoring devices **5124** can be utilized by the surgical hub **5104** to derive the same or different contextual information about the particular surgical procedure being performed.

[0976] FIGS. **84A-J** depict logic flow diagrams for processes for deriving **5008a**, **5008b**, **5008c**, **5008d** contextual information from various modular devices, as discussed above with respect to the processes **5000a**, **5000b**, **5000c**, **5000d** depicted in FIGS. **82A-D**. The derived contextual information in FIGS. **84A-C** is the procedure type. The procedure type can correspond to techniques utilized during the surgical procedure (e.g., a segmentectomy), the category of the surgical procedure (e.g., a laparoscopic procedure), the organ, tissue, or other structure being operated on, and other characteristics to identify the particular surgical procedure (e.g., the procedure utilizes insufflation). The derived contextual information in FIGS. **84D-G** is the particular step of the surgical procedure that is being performed. The derived contextual information in FIGS. **84H-J** is the patient's status. It can be noted that the patient's status can also correspond to the particular step of the surgical procedure that is being performed (e.g., determining that the patient's status has changed from not being under anesthesia to being under anesthesia can indicate that the step of the surgical procedure of placing the patient under anesthesia was carried out by the surgical staff). As with the process **5000a** depicted in FIG. **82A**, the processes illustrated in FIGS. **84A-J** can, in one exemplification, be executed by a control circuit of the surgical hub **5104**. In the following descriptions of the processes illustrated in FIGS. **84A-J**, reference should also be made to FIG. **83A**.

[0977] FIG. **84A** illustrates a logic flow diagram of a process **5111** for determining a procedure type according to smoke evacuator **5106** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5111** receives **5113** perioperative data from the smoke evacuator **5106** and then determines **5115** whether the smoke evacuator **5106** is activated based thereon. If the smoke evacuator **5106** is not activated, then the process **5111** continues along the NO branch and the control circuit of the surgical hub **5104** continues monitoring for the receipt of smoke evacuator **5106** perioperative data. If the smoke evacuator **5106** is activated, then the process **5111** continues along the YES branch and the control circuit of the surgical hub **5104** determines **5117** whether there is a pressure differential between an internal upstream pressure sensor  $P_3$  (FIG. **83B**) and an external or ambient pressure sensor  $P_1$  (FIG. **83B**). If there is a pressure differential (i.e., the internal upstream pressure of the smoke evacuator **5106** is greater than the ambient pressure of the operating theater), then the process **5111** continues along the YES branch and the control circuit determines **5119** that the surgical procedure is an insufflation-utilizing procedure. If there is not a pressure differential, then the process **5111** continues along the NO branch and the control circuit determines **5121** that the surgical procedure is not an insufflation-utilizing procedure.

[0978] FIG. **84B** illustrates a logic flow diagram of a process **5123** for determining a procedure type according to smoke evacuator **5106**, insufflator **5110**, and medical imaging device **5108** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5123** receives **5125**, **5127**, **5129** perioperative data

from the smoke evacuator **5106**, insufflator **5110**, and medical imaging device **5108** and then determines **5131** whether all of the devices are activated or paired with the surgical hub **5104**. If all of these devices are not activated or paired with the surgical hub **5104**, then the process **5123** continues along the NO branch and the control circuit determines **5133** that the surgical procedure is not a VATS procedure. If all of the aforementioned devices are activated or paired with the surgical hub **5104**, then the process **5123** continues along the YES branch and the control circuit determines **5135** that the surgical procedure is a VATS procedure. The control circuit can make this determination based upon the fact that all of these devices are required for a VATS procedure; therefore, if not all of these devices are being utilized in the surgical procedure, it cannot be a VATS procedure.

[0979] FIG. **84C** illustrates a logic flow diagram of a process **5137** for determining a procedure type according to medical imaging device **5108** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5137** receives **5139** perioperative data from the medical imaging device **5108** and then determines **5141** whether the medical imaging device **5108** is transmitting an image or video feed. If the medical imaging device **5108** is not transmitting an image or video feed, then the process **5137** continues along the NO branch and the control circuit determines **5143** that the surgical procedure is not a VATS procedure. If the medical imaging device **5108** is not transmitting an image or video feed, then the process **5137** continues along the YES branch and the control circuit determines **5145** that the surgical procedure is a VATS procedure. In one exemplification, the control circuit of the surgical hub **5104** can execute the process **5137** depicted in FIG. **84C** in combination with the process **5123** depicted in FIG. **84B** in order to confirm or increase the confidence in the contextual information derived by both processes **5123**, **5137**. If there is a discontinuity between the determinations of the processes **5123**, **5137** (e.g., the medical imaging device **5108** is transmitting a feed, but not all of the requisite devices are paired with the surgical hub **5104**), then the surgical hub **5104** can execute additional processes to come to a final determination that resolves the discontinuities between the processes **5123**, **5137** or display an alert or feedback to the surgical staff as to the discontinuity.

[0980] FIG. **84D** illustrates a logic flow diagram of a process **5147** for determining a procedural step according to insufflator **5110** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5147** receives **5149** perioperative data from the insufflator **5110** and then determines **5151** whether there is a pressure differential between the surgical site and the ambient environment of the operating theater. In one exemplification, the insufflator **5110** perioperative data can include a surgical site pressure (e.g., the intra-abdominal pressure) sensed by a first pressure sensor associated with the insufflator **5110**, which can be compared against a pressure sensed by a second pressure sensor configured to detect the ambient pressure. The first pressure sensor can be configured to detect an intra-abdominal pressure between 0-10 mmHg, for example. If there is a pressure differential, then the process **5147** continues along the YES branch and the control circuit determines **5153** that an insufflation-utilizing step of the surgical procedure is being performed. If there is not a pressure differential, then the process **5147** continues along the NO branch and the control circuit

determines **5155** that an insufflation-utilizing step of the surgical procedure is not being performed.

[0981] FIG. **84E** illustrates a logic flow diagram of a process **5157** for determining a procedural step according to energy generator **5112** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5157** receives **5159** perioperative data from the energy generator **5112** and then determines **5161** whether the energy generator **5112** is in the sealing mode. In various exemplifications, the energy generator **5112** can include two modes: a sealing mode corresponding to a first energy level and a cut/coagulation mode corresponding to a second energy level. If the energy generator **5112** is not in the sealing mode, then the process **5157** proceeds along the NO branch and the control circuit determines **5163** that a dissection step of the surgical procedure is being performed. The control circuit can make this determination **5163** because if the energy generator **5112** is not on the sealing mode, then it must thus be on the cut/coagulation mode for energy generators **5112** having two modes of operation. The cut/coagulation mode of the energy generator **5112** corresponds to a dissection procedural step because it provides an appropriate degree of energy to the ultrasonic surgical instrument or RF electro-surgical instrument to dissect tissue. If the energy generator **5112** is in the sealing mode, then the process **5157** proceeds along the YES branch and the control circuit determines **5165** that a ligation step of the surgical procedure is being performed. The sealing mode of the energy generator **5112** corresponds to a ligation procedural step because it provides an appropriate degree of energy to the ultrasonic surgical instrument or RF electro-surgical instrument to ligate vessels.

[0982] FIG. **84F** illustrates a logic flow diagram of a process **5167** for determining a procedural step according to energy generator **5112** perioperative data. In various aspects, previously received perioperative data and/or previously derived contextual information can also be considered by processes in deriving subsequent contextual information. This allows the situational awareness system of the surgical hub **5104** to derive additional and/or increasingly detailed contextual information about the surgical procedure as the procedure is performed. In this exemplification, the process **5167** determines **5169** that a segmentectomy procedure is being performed. This contextual information can be derived by this process **5167** or other processes based upon other received perioperative data and/or retrieved from a memory. Subsequently, the control circuit receives **5171** perioperative data from the energy generator **5112** indicating that a surgical instrument is being fired and then determines **5173** whether the energy generator **5112** was utilized in a previous step of the procedure to fire the surgical instrument. The control circuit can determine **5173** whether the energy generator **5112** was previously utilized in a prior step of the procedure by retrieving a list of the steps that have been performed in the current surgical procedure from a memory, for example. In such exemplifications, when the surgical hub **5104** determines that a step of the surgical procedure has been performed or completed by the surgical staff, the surgical hub **5104** can update a list of the procedural steps that have been performed, which can then be subsequently retrieved by the control circuit of the surgical hub **5104**. In one exemplification, the surgical hub **5104** can distinguish between sequences of firings of the surgical instrument as corresponding to separate steps of the surgical procedure

according to the time delay between the sequences of firings, whether any intervening actions were taken or modular devices **5102** were utilized by the surgical staff, or other factors that the situational awareness system can detect. If the energy generator **5112** has not been previously utilized during the course of the segmentectomy procedure, the process **5167** proceeds along the NO branch and the control circuit determines **5175** that the step of dissecting tissue to mobilize the patient's lungs is being performed by the surgical staff. If the energy generator **5112** has been previously utilized during the course of the segmentectomy procedure, the process **5167** proceeds along the YES branch and the control circuit determines **5177** that the step of dissecting nodes is being performed by the surgical staff. An ultrasonic surgical instrument or RF electro-surgical instrument is utilized twice during the course of an example of a segmentectomy procedure (e.g., FIG. **86**); therefore, the situational awareness system of the surgical hub **5104** executing the process **5167** can distinguish between which step the utilization of the energy generator **5112** indicates is currently being performed based upon whether the energy generator **5112** was previously utilized.

[0983] FIG. **84G** illustrates a logic flow diagram of a process **5179** for determining a procedural step according to stapler perioperative data. As described above with respect to the process **5167** illustrated in FIG. **84F**, the process **5179** utilizes previously received perioperative data and/or previously derived contextual information in deriving subsequent contextual information. In this exemplification, the process **5179** determines **5181** that a segmentectomy procedure is being performed. This contextual information can be derived by this process **5179** or other processes based upon other received perioperative data and/or retrieved from a memory. Subsequently, the control circuit receives **5183** perioperative data from the surgical stapling instrument (i.e., stapler) indicating that the surgical stapling instrument is being fired and then determines **5185** whether the surgical stapling instrument was utilized in a previous step of the surgical procedure. As described above, the control circuit can determine **5185** whether the surgical stapling instrument was previously utilized in a prior step of the procedure by retrieving a list of the steps that have been performed in the current surgical procedure from a memory, for example. If the surgical stapling instrument has not been utilized previously, then the process **5179** proceeds along the NO branch and the control circuit determines **5187** that the step of ligating arteries and veins is being performed by the surgical staff. If the surgical stapling instrument has been previously utilized during the course of the segmentectomy procedure, the process **5179** proceeds along the YES branch and the control circuit determines **5189** that the step of transecting parenchyma is being performed by the surgical staff. A surgical stapling instrument is utilized twice during the course of an example of a segmentectomy procedure (e.g., FIG. **86**); therefore, the situational awareness system of the surgical hub **5104** executing the process **5179** can distinguish between which step the utilization of the surgical stapling instrument indicates is currently being performed based upon whether the surgical stapling instrument was previously utilized.

[0984] FIG. **84H** illustrates a logic flow diagram of a process **5191** for determining a patient status according to ventilator **5110**, pulse oximeter **5114**, BP monitor **5116**, and/or EKG monitor **5120** perioperative data. In this exem-

plification, the control circuit of the surgical hub **5104** executing the process **5191** receives **5193**, **5195**, **5197**, **5199** perioperative data from each of the ventilator **5110**, pulse oximeter **5114**, BP monitor **5116**, and/or EKG monitor **5120** and then determines whether one or more values of the physiological parameters sensed by each of the devices fall below a threshold for each of the physiological parameters. The threshold for each physiological parameter can correspond to a value that corresponds to a patient being under anesthesia. In other words, the control circuit determines **5201** whether the patient's respiration rate, oxygen saturation, blood pressure, and/or heart rate indicate that the patient is under anesthesia according data sensed by the respective modular device **5102** and/or patient monitoring devices **5124**. In one exemplification, if the all of the values from the perioperative data are below their respective thresholds, then the process **5191** proceeds along the YES branch and the control circuit determines **5203** that the patient is under anesthesia. In another exemplification, the control circuit can determine **5203** that the patient is under anesthesia if a particular number or ratio of the monitored physiological parameters indicate that the patient is under anesthesia. Otherwise, the process **5191** proceeds along the NO branch and the control circuit determines **5205** that the patient is not under anesthesia.

[0985] FIG. **84I** illustrates a logic flow diagram of a process **5207** for determining a patient status according to pulse oximeter **5114**, BP monitor **5116**, and/or EKG monitor **5120** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5207** receives **5209**, **5211**, **5213** (or attempts to receive) perioperative data the pulse oximeter **5114**, BP monitor **5116**, and/or EKG monitor **5120** and then determines **5215** whether at least one of the devices is paired with the surgical hub **5104** or the surgical hub **5104** is otherwise receiving data therefrom. If the control circuit is receiving data from at least one of these patient monitoring devices **5124**, the process **5207** proceeds along the YES branch and the control circuit determines **5217** that the patient is in the operating theater. The control circuit can make this determination because the patient monitoring devices **5214** connected to the surgical hub **5104** must be in the operating theater and thus the patient must likewise be in the operating theater. If the control circuit is not receiving data from at least one of these patient monitoring devices **5124**, the process **5207** proceeds along the NO branch and the control circuit determines **5219** that the patient is not in the operating theater.

[0986] FIG. **84J** illustrates a logic flow diagram of a process **5221** for determining a patient status according to ventilator **5110** perioperative data. In this exemplification, the control circuit of the surgical hub **5104** executing the process **5221** receives **5223** perioperative data from the ventilator **5110** and then determines **5225** whether the patient's airway volume has decreased or is decreasing. In one exemplification, the control circuit determines **5225** whether the patient's airway volume falls below a particular threshold value indicative of a lung having collapsed or been deflated. In another exemplification, the control circuit determines **5225** whether the patient's airway volume falls below an average or baseline level by a threshold amount. If the patient's airway volume has not decreased sufficiently, the process **5221** proceeds along the NO branch and the control circuit determines **5227** that the patient's lung is not

deflated. If the patient's airway volume has decreased sufficiently, the process **5221** proceeds along the YES branch and the control circuit determines **5229** that the patient's lung is not deflated.

[0987] In one exemplification, the surgical system **5100** can further include various scanners that can be paired with the surgical hub **5104** to detect and record objects and individuals that enter and exit the operating theater. FIG. **85A** illustrates a scanner **5128** paired with a surgical hub **5104** that is configured to scan a patient wristband **5130**. In one aspect, the scanner **5128** includes, for example, a barcode reader or a radio-frequency identification (RFID) reader that is able to read patient information from the patient wristband **5130** and then transmit that information to the surgical hub **5104**. The patient information can include the surgical procedure to be performed or identifying information that can be cross-referenced with the hospital's EMR database **5122** by the surgical hub **5104**, for example. FIG. **85B** illustrates a scanner **5132** paired with a surgical hub **5104** that is configured to scan a product list **5134** for a surgical procedure. The surgical hub **5104** can utilize data from the scanner **5132** regarding the number, type, and mix of items to be used in the surgical procedure to identify the type of surgical procedure being performed. In one exemplification, the scanner **5132** includes a product scanner (e.g., a barcode reader or an RFID reader) that is able to read the product information (e.g., name and quantity) from the product itself or the product packaging as the products are brought into the operating theater and then transmit that information to the surgical hub **5104**. In another exemplification, the scanner **5132** includes a camera (or other visualization device) and associated optical character recognition software that is able to read the product information from a product list **5134**. The surgical hub **5104** can be configured to cross-reference the list of items indicated by the received data with a lookup table or database of items utilized for various types of surgical procedures in order to infer the particular surgical procedure that is to be (or was) performed. As shown in FIG. **85B**, the illustrative product list **5134** includes ring forceps, rib spreaders, a powered vascular stapler (PVS), and a thoracic wound protector. In this example, the surgical hub **5104** can infer that the surgical procedure is a thoracic procedure from this data since these products are only utilized in thoracic procedures. In sum, the scanner(s) **5128**, **5132** can provide serial numbers, product lists, and patient information to the surgical hub **5104**. Based on this data regarding what devices and instruments are being utilized and the patient's medical information, the surgical hub **5104** can determine additional contextual information regarding the surgical procedure.

#### Situational Awareness

[0988] Situational awareness is the ability of some aspects of a surgical system to determine or infer information related to a surgical procedure from data received from databases and/or instruments. The information can include the type of procedure being undertaken, the type of tissue being operated on, or the body cavity that is the subject of the procedure. With the contextual information related to the surgical procedure, the surgical system can, for example, improve the manner in which it controls the modular devices (e.g., a robotic arm and/or robotic surgical tool) that are

connected to it and provide contextualized information or suggestions to the surgeon during the course of the surgical procedure.

[0989] In order to assist in the understanding of the process 5000a illustrated in FIG. 82A and the other concepts discussed above, FIG. 86 illustrates a timeline 5200 of an illustrative surgical procedure and the contextual information that a surgical hub 5104 can derive from the data received from the data sources 5126 at each step in the surgical procedure. In the following description of the timeline 5200 illustrated in FIG. 86, reference should also be made to FIG. 81. The timeline 5200 depicts the typical steps that would be taken by the nurses, surgeons, and other medical personnel during the course of a lung segmentectomy procedure, beginning with setting up the operating theater and ending with transferring the patient to a post-operative recovery room. The situationally aware surgical hub 5104 receives data from the data sources 5126 throughout the course of the surgical procedure, including data generated each time medical personnel utilize a modular device 5102 that is paired with the surgical hub 5104. The surgical hub 5104 can receive this data from the paired modular devices 5102 and other data sources 5126 and continually derive inferences (i.e., contextual information) about the ongoing procedure as new data is received, such as which step of the procedure is being performed at any given time. The situational awareness system of the surgical hub 5104 is able to, for example, record data pertaining to the procedure for generating reports (e.g., see FIGS. 90-101), verify the steps being taken by the medical personnel, provide data or prompts (e.g., via a display screen) that may be pertinent for the particular procedural step, adjust modular devices 5102 based on the context (e.g., activate monitors, adjust the FOV of the medical imaging device, or change the energy level of an ultrasonic surgical instrument or RF electrosurgical instrument), and take any other such action described above.

[0990] As the first step 5202 in this illustrative procedure, the hospital staff members retrieve the patient's EMR from the hospital's EMR database. Based on select patient data in the EMR, the surgical hub 5104 determines that the procedure to be performed is a thoracic procedure. Second 5204, the staff members scan the incoming medical supplies for the procedure. The surgical hub 5104 cross-references the scanned supplies with a list of supplies that are utilized in various types of procedures and confirms that the mix of supplies corresponds to a thoracic procedure (e.g., as depicted in FIG. 85B). Further, the surgical hub 5104 is also able to determine that the procedure is not a wedge procedure (because the incoming supplies either lack certain supplies that are necessary for a thoracic wedge procedure or do not otherwise correspond to a thoracic wedge procedure). Third 5206, the medical personnel scan the patient band (e.g., as depicted in FIG. 85A) via a scanner 5128 that is communicably connected to the surgical hub 5104. The surgical hub 5104 can then confirm the patient's identity based on the scanned data. Fourth 5208, the medical staff turns on the auxiliary equipment. The auxiliary equipment being utilized can vary according to the type of surgical procedure and the techniques to be used by the surgeon, but in this illustrative case they include a smoke evacuator, insufflator, and medical imaging device. When activated, the auxiliary equipment that are modular devices 5102 can automatically pair with the surgical hub 5104 that is located

within a particular vicinity of the modular devices 5102 as part of their initialization process. The surgical hub 5104 can then derive contextual information about the surgical procedure by detecting the types of modular devices 5102 that pair with it during this pre-operative or initialization phase. In this particular example, the surgical hub 5104 determines that the surgical procedure is a VATS procedure based on this particular combination of paired modular devices 5102. Based on the combination of the data from the patient's EMR, the list of medical supplies to be used in the procedure, and the type of modular devices 5102 that connect to the hub, the surgical hub 5104 can generally infer the specific procedure that the surgical team will be performing. Once the surgical hub 5104 knows what specific procedure is being performed, the surgical hub 5104 can then retrieve the steps of that procedure from a memory or from the cloud and then cross-reference the data it subsequently receives from the connected data sources 5126 (e.g., modular devices 5102 and patient monitoring devices 5124) to infer what step of the surgical procedure the surgical team is performing. Fifth 5210, the staff members attach the EKG electrodes and other patient monitoring devices 5124 to the patient. The EKG electrodes and other patient monitoring devices 5124 are able to pair with the surgical hub 5104. As the surgical hub 5104 begins receiving data from the patient monitoring devices 5124, the surgical hub 5104 thus confirms that the patient is in the operating theater, as described in the process 5207 depicted in FIG. 84I, for example. Sixth 5212, the medical personnel induce anesthesia in the patient. The surgical hub 5104 can infer that the patient is under anesthesia based on data from the modular devices 5102 and/or patient monitoring devices 5124, including EKG data, blood pressure data, ventilator data, or combinations thereof, as described in the process 5191 depicted in FIG. 84H, for example. Upon completion of the sixth step 5212, the pre-operative portion of the lung segmentectomy procedure is completed and the operative portion begins.

[0991] Seventh 5214, the patient's lung that is being operated on is collapsed (while ventilation is switched to the contralateral lung). The surgical hub 5104 can infer from the ventilator data that the patient's lung has been collapsed, as described in the process 5221 depicted in FIG. 84J, for example. The surgical hub 5104 can infer that the operative portion of the procedure has commenced as it can compare the detection of the patient's lung collapsing to the expected steps of the procedure (which can be accessed or retrieved previously) and thereby determine that collapsing the lung is the first operative step in this particular procedure. Eighth 5216, the medical imaging device 5108 (e.g., a scope) is inserted and video from the medical imaging device is initiated. The surgical hub 5104 receives the medical imaging device data (i.e., video or image data) through its connection to the medical imaging device. Upon receipt of the medical imaging device data, the surgical hub 5104 can determine that the laparoscopic portion of the surgical procedure has commenced. Further, the surgical hub 5104 can determine that the particular procedure being performed is a segmentectomy, as opposed to a lobectomy (note that a wedge procedure has already been discounted by the surgical hub 5104 based on data received at the second step 5204 of the procedure). The data from the medical imaging device 124 (FIG. 2) can be utilized to determine contextual information regarding the type of procedure being performed in a number of different ways, including by determining the

angle at which the medical imaging device is oriented with respect to the visualization of the patient's anatomy, monitoring the number or medical imaging devices being utilized (i.e., that are activated and paired with the surgical hub 5104), and monitoring the types of visualization devices utilized. For example, one technique for performing a VATS lobectomy places the camera in the lower anterior corner of the patient's chest cavity above the diaphragm, whereas one technique for performing a VATS segmentectomy places the camera in an anterior intercostal position relative to the segmental fissure. Using pattern recognition or machine learning techniques, for example, the situational awareness system can be trained to recognize the positioning of the medical imaging device according to the visualization of the patient's anatomy. As another example, one technique for performing a VATS lobectomy utilizes a single medical imaging device, whereas another technique for performing a VATS segmentectomy utilizes multiple cameras. As yet another example, one technique for performing a VATS segmentectomy utilizes an infrared light source (which can be communicably coupled to the surgical hub as part of the visualization system) to visualize the segmental fissure, which is not utilized in a VATS lobectomy. By tracking any or all of this data from the medical imaging device 5108, the surgical hub 5104 can thereby determine the specific type of surgical procedure being performed and/or the technique being used for a particular type of surgical procedure.

[0992] Ninth 5218, the surgical team begins the dissection step of the procedure. The surgical hub 5104 can infer that the surgeon is in the process of dissecting to mobilize the patient's lung because it receives data from the RF or ultrasonic generator indicating that an energy instrument is being fired. The surgical hub 5104 can cross-reference the received data with the retrieved steps of the surgical procedure to determine that an energy instrument being fired at this point in the process (i.e., after the completion of the previously discussed steps of the procedure) corresponds to the dissection step. Tenth 5220, the surgical team proceeds to the ligation step of the procedure. The surgical hub 5104 can infer that the surgeon is ligating arteries and veins because it receives data from the surgical stapling and cutting instrument indicating that the instrument is being fired. Similarly to the prior step, the surgical hub 5104 can derive this inference by cross-referencing the receipt of data from the surgical stapling and cutting instrument with the retrieved steps in the process. Eleventh 5222, the segmentectomy portion of the procedure is performed. The surgical hub 5104 can infer that the surgeon is transecting the parenchyma based on data from the surgical stapling and cutting instrument, including data from its cartridge. The cartridge data can correspond to the size or type of staple being fired by the instrument, for example. As different types of staples are utilized for different types of tissues, the cartridge data can thus indicate the type of tissue being stapled and/or transected. In this case, the type of staple being fired is utilized for parenchyma (or other similar tissue types), which allows the surgical hub 5104 to infer that the segmentectomy portion of the procedure is being performed. Twelfth 5224, the node dissection step is then performed. The surgical hub 5104 can infer that the surgical team is dissecting the node and performing a leak test based on data received from the generator indicating that an RF or ultrasonic instrument is being fired. For this particular procedure, an RF or ultrasonic instrument being utilized after paren-

chyma was transected corresponds to the node dissection step, which allows the surgical hub 5104 to make this inference. It should be noted that surgeons regularly switch back and forth between surgical stapling/cutting instruments and surgical energy (i.e., RF or ultrasonic) instruments depending upon the particular step in the procedure because different instruments are better adapted for particular tasks. Therefore, the particular sequence in which the stapling/cutting instruments and surgical energy instruments are used can indicate what step of the procedure the surgeon is performing. Upon completion of the twelfth step 5224, the incisions are closed up and the post-operative portion of the procedure begins.

[0993] Thirteenth 5226, the patient's anesthesia is reversed. The surgical hub 5104 can infer that the patient is emerging from the anesthesia based on the ventilator data (i.e., the patient's breathing rate begins increasing), for example. Lastly, the fourteenth step 5228 is that the medical personnel remove the various patient monitoring devices 5124 from the patient. The surgical hub 5104 can thus infer that the patient is being transferred to a recovery room when the hub loses EKG, BP, and other data from the patient monitoring devices 5124. As can be seen from the description of this illustrative procedure, the surgical hub 5104 can determine or infer when each step of a given surgical procedure is taking place according to data received from the various data sources 5126 that are communicably coupled to the surgical hub 5104.

[0994] In addition to utilizing the patient data from EMR database(s) to infer the type of surgical procedure that is to be performed, as illustrated in the first step 5202 of the timeline 5200 depicted in FIG. 86, the patient data can also be utilized by a situationally aware surgical hub 5104 to generate control adjustments for the paired modular devices 5102. FIG. 87A illustrates a flow diagram depicting the process 5240 of importing patient data stored in an EMR database 5250 and deriving inferences 5256 therefrom, in accordance with at least one aspect of the present disclosure. Further, FIG. 87B illustrates a flow diagram depicting the process 5242 of determining control adjustments 5264 corresponding to the derived inferences 5256 from FIG. 87A, in accordance with at least one aspect of the present disclosure. In the following description of the processes 5240, 5242, reference should also be made to FIG. 81.

[0995] As shown in FIG. 87A, the surgical hub 5104 retrieves the patient information (e.g., EMR) stored in a database 5250 to which the surgical hub 5104 is communicably connected. The unredacted portion of the patient data is removed 5252 from the surgical hub 5104, leaving anonymized, stripped patient data 5254 related to the patient's condition and/or the surgical procedure to be performed. The unredacted patient data is removed 5252 in order to maintain patient anonymity for the processing of the data (including if the data is uploaded to the cloud for processing and/or data tracking for reports). The stripped patient data 5254 can include any medical conditions that the patient is suffering from, the patient's medical history (including previous treatments or procedures), medication that the patient is taking, and other such medically relevant details. The control circuit of the surgical hub 5104 can then derive various inferences 5256 from the stripped patient data 5254, which can in turn be utilized by the surgical hub 5104 to derive various control adjustments for the paired modular devices 5102. The derived inferences 5256 can be based upon individual pieces



of data or combinations of pieces of data. Further, the derived inferences **5256** may, in some cases, be redundant with each other as some data may lead to the same inference. By integrating each patient's stripped patient data **5254** into the situational awareness system, the surgical hub **5104** is thus able to generate pre-procedure adjustments to optimally control each of the modular devices **5102** based on the unique circumstances associated with each individual patient. In the illustrated example, the stripped patient data **5254** includes that (i) the patient is suffering from emphysema, (ii) has high blood pressure, (iii) is suffering from a small cell lung cancer, (iv) is taking warfarin (or another blood thinner), and/or (v) has received radiation pretreatment. In the illustrated example, the inferences **5256** derived from the stripped patient data **5254** include that (i) the lung tissue will be more fragile than normal lung tissue, (ii) hemostasis issues are more likely, (iii) the patient is suffering from a relatively aggressive cancer, (iv) hemostasis issues are more likely, and (v) the lung tissue will be stiffer and more prone to fracture, respectively.

[0996] After the control circuit of the surgical hub **5104** receives or identifies the implications **5256** that are derived from anonymized patient data, the control circuit of the surgical hub **5104** is configured to execute a process **5242** to control the modular devices **5102** in a manner consistent with the derived implications **5256**. In the example shown in FIG. **87B**, the control circuit of the surgical hub **5104** interprets how the derived implications **5256** impacts the modular devices **5102** and then communicates corresponding control adjustments to each of the modular devices **5102**. In the example shown in FIG. **87B**, the control adjustments include (i) adjusting the compression rate threshold parameter of the surgical stapling and cutting instrument, (ii) adjusting the visualization threshold value of the surgical hub **5104** to quantify bleeding via the visualization system **108** (FIG. **2**) (this adjustment can apply to the visualization system **108** itself or as an internal parameter of the surgical hub **5104**), (iii) adjusts the power and control algorithms of the combo generator module **140** (FIG. **3**) for the lung tissue and vessel tissue types, (iv) adjusts the margin ranges of the medical imaging device **124** (FIG. **2**) to account for the aggressive cancer type, (v) notifies the surgical stapling and cutting instrument of the margin parameter adjustment needed (the margin parameter corresponds to the distance or amount of tissue around the cancer that will be excised), and (vi) notifies the surgical stapling and cutting instrument that the tissue is potentially fragile, which causes the control algorithm of the surgical stapling and cutting instrument to adjust accordingly. Furthermore, the data regarding the implications **5256** derived from the anonymized patient data **5254** is considered by the situational awareness system to infer contextual information **5260** regarding the surgical procedure being performed. In the example shown in FIG. **87B**, the situational awareness system further infers that the procedure is a thoracic lung resection **5262**, e.g., segmentectomy.

[0997] Determining where inefficiencies or ineffectiveness may reside in a medical facility's practice can be challenging because medical personnel's efficiency in completing a surgical procedure, correlating positive patient outcomes with particular surgical teams or particular techniques in performing a type of surgical procedure, and other performance measures are not easily quantified using legacy systems. As one solution, the surgical hubs can be employed

to track and store data pertaining to the surgical procedures that the surgical hubs are being utilized in connection with and generate reports or recommendations related to the tracked data. The tracked data can include, for example, the length of time spent during a particular procedure, the length of time spent on a particular step of a particular procedure, the length of downtime between procedures, modular device(s) (e.g., surgical instruments) utilized during the course of a procedure, and the number and type of surgical items consumed during a procedure (or step thereof). Further, the tracked data can include, for example, the operating theater in which the surgical hub is located, the medical personnel associated with the particular event (e.g., the surgeon or surgical team performing the surgical procedure), the day and time at which the particular event(s) occurred, and patient outcomes. This data can be utilized to create performance metrics, which can be utilized to detect and then ultimately address inefficiencies or ineffectiveness within a medical facility's practice. In one exemplification, the surgical hub includes a situational awareness system, as described above, that is configured to infer or determine information regarding a particular event (e.g., when a particular step of a surgical procedure is being performed and/or how long the step took to complete) based on data received from data sources connected to the surgical hub (e.g., paired modular devices). The surgical hub can then store this tracked data to provide reports or recommendations to users.

#### Aggregation and Reporting of Surgical Hub Data

[0998] FIG. **88** illustrates a block diagram of a computer-implemented interactive surgical system **5700**, in accordance with at least one aspect of the present disclosure. The system **5700** includes a number of surgical hubs **5706** that, as described above, are able to detect and track data related to surgical procedures that the surgical hubs **5706** (and the modular devices paired to the surgical hubs **5706**) are utilized in connection with. In one exemplification, the surgical hubs **5706** are connected to form local networks such that the data being tracked by the surgical hubs **5706** is aggregated together across the network. The networks of surgical hubs **5706** can be associated with a medical facility, for example. The data aggregated from the network of surgical hubs **5706** can be analyzed to provide reports on data trends or recommendations. For example, the surgical hubs **5706** of a first medical facility **5704a** are communicably connected to a first local database **5708a** and the surgical hubs **5706** of a second medical facility **5704b** are communicably connected to a second local database **5708b**. The network of surgical hubs **5706** associated with the first medical facility **5704a** can be distinct from the network of surgical hubs **5706** associated with the second medical facility **5704b**, such that the aggregated data from each network of surgical hubs **5706** corresponds to each medical facility **5704a**, **5704b** individually. A surgical hub **5706** or another computer terminal communicably connected to the database **5708a**, **5708b** can be configured to provide reports or recommendations based on the aggregated data associated with the respective medical facility **5704a**, **5704b**. In this exemplification, the data tracked by the surgical hubs **5706** can be utilized to, for example, report whether a particular incidence of a surgical procedure deviated from the average in-network time to complete the particular procedure type. [0999] In another exemplification, each surgical hub **5706** is configured to upload the tracked data to the cloud **5702**,

which then processes and aggregates the tracked data across multiple surgical hubs 5706, networks of surgical hubs 5706, and/or medical facilities 5704a, 5704b that are connected to the cloud 5702. Each surgical hub 5706 can then be utilized to provide reports or recommendations based on the aggregated data. In this exemplification, the data tracked by the surgical hubs 5706 can be utilized to, for example, report whether a particular incidence of a surgical procedure deviated from the average global time to complete the particular procedure type.

[1000] In another exemplification, each surgical hub 5706 can further be configured to access the cloud 5702 to compare locally tracked data to global data aggregated from all of the surgical hubs 5706 that are communicably connected to the cloud 5702. Each surgical hub 5706 can be configured to provide reports or recommendations based on the comparison between the tracked local data relative to local (i.e., in-network) or global norms. In this exemplification, the data tracked by the surgical hubs 5706 can be utilized to, for example, report whether a particular incidence of a surgical procedure deviated from either the average in-network time or the average global time to complete the particular procedure type.

[1001] In one exemplification, each surgical hub 5706 or another computer system local to the surgical hub 5706 is configured to locally aggregate the data tracked by the surgical hubs 5706, store the tracked data, and generate reports and/or recommendations according to the tracked data in response to queries. In cases where the surgical hub 5706 is connected to a medical facility network (which may include additional surgical hubs 5706), the surgical hub 5706 can be configured to compare the tracked data with the bulk medical facility data. The bulk medical facility data can include EMR data and aggregated data from the local network of surgical hubs 5706. In another exemplification, the cloud 5702 is configured to aggregate the data tracked by the surgical hubs 5706, store the tracked data, and generate reports and/or recommendations according to the tracked data in response to queries.

[1002] Each surgical hub 5706 can provide reports regarding trends in the data and/or provide recommendations on improving the efficiency or effectiveness of the surgical procedures being performed. In various exemplifications, the data trends and recommendations can be based on data tracked by the surgical hub 5706 itself, data tracked across a local medical facility network containing multiple surgical hubs 5706, or data tracked across a number of surgical hubs 5706 communicably connected to a cloud 5702. The recommendations provided by the surgical hub 5706 can describe, for example, particular surgical instruments or product mixes to utilize for particular surgical procedures based on correlations between the surgical instruments/product mixes and patient outcomes and procedural efficiency. The reports provided by the surgical hub 5706 can describe, for example, whether a particular surgical procedure was performed efficiently relative to local or global norms, whether a particular type of surgical procedure being performed at the medical facility is being performed efficiently relative to global norms, and the average time taken to complete a particular surgical procedure or step of a surgical procedure for a particular surgical team.

[1003] In one exemplification, each surgical hub 5706 is configured to determine when operating theater events occur (e.g., via a situational awareness system) and then track the

length of time spent on each event. An operating theater event is an event that a surgical hub 5706 can detect or infer the occurrence of. An operating theater event can include, for example, a particular surgical procedure, a step or portion of a surgical procedure, or downtime between surgical procedures. The operating theater events can be categorized according to an event type, such as a type of surgical procedure being performed, so that the data from individual procedures can be aggregated together to form searchable data sets. FIG. 90 illustrates an example of a diagram 5400 depicting the data tracked by the surgical hubs 5706 being parsed to provide increasingly detailed metrics related to surgical procedures or the use of the surgical hub 5706 (as depicted further in FIGS. 91-95) for an illustrative data set. In one exemplification, the surgical hub 5706 is configured to determine whether a surgical procedure is being performed and then track both the length of time spent between procedures (i.e., downtime) and the time spent on the procedures themselves. The surgical hub 5706 can further be configured to determine and track the time spent on each of the individual steps taken by the medical personnel (e.g., surgeons, nurses, orderlies) either between or during the surgical procedures. The surgical hub can determine when surgical procedures or different steps of surgical procedures are being performed via a situational awareness system, which is described in further detail above.

[1004] FIG. 89 illustrates a logic flow diagram of a process 5300 for tracking data associated with an operating theater event. In the following description, description of the process 5300, reference should also be made to FIG. 88. In one exemplification, the process 5300 can be executed by a control circuit of a surgical hub 206, as depicted in FIG. 10 (processor 244). In yet another exemplification, the process 5300 can be executed by a distributed computing system including a control circuit of a surgical hub 206 in combination with a control circuit of a modular device, such as the microcontroller 461 of the surgical instrument depicted FIG. 12, the microcontroller 620 of the surgical instrument depicted in FIG. 16, the control circuit 710 of the robotic surgical instrument 700 depicted in FIG. 17, the control circuit 760 of the surgical instruments 750, 790 depicted in FIGS. 18 and 19, or the controller 838 of the generator 800 depicted in FIG. 20. For economy, the following description of the process 5300 will be described as being executed by the control circuit of a surgical hub 5706; however, it should be understood that the description of the process 5300 encompasses all of the aforementioned exemplifications.

[1005] The control circuit of the surgical hub 5706 executing the process 5300 receives 5302 perioperative data from the modular devices and other data sources (e.g., databases and patient monitoring devices) that are communicably coupled to the surgical hub 5706. The control circuit then determines 5304 whether an event has occurred via, for example, a situational awareness system that derives contextual information from the received 5302 data. The event can be associated with an operating theater in which the surgical hub 5706 is being used. The event can include, for example, a surgical procedure, a step or portion of a surgical procedure, or downtime between surgical procedures or steps of a surgical procedure. Furthermore, the control circuit tracks data associated with the particular event, such as the length of time of the event, the surgical instruments and/or other medical products utilized during the course of the event, and the medical personnel associated with the

event. The surgical hub **5706** can further determine this information regarding the event via, for example, the situational awareness system.

**[1006]** For example, the control circuit of a situationally aware surgical hub **5706** could determine that anesthesia is being induced in a patient through data received from one or more modular devices **5102** (FIG. **81**) and/or patient monitoring devices **5124** (FIG. **81**). The control circuit could then determine that the operative portion of the surgical procedure has begun upon detecting that an ultrasonic surgical instrument or RF electrosurgical instrument has been activated. The control circuit could thus determine the length of time for the anesthesia inducement step according to the difference in time between the beginning of that particular step and the beginning of the first step in the operative portion of the surgical procedure. Likewise, the control circuit could determine how long the particular operative step in the surgical procedure took according to when the control circuit detects the subsequent step in the procedure begins. Further, the control circuit could determine how long the overall operative portion of the surgical procedure took according to when the control circuit detects that the final operative step in the procedure ends. The control circuit can also determine what surgical instruments (and other modular devices **5102**) are being utilized during the course of each step in the surgical procedure by tracking the activation and/or use of the instruments during each of the steps. The control circuit can also detect the completion of the surgical procedure by, for example, detecting when the patient monitoring devices **5124** have been removed from the patient (as in step fourteen **5228** of FIG. **86**). The control circuit can then track the downtime between procedures according to when the control circuit infers that the subsequent surgical procedure has begun.

**[1007]** The control circuit executing the process **5300** then aggregates **5306** the data associated with the event according to the event type. In one exemplification, the aggregated **5306** data can be stored in a memory **249** (FIG. **10**) of the surgical hub **5706**. In another exemplification, the control circuit is configured to upload the data associated with the event to the cloud **5702**, whereupon the data is aggregated **5306** according to the event type for all of the data uploaded by each of the surgical hubs **5706** connected to the cloud **5702**. In yet another exemplification, the control circuit is configured to upload the data associated with the event to a database associated with a local network of the surgical hubs **5706**, whereupon the data is aggregated **5306** according to the event type for all of the data uploaded across the local network of surgical hubs **5706**.

**[1008]** In one exemplification, the control circuit is further configured to compare the data associated with the event type to baseline data associated with the event type. The baseline data can correspond to, for example, average values associated with the particular event type for a particular hospital, network of hospitals, or across the entirety of the cloud **5702**. The baseline data can be stored on the surgical hub **5706** or retrieved by the surgical **5706** as the perioperative data is received **5302** thereby.

**[1009]** Aggregating **5306** the data from each of the events according to the event type allows individual incidents of the event type to thereafter be compared against the historical or aggregated data to determine when deviations from the norm for an event type occur. The control circuit further determines **5308** whether it has received a query. If the control

circuit does not receive a query, then the process **5300** continues along the NO branch and loops back to continue receiving **5302** data from the data sources. If the control circuit does receive a query for a particular event type, the process **5300** continues along the YES branch and the control circuit then retrieves the aggregated data for the particular event type and displays **5310** the appropriate aggregated data corresponding to the query. In various exemplifications, the control circuit can retrieve the appropriate aggregated data from the memory of the surgical hub **5706**, the cloud **5702**, or a local database **5708a**, **5708b**.

**[1010]** In one example, the surgical hub **5706** is configured to determine a length of time for a particular procedure via the aforementioned situational awareness system according to data received from one or more modular devices utilized in the performance of the surgical procedure (and other data sources). Each time a surgical procedure is completed, the surgical hub **5706** uploads or stores the length of time required to complete the particular type of surgical procedure, which is then aggregated with the data from every other instance of the type of procedure. In some aspects, the surgical hub **5706**, cloud **5702**, and/or local database **5708a**, **5708b** can then determine an average or expected procedure length for the particular type of procedure from the aggregated data. When the surgical hub **5706** receives a query as to the particular type of procedure thereafter, the surgical hub **5706** can then provide feedback as to the average (or expected) procedure length or compare an individual incidence of the procedure type to the average procedure length to determine whether the particular incidence deviates therefrom.

**[1011]** In some aspects, the surgical hub **5706** can be configured to automatically compare each incidence of an event type to average or expected norms for the event type and then provide feedback (e.g., display a report) when a particular incidence of the event type deviates from the norm. For example, the surgical hub **5706** can be configured to provide feedback whenever a surgical procedure (or a step of the surgical procedure) deviates from the expected length of time to complete the surgical procedure (or the step of the surgical procedure) by more than a set amount.

**[1012]** Referring back to FIG. **90**, the surgical hub **5706** could be configured to track, store, and display data regarding the number of patients operated on (or procedures completed) per day per operating theater (bar graph **5402** depicted further in FIG. **91**), for example. The surgical hub **5706** could be configured to further parse the number of patients operated on (or procedures completed) per day per operating theater and can be further parsed according to the downtime between the procedures on a given day (bar graph **5404** depicted further in FIG. **92**) or the average procedure length on a given day (bar graph **5408** depicted further in FIG. **94**). The surgical hub **5706** can be further configured to provide a detailed breakdown of the downtime between procedures according to, for example, the number and length of the downtime time periods and the subcategories of the actions or steps during each time period (bar graph **5406** depicted further in FIG. **93**). The surgical hub **5706** can be further configured to provide a detailed breakdown of the average procedure length on a given day according to each individual procedure and the subcategory of actions or steps during each procedure (bar graph **5410** depicted further in FIG. **95**). The various graphs shown in FIGS. **90-95** can represent data tracked by the surgical hub **5706** and can

further be generated automatically or displayed by the surgical hub 5706 in response to queries submitted by users.

[1013] FIG. 91 illustrates an example bar graph 5402 depicting the number of patients 5420 operated on relative to the days of the week 5422 for different operating rooms 5424, 5426. The surgical hub 5706 can be configured to provide (e.g., via a display) the number of patients 5420 operated on or procedures that are completed in connection with each surgical hub 5706, which can be tracked through a situational awareness system or accessing the hospital's EMR database, for example. In one exemplification, the surgical hub 5706 can further be configured to collate this data from different surgical hubs 5706 within the medical facility that are communicably connected together, which allows each individual surgical hub to present the aggregated data of the medical facility on a hub-by-hub or operating theater-by-theater basis. In one exemplification, the surgical hub 5706 can be configured to compare one or more tracked metrics to a threshold value (which may be unique to each tracked metric). When at least one of the tracked metrics exceeds the threshold value (i.e., either increases above or drops below the threshold value, as appropriate for the particular tracked metric), then the surgical hub 5706 provides a visual, audible, or tactile alert to notify a user of such. For example, the surgical hub 5706 can be configured to indicate when the number of patients or procedures deviates from an expected, average, or threshold value. For example, FIG. 91 depicts the number of patients on Tuesday 5428 and Thursday 5430 for a first operating theater 5424 as being highlighted for being below expectation. Conversely, no days are highlighted for a second operating theater 5426 for this particular week, which means in this context that the number of patients for each day falls within expectations.

[1014] FIG. 92 illustrates a bar graph 5404 depicting the total downtime between procedures 5432 relative to the days of a week 5434 for a particular operating room. The surgical hub 5706 can be configured to track the length of downtime between surgical procedures through a situational awareness system, for example. The situational awareness system can detect or infer when each particular downtime instance is occurring and then track the length of time for each instance of downtime. The surgical hub 5706 can thereby determine the total downtime 5432 for each day of the week 5434 by summing the downtime instances for each particular day. In one exemplification, the surgical hub 5706 can be configured to provide an alert when the total length of downtime on a given day (or another unit of time) deviates from an expected, average, or threshold value. For example, FIG. 92 depicts the total downtime 5432 on Tuesday 5436 and Friday 5438 as being highlighted for deviating from an expected length of time.

[1015] FIG. 93 illustrates a bar graph 5406 depicting the total downtime 5432 per day of the week 5434 as depicted in FIG. 92 broken down according to each individual downtime instance. The number of downtime instances and the length of time for each downtime instance can be represented within each day's total downtime. For example, on Tuesday in the first operating theater (OR1) there were four instances of downtime between procedures and the magnitude of the first downtime instance indicates that it was longer than the other three instances. In one exemplification, the surgical hub 5706 is configured to further indicate the particular actions or steps taken during a

selected downtime instance. For example, in FIG. 93, Thursday's second downtime instance 5440 has been selected, which then causes a callout 5442 to be displayed indicating that this particular downtime instance consisted of performing the initial set-up of the operating theater, administering anesthesia, and prepping the patient. As with the downtime instances themselves, the relative size or length of the actions or steps within the callout 5442 can correspond to the length of time for each particular action or step. The detail views for the downtime instances can be displayed when a user selects the particular instance, for example.

[1016] FIG. 94 illustrates a bar graph 5408 depicting the average procedure length 5444 relative to the days of a week 5446 for a particular operating theater. The surgical hub 5706 can be configured to track the average procedure length through a situational awareness system, for example. The situational awareness system can detect or infer when each particular step of a surgical procedure is occurring (see FIG. 86, for example) and then track the length of time for each of the steps. The surgical hub 5706 can thereby determine the total downtime 5432 for each day of the week 5434 by summing the lengths of the downtime instances for the particular day. In one exemplification, the surgical hub 5706 can be configured to indicate when the average procedure length deviates from an expected value. For example, FIG. 94 depicts Thursday's average procedure length 5448 for the first operating room (OR1) as being highlighted for deviating from an expected length of time.

[1017] FIG. 95 illustrates a bar graph 5410 depicting the procedure lengths 5450 relative to procedure types 5452. The depicted procedure lengths 5450 can either represent the average procedure lengths for particular types of procedures or the procedure lengths for each individual procedure performed on a given day in a given operating theater. The procedure lengths 5450 for different procedure types 5452 can then be compared. Further, the average lengths for the steps in a procedure type 5452 or the length for each particular step in a particular procedure can be displayed when a procedure is selected. Further, the procedure types 5452 can be tagged with various identifiers for parsing and comparing different data sets. For example, in FIG. 95 the first procedure 5454 corresponds to a colorectal procedure (specifically, a low anterior resection) where there was a preoperative identification of abdominal adhesions. The second procedure 5456 corresponds to a thoracic procedure (specifically, a segmentectomy). It should be noted again that the procedures depicted in FIG. 95 can represent the lengths of time for individual procedures or the average lengths of time for all of the procedures for the given procedure types. Each of the procedures can further be broken down according to the length of time for each step in the procedure. For example, FIG. 95 depicts the second procedure 5456 (a thoracic segmentectomy) as including an icon or graphical representation 5458 of the length of time spent on the dissect vessels, ligate (the vessels), nodal dissection, and closing steps of the surgical procedure. As with the procedure lengths themselves, the relative size or length of the steps within the graphical representation 5442 can correspond to the length of time for each particular step of the surgical procedure. The detail views for the steps of the surgical procedures can be displayed when a user selects the particular procedure, for example. In one exemplification, the surgical hub 5706 can be configured to identify when a length of time to complete a given step in the

procedure deviates from an expected length of time. For example, FIG. 95 depicts the nodal dissection step as being highlighted for deviating from an expected length of time.

[1018] In one exemplification, an analytics package of the surgical hub 5706 can be configured to provide the user with usage data and results correlations related to the surgical procedures (or downtime between procedures). For example, the surgical hub 5706 can be configured to display methods or suggestions to improve the efficiency or effectiveness of a surgical procedure. As another example, the surgical hub 5706 can be configured to display methods to improve cost allocation. FIGS. 96-101 depict examples of various metrics that can be tracked by the surgical hub 5706, which can then be utilized to provide medical facility personnel suggestions for inventory utilization or technique outcomes. For example, a surgical hub 5706 could provide a surgeon with a suggestion pertaining to a particular technique outcome prior to or at the beginning of a surgical procedure based on the metrics tracked by the surgical hub 5706.

[1019] FIG. 96 illustrates a bar graph 5460 depicting the average completion time 5462 for particular procedural steps 5464 for different types of thoracic procedures. The surgical hub 5706 can be configured to track and store historical data for different types of procedures and calculate the average time to complete the procedure (or an individual step thereof). For example, FIG. 96 depicts the average completion time 5462 for thoracic segmentectomy 5466, wedge 5468, and lobectomy 5470 procedures. For each type of procedure, the surgical hub 5706 can track the average time to complete each step thereof. In this particular example, the dissection, vessel transection, and node dissection steps are indicated for each type of procedure. In addition to tracking and providing the average time for the steps of the procedure types, the surgical hub 5706 can additionally track other metrics or historical data, such as the complication rate for each procedure type (i.e., the rate of procedures having at least one complication as defined by the surgical hub 5706 or the surgeon). Additional tracked metrics for each procedure type, such as the complication rate, can also be depicted for comparison between the different procedure types.

[1020] FIG. 97 illustrates a bar graph 5472 depicting the procedure time 5474 relative to procedure types 5476. The surgical hub 5706 can be configured to track and store historical data or metrics for different procedure types 5476 or classes, which can encompass multiple subtypes of procedures. For example, FIG. 97 depicts the procedure time 5474 for surgical procedures classified as a thoracic 5478, bariatric 5480, or colorectal 5482 procedure. In various exemplifications, the surgical hub 5706 can output the procedure time 5474 for the procedure classifications expressed in terms of either the total length of time or the average time spent on the given procedure types 5476. The analytics package of the surgical hub 5706 can, for example, provide this data to the surgeons, hospital officials, or medical personnel to track the efficiency of the queried procedures. For example, FIG. 97 depicts bariatric procedures 5480 as taking a lower average time (i.e., being more time efficient) than either thoracic procedures 5478 or colorectal procedures 5482.

[1021] FIG. 98 illustrates a bar graph 5484 depicting operating room downtime 5486 relative to the time of day 5488. Relatedly, FIG. 99 illustrates a bar graph 5494 depict-

ing operating room downtime 5496 relative to the day of the week 5498. Operating room downtime 5486, 5496 can be expressed in, for example, a length of a unit of time or relative utilization (i.e., percentage of time that the operating room is in use). The operating room downtime data can encompass an individual operating room or an aggregation of multiple operating rooms at a medical facility. As discussed above, a surgical hub 5706 can be configured to track whether a surgical procedure is being performed in the operating theater associated with the surgical hub 5706 (including the length of time that a surgical procedure is or is not being performed) utilizing a situational awareness system, for example. As shown in FIGS. 98 and 99, the surgical hub 5706 can provide an output (e.g., bar graphs 5484, 5494 or other graphical representations of data) depicting the tracked data pertaining to when the operating room is being utilized (i.e., when a surgical procedure is being performed) and/or when there is downtime between procedures. Such data can be utilized to identify ineffectiveness or inefficiencies in performing surgical procedures, cleaning or preparing operating theaters for surgery, scheduling, and other metrics associated with operating theater use. For example, FIG. 98 depicts a comparative increase in operating room downtime 5486 at a first instance 5490 from 11:00 a.m.-12:00 p.m. and a second instance 5492 from 3:00-4:00 p.m. As another example, FIG. 99 depicts a comparative increase in operating room downtime 5496 on Mondays 5500 and Fridays 5502. In various exemplifications, the surgical hub 5706 can provide operating theater downtime data for a particular instance (i.e., a specific time, day, week, etc.) or an average operating theater downtime data for a category of instances (i.e., aggregated data for a day, time, week, etc.). Hospital officials or other medical personnel thus could use this data to identify specific instances where an inefficiency may have occurred or identify trends in particular days and/or times of day where there may be inefficiencies. From such data, the hospital officials or other medical personnel could then investigate to identify the specific reasons for these increased downtimes and take corrective action to address the identified reason.

[1022] In various exemplifications, the surgical hub 5706 can be configured to display data in response to queries in a variety of different formats (e.g., bar graphs, pie graphs, infographics). FIG. 100 illustrates a pair of pie charts depicting the percentage of time that the operating theater is utilized. The operating theater utilization percentage can encompass an individual operating theater or an aggregation of multiple operating theaters (e.g., the operating rooms at a medical facility or every operating room for all medical facilities having surgical hubs 5706 connected to the cloud 5702). As discussed above, a surgical hub 5706 can be configured to determine when a surgical procedure is or is not being performed (i.e., whether the operating theater associated with the surgical hub 5706 is being utilized) using a situational awareness system, for example. In addition to expressing operating theater utilization in terms of an average or absolute amount for different time periods (as depicted in FIGS. 98-99), the surgical hub 5706 can additionally express operating theater utilization in terms of a percentage or relative amount compared to a maximum possible utilization. As above, the operating theater utilization can be parsed for particular time periods, including the overall utilization (i.e., the total historical percentage of time in use) for the particular operating theater (or groups of

operating theaters) or the utilization over the span of a particular time period. As shown in FIG. 100, a first pie chart 5504 depicts the overall operating theater utilization 5508 (85%) and a second pie chart 5506 depicts the operating theater utilization for the prior week 5510 (75%). Hospital officials and other medical personnel could use this data to identify that there may have been some inefficiency that occurred in the prior week that caused the particular operating theater (or group of operating theaters) to be utilized less efficiently compared to the historical average so that further investigations can be carried out to identify the specific reasons for this decreased utilization.

[1023] In some exemplifications, the surgical hub 5706 is configured to track detect and track the number of surgical items that are utilized during the course of a surgical procedure. This data can then be aggregated and displayed (either automatically or in response to a query) according to, for example, a particular time period (e.g., per day or per week) or for a particular surgical procedure type (e.g., thoracic procedures or abdominal procedures). FIG. 101 illustrates a bar graph 5512 depicting consumed and unused surgical items 5514 relative to procedure type 5516. The surgical hub 5706 can be configured to determine or infer what surgical items are being consumed during the course of each surgical procedure via a situational awareness system. The situational awareness system can determine or receive the list of surgical items to be used in a procedure (e.g., see FIG. 85B), determine or infer when each procedure (and steps thereof) begins and ends, and determine when a particular surgical item is being utilized according to the procedural step being performed. The inventory of surgical items that are consumed or unused during the course of a surgical procedure can be represented in terms of the total number of surgical items or the average number of surgical items per procedure type 5516, for example. The consumed surgical items can include non-reusable items that are utilized during the course of a surgical procedure. The unused surgical items can include additional items that are not utilized during the procedure(s) or scrap items. The procedure type can correspond to broad classifications of procedures or a specific procedure type or technique for performing a procedure type. For example, in FIG. 101 the procedure types 5516 being compared are thoracic, colorectal, and bariatric procedures. For each of these procedure types 5516, the average number of consumed and unused surgical items 5514 are both provided. In one aspect, the surgical hub 5706 can be configured to further parse the consumed and/or unused surgical items 5514 by the specific item type. In one exemplification, the surgical hub 5706 can provide a detailed breakdown of the surgical items 5514 making up each item category for each surgical procedure type 5516 and graphically represent the different categories of surgical items 5514. For example, in FIG. 101, the unused surgical items are depicted in dashed lines and the consumed surgical items are depicted in solid lines. In one exemplification, the surgical hub 5706 is configured to further indicate the specific within a category for a particular procedure type 5516. For example, in FIG. 93, the consumed items category for the thoracic procedure type has been selected, which then causes a callout 5520 to be displayed listing the particular surgical items in the category: stapler cartridges, sponges, saline, fibrin sealants, surgical sutures, and stapler buttress material. Furthermore, the callout 5520 can be configured to provide the quantities of the listed items in the category,

which may be the average or absolute quantities of the items (either consumed or unused) for the particular procedure type.

[1024] In one exemplification, the surgical hub 5706 can be configured to aggregate tracked data in a redacted format (i.e., with any patient-identifying information stripped out). Such bulk data can be utilized for academic or business analysis purposes. Further, the surgical hub 5706 can be configured to upload the redacted or anonymized data to a local database of the medical facility in which the surgical hub 5706 is located, an external database system, or the cloud 5702, whereupon the anonymized data can be accessed by user/client applications on demand. The anonymized data can be utilized to compare outcomes and efficiencies within a hospital or between geographic regions, for example.

[1025] The process 5300 depicted in FIG. 89 improves scheduling efficiency by allowing the surgical hubs 5706 to automatically store and provide granular detail on correlations between lengths of time required for various procedures according to particular days, particular types of procedures, particular hospital staff members, and other such metrics. This process 5300 also reduces surgical item waste by allowing the surgical hubs 5706 to provide alerts when the amount of surgical items being consumed, either on a per-procedure basis or as a category, are deviating from the expected amounts. Such alerts can be provided either automatically or in response to receiving a query.

[1026] FIG. 102 illustrates a logic flow diagram of a process 5350 for storing data from the modular devices and patient information database for comparison. In the following description, description of the process 5350, reference should also be made to FIG. 88. In one exemplification, the process 5350 can be executed by a control circuit of a surgical hub 206, as depicted in FIG. 10 (processor 244). In yet another exemplification, the process 5350 can be executed by a distributed computing system including a control circuit of a surgical hub 206 in combination with a control circuit of a modular device, such as the microcontroller 461 of the surgical instrument depicted FIG. 12, the microcontroller 620 of the surgical instrument depicted in FIG. 16, the control circuit 710 of the robotic surgical instrument 700 depicted in FIG. 17, the control circuit 760 of the surgical instruments 750, 790 depicted in FIGS. 18 and 19, or the controller 838 of the generator 800 depicted in FIG. 20. For economy, the following description of the process 5350 will be described as being executed by the control circuit of a surgical hub 5706; however, it should be understood that the description of the process 5350 encompasses all of the aforementioned exemplifications.

[1027] The control circuit executing the process 5350 receives data from the data sources, such as the modular device(s) and the patient information database(s) (e.g., EMR databases) that are communicably coupled to the surgical hub 5706. The data from the modular devices can include, for example, usage data (e.g., data pertaining to how often the modular device has been utilized, what procedures the modular device has been utilized in connection with, and who utilized the modular devices) and performance data (e.g., data pertaining to the internal state of the modular device and the tissue being operated on). The data from the patient information databases can include, for example, patient data (e.g., data pertaining to the patient's age, sex, and medical history) and patient outcome data (e.g., data

pertaining to the outcomes from the surgical procedure). In some exemplifications, the control circuit can continuously receive 5352 data from the data sources before, during, or after a surgical procedure.

[1028] As the data is received 5352, the control circuit aggregates 5354 the data in comparison groups of types of data. In other words, the control circuit causes a first type of data to be stored in association with a second type of data. However, more than two different types of data can be aggregated 5354 together into a comparison group. For example, the control circuit could store a particular type of performance data for a particular type of modular device (e.g., the force to fire for a surgical cutting and stapling instrument or the characterization of the energy expended by an RF or ultrasonic surgical instrument) in association with patient data, such as sex, age (or age range), a condition (e.g., emphysema) associated with the patient. In one exemplification, when the data is aggregated 5354 into comparison groups, the data is anonymized such that all patient-identifying information is removed from the data. This allows the data aggregated 5354 into comparison groups to be utilized for studies, without compromising confidential patient information. The various types of data can be aggregated 5354 and stored in association with each other in lookup tables, arrays, and other such formats. In one exemplification, the received 5352 data is automatically aggregated 5354 into comparison groups. Automatically aggregating 5354 and storing the data allows the surgical hub 5706 to quickly return results for queries and the groups of data to be exported for analysis according to specifically desired data types.

[1029] When the control circuit receives 5356 a query for a comparison between two or more of the tracked data types, the process 5350 proceeds along the YES branch. The control circuit then retrieves the particular combination of the data types stored in association with each other and then displays 5358 a comparison (e.g., a graph or other graphical representation of the data) between the subject data types. If the control circuit does not receive 5356 a query, the process 5350 continues along the NO branch and the control circuit continues receiving 5352 data from the data sources.

[1030] In one exemplification, the control circuit can be configured to automatically quantify a correlation between the received 5352 data types. In such aspects, the control circuit can calculate a correlation coefficient (e.g., the Pearson's coefficient) between pairs of data types. In one aspect, the control circuit can be configured to automatically display a report providing suggestions or other feedback if the quantified correlation exceeds a particular threshold value. In one aspect, the control circuit of the surgical hub 5706 can be configured to display a report on quantified correlations exceeding a particular threshold value upon receiving a query or request from a user.

[1031] In one exemplification, a surgical hub 5706 can compile information on procedures that the surgical hub 5706 was utilized in the performance of, communicate with other surgical hubs 5706 within its network (e.g., a local network of a medical facility or a number of surgical hubs 5706 connected by the cloud 5702), and compare results between type of surgical procedures or particular operating theaters, doctors, or departments. Each surgical hub 5706 can calculate and analyze utilization, efficiency, and comparative results (relative to all surgical hubs 5706 across a hospital network, a region, etc.). For example, the surgical

hub 5706 can display efficiency and comparative data, including operating theater downtime, operating theater clean-up and recycle time, step-by-step completion timing for procedures (including highlighting which procedural steps take the longest, for example), average times for surgeons to complete procedures (including parsing the completion times on a procedure-by-procedure basis), historical completion times (e.g., for completing classes of procedures, specific procedures, or specific steps within a procedure), and/or operating theater utilization efficiency (i.e., the time efficiency from a procedure to a subsequent procedure). The data that is accessed and shared across networks by the surgical hubs 5706 can include the anonymized data aggregated into comparison groups, as discussed above.

[1032] For example, the surgical hub 5706 can be utilized to perform studies of performance by instrument type or cartridge type for various procedures. As another example, the surgical hub 5706 can be utilized to perform studies on the performance of individual surgeons. As yet another example, the surgical hub 5706 can be utilized to perform studies on the effectiveness of different surgical procedures according to patients' characteristics or disease states.

[1033] In another exemplification, a surgical hub 5706 can provide suggestions on streamlining processes based on tracked data. For example, the surgical hub 5706 can suggest different product mixes according to the length of certain procedures or steps within a procedure (e.g., suggest a particular item that is more appropriate for long procedure steps), suggest more cost effective product mixes based on the utilization of items, and/or suggest kitting or pre-grouping certain items to lower set-up time. In another exemplification, a surgical hub 5706 can compare operating theater utilization across different surgical groups in order to better balance high volume surgical groups with surgical groups that have more flexible bandwidth. In yet another aspect, the surgical hub 5706 could be put in a forecasting mode that would allow the surgical hub 5706 to monitor upcoming procedure preparation and scheduling, then notify the administration or department of upcoming bottlenecks or allow them to plan for scalable staffing. The forecasting mode can be based on, for example, the anticipated future steps of the current surgical procedure that is being performed using the surgical hub 5706, which can be determined by a situational awareness system.

[1034] In another exemplification, a surgical hub 5706 can be utilized as a training tool to allow users to compare their procedure timing to other types of individuals or specific individuals within their department (e.g., a resident could compare his or her timing to a particular specialist or the average time for a specialist within the hospital) or the department average times. For example, users could identify what steps of a surgical procedure they are spending an inordinate amount of time on and, thus, what steps of the surgical procedure that they need to improve upon.

[1035] In one exemplification, all processing of stored data is performed locally on each surgical hub 5706. In another exemplification, each surgical hub 5706 is part of a distributed computing network, wherein each individual surgical hub 5706 compiles and analyzes its stored data and then communicates the data to the requesting surgical hub 5706. A distributed computing network could permit fast parallel processing. In another exemplification, each surgical hub 5706 is communicably connected to a cloud 5702,

which can be configured to receive the data from each surgical hub 5706 and then perform the necessary processing (data aggregation, calculations, and so on) on the data.

[1036] The process 5350 depicted in FIG. 102 improves the ability to determine when procedures are being performed inefficiently by allowing the surgical hubs 5706 to provide alerts when particular procedures, either on a per-procedure basis or as category, are deviating from the expected times to complete the procedures. Such alerts can be provided either automatically or in response to receiving a query. This process 5350 also improves the ability to perform studies on what surgical instruments and surgical procedure techniques provide the best patient outcomes by automatically tracking and indexing such data in easily-retrievable and reportable formats.

[1037] Some systems described herein offload the data processing that controls the modular devices (e.g., surgical instruments) from the modular devices themselves to an external computing system (e.g., a surgical hub) and/or a cloud. However in some exemplifications, some modular devices can sample data (e.g., from the sensors of the surgical instruments) at a faster rate than the rate at which the data can be transmitted to and processed by a surgical hub. As one solution, the surgical hub and the surgical instruments (or other modular devices) can utilize a distributed computing system where at least a portion of the data processing is performed locally on the surgical instrument. This can avoid data or communication bottlenecks between the instrument and the surgical hub by allowing the onboard processor of the surgical instrument to handle at least some of the data processing when the data sampling rate is exceeding the rate at which the data can be transmitted to the surgical hub. In some exemplifications, the distributed computing system can cease distributing the processing between the surgical hub and the surgical instrument and instead have the processing be executed solely onboard the surgical instrument. The processing can be executed solely by the surgical instrument in situations where, for example, the surgical hub needs to allocate its processing capabilities to other tasks or the surgical instrument is sampling data at a very high rate and it has the capabilities to execute all of the data processing itself.

[1038] Similarly, the data processing for controlling the modular devices, such as surgical instruments, can be taxing for an individual surgical hub to perform. If the surgical hub's processing of the control algorithms for the modular devices cannot keep pace with the use of the modular devices, then the modular devices will not perform adequately because their control algorithms will either not be updated as needed or the updates to the control algorithms will lag behind the actual use of the instrument. As one solution, the surgical hubs can be configured to utilize a distributed computing system where at least a portion of the processing is performed across multiple separate surgical hubs. This can avoid data or communication bottlenecks between the modular devices and the surgical hub by allowing each surgical hub to utilize the networked processing power of multiple surgical hubs, which can increase the rate at which the data is processed and thus the rate at which the control algorithm adjustments can be transmitted by the surgical hub to the paired modular devices. In addition to distributing the computing associated with controlling the various modular devices connected to the surgical hubs, a distributed computing system can also dynamically shift

computing resources between multiple surgical hubs in order to analyze tracked data in response to queries from users and perform other such functions. The distributed computing system for the surgical hubs can further be configured to dynamically shift data processing resources between the surgical hubs when any particular surgical hub becomes overtaxed.

[1039] The modular devices that are communicably connectable to the surgical hub can include sensors, memories, and processors that are coupled to the memories and configured to receive and analyze data sensed by the sensors. The surgical hub can further include a processor coupled to a memory that is configured to receive (through the connection between the modular device and the surgical hub) and analyze the data sensed by the sensors of the modular device. In one exemplification, the data sensed by the modular device is processed externally to the modular device (e.g., external to a handle assembly of a surgical instrument) by a computer that is communicably coupled to the modular device. For example, the advanced energy algorithms for controlling the operation of a surgical instrument can be processed by an external computing system, rather than on a controller embedded in the surgical instrument (such as instrument using an Advanced RISC Machine (ARM) processor). The external computer system processing the data sensed by the modular devices can include the surgical hub to which the modular devices are paired and/or a cloud computing system. In one exemplification, data sampled at a particular rate (e.g., 20 Ms/sec) and a particular resolution (e.g., 12 bits resolution) by a surgical instrument is decimated and then transmitted over a link to the surgical hub to which the surgical instrument is paired. Based on this received data, the control circuit of the surgical hub then determines the appropriate control adjustments for the surgical instrument, such as controlling power for an ultrasonic surgical instrument or RF electrosurgical instrument, setting motor termination points for a motor-driven surgical instrument, and so on. The control adjustments are then transmitted to the surgical instrument for application thereon.

#### Distributed Processing

[1040] FIG. 103 illustrates a diagram of a distributed computing system 5600. The distributed computing system 5600 includes a set of nodes 5602a, 5602b, 5602c that are communicably coupled by a distributed multi-party communication protocol such that they execute a shared or distributed computer program by passing messages therebetween. Although three nodes 5602a, 5602b, 5602c are depicted, the distributed computing system 5600 can include any number of nodes 5602a, 5602b, 5602c that are communicably connected together. Each of the nodes 5602a, 5602b, 5602c comprises a respective memory 5606a, 5606b, 5606c and processor 5604a, 5604b, 5604c coupled thereto. The processors 5604a, 5604b, 5604c execute the distributed multi-party communication protocol, which is stored at least partially in the memories 5606a, 5606b, 5606c. Each node 5602a, 5602b, 5602c can represent either a modular device or a surgical hub. Therefore, the depicted diagram represents aspects wherein various combinations of surgical hubs and/or modular devices are communicably coupled. In various exemplifications, the distributed computing system 5600 can be configured to distribute the computing associated with controlling the modular device(s) (e.g., advanced energy algorithms) over the modular device(s) and/or the surgical



hub(s) to which the modular device(s) are connected. In other words, the distributed computing system **5600** embodies a distributed control system for controlling the modular device(s) and/or surgical hub(s).

[1041] In some exemplifications, the modular device(s) and surgical hub(s) utilize data compression for their communication protocols. Wireless data transmission over sensor networks can consume a significant amount of energy and/or processing resources compared to data computation on the device itself. Thus data compression can be utilized to reduce the data size at the cost of extra processing time on the device. In one exemplification, the distributed computing system **5600** utilizes temporal correlation for sensing data, data transformation from one dimension to two dimension, and data separation (e.g., upper 8 bit and lower 8 bit data). In another exemplification, the distributed computing system **5600** utilizes a collection tree protocol for data collection from different nodes **5602a**, **5602b**, **5602c** having sensors (e.g., modular devices) to a root node. In yet another aspect, the distributed computing system **5600** utilizes first-order prediction coding to compress the data collected by the nodes **5602a**, **5602b**, **5602c** having sensors (e.g., modular devices), which can minimize the amount of redundant information and greatly reduce the amount of data transmission between the nodes **5602a**, **5602b**, **5602c** of the network. In yet another exemplification, the distributed computing system **5600** is configured to transmit only the electroencephalogram (EEG) features. In still yet another exemplification, the distributed computing system **5600** can be configured to transmit only the complex data features that are pertinent to the surgical instrument detection, which can save significant power in wireless transmission. Various other exemplifications can utilize combinations of the aforementioned data compression techniques and/or additional techniques of data compression.

[1042] FIG. 104 illustrates a logic flow diagram of a process **5650** for shifting distributed computing resources. In the following description of the **5650**, reference should also be made to FIG. 103. In one exemplification, the process **5650** can be executed by a distributed computing system including a control circuit of a surgical hub **206**, as depicted in FIG. 10 (processor **244**), in combination with a control circuit of a second surgical hub **206** and/or a control circuit of a modular device, such as the microcontroller **461** of the surgical instrument depicted FIG. 12, the microcontroller **620** of the surgical instrument depicted in FIG. 16, the control circuit **710** of the robotic surgical instrument **700** depicted in FIG. 17, the control circuit **760** of the surgical instruments **750**, **790** depicted in FIGS. 18 and 19, or the controller **838** of the generator **800** depicted in FIG. 20. For economy, the following description of the process **5650** will be described as being executed by the control circuits of one or more nodes; however, it should be understood that the description of the process **5650** encompasses all of the aforementioned exemplifications.

[1043] The control circuits of each node execute **5652** a distributed control program in synchrony. As the distributed control program is being executed across the network of nodes, at least one of the control circuits monitors for a command instructing the distributed computing system to shift from a first mode, wherein the distributed computing program is executed across the network of nodes, to a second mode, wherein the control program is executed by a single node. In one exemplification, the command can be

transmitted by a surgical hub in response to the surgical hub's resources being needed for an alternative computing task. In another exemplification, the command can be transmitted by a modular device in response to the rate at which the data is sampled by the modular device outpacing the rate at which the sampled data can be communicated to the other nodes in the network. If a control circuit determines that an appropriate command has been received **5654**, the process **5650** continues along the YES branch and the distributed computing system **5600** shifts to a single node executing **5656** the program. For example, the distributed computing system **5600** shifts the distributed computing program from being executed by both a modular device and a surgical hub to being executed solely by the modular device. As another example, the distributed computing system **5600** shifts the distributed computing program from being executed by both a first surgical hub and a second surgical hub to being executed solely by the first surgical hub. If no control circuit determines that an appropriate command has been received **5654**, the process continues along the NO branch and the control circuits of the network of nodes continues executing **5652** the distributed computing program across the network of nodes.

[1044] In the event that the program has been shifted to being executed **5656** by a single node, the control circuit of the particular node solely executing the distributed program and/or a control circuit of another node within the network (which previously was executing the distributed program) monitors for a command instructing the node to re-distribute the processing of the program across the distributed computing system. In other words, the node monitors for a command to re-initiate the distributed computing system. In one exemplification, the command to re-distribute the processing across the network can be generated when the sampling rate of the sensor is less than the data communication rate between the modular device and the surgical hub. If a control circuit receives **5658** an appropriate command to re-distribute the processing, then the process **5650** proceeds along the YES branch and the program is once again executed **5652** across the node network. If a control circuit has not received **5658** an appropriate command, then the node continues singularly executing **5656** the program.

[1045] The process **5650** depicted in FIG. 104 eliminates data or communication bottlenecks in controlling modular devices by utilizing a distributed computing architecture that can shift computing resources either between the modular devices and surgical hubs or between the surgical hubs as needed. This process **5650** also improves the modular devices' data processing speed by allowing the processing of the modular devices' control adjustments to be executed at least in part by the modular devices themselves. This process **5650** also improves the surgical hubs' data processing speed by allowing the surgical hubs to shift computing resources between themselves as necessary.

[1046] It can be difficult during video-assisted surgical procedures, such as laparoscopic procedures, to accurately measure sizes or dimensions of features being viewed through a medical imaging device due to distortive effects caused by the device's lens. Being able to accurately measure sizes and dimensions during video-assisted procedures could assist a situational awareness system for a surgical hub by allowing the surgical hub to accurately identify organs and other structures during video-assisted surgical procedures. As one solution, a surgical hub could be configured to

automatically calculate sizes or dimensions of structures (or distances between structures) during a surgical procedure by comparing the structures to markings affixed to devices that are intended to be placed within the FOV of the medical imaging device during a surgical procedure. The markings can represent a known scale, which can then be utilized to make measurements by comparing the unknown measured length to the known scale.

[1047] In one exemplification, the surgical hub is configured to receive image or video data from a medical imaging device paired with the surgical hub. When a surgical instrument bearing a calibration scale is within the FOV of the medical imaging device, the surgical hub is able to measure organs and other structures that are likewise within the medical imaging device's FOV by comparing the structures to the calibration scale. The calibration scale can be positioned on, for example, the distal end of a surgical instrument.

[1048] FIG. 105 illustrates a diagram of an imaging system 5800 and a surgical instrument 5806 bearing a calibration scale 5808. The imaging system 5800 includes a medical imaging device 5804 that is paired with a surgical hub 5802. The surgical hub 5802 can include a pattern recognition system or a machine learning system configured to recognize features in the FOV from image or video data received from the medical imaging device 5804. In one exemplification, a surgical instrument 5806 (e.g., a surgical cutting and stapling instrument) that is intended to enter the FOV of the medical imaging device 5804 during a surgical procedure includes a calibration scale 5808 affixed thereon. The calibration scale 5808 can be positioned on the exterior surface of the surgical instrument 5806, for example. In aspects wherein the surgical instrument 5806 is a surgical cutting and stapling instrument, the calibration scale 5808 can be positioned along the exterior surface of the anvil. The calibration scale 5808 can include a series of graphical markings separated at fixed and/or known intervals. The distance between the end or terminal markings of the calibration scale 5808 can likewise be a set distance  $L$  (e.g., 35 mm). In one exemplification, the end markings (e.g., the most proximal marking and the most distal marking) of the calibration scale 5808 are differentiated from the intermediate markings in size, shape, color, or another such fashion. This allows the image recognition system of the surgical hub 5802 to identify the end markings separately from the intermediate markings. The distance(s) between the markings can be stored in a memory or otherwise retrieved by the surgical hub 5802. The surgical hub 5802 can thus measure lengths or sizes of structures relative to the provided calibration scale 5808. In FIG. 105, for example, the surgical hub 5802 can calculate that the artery 5810a has a diameter or width of  $D_1$  (e.g., 17.0 mm), the vein 5810b has a diameter or width of  $D_2$  (e.g., 17.5 mm), and the distance between the vessels is  $D_3$  (e.g., 20 mm) by comparing the visualizations of these distances  $D_1$ ,  $D_2$ ,  $D_3$  to the known length  $L$  of the calibration scale 5808 positioned on the surgical instrument 5806 within the FOV of the medical imaging device 5804. The surgical hub 5802 can recognize the presence of the vessels 5810a, 5810b via an image recognition system. In some exemplifications, the surgical hub 5802 can be configured to automatically measure and display the size or dimension of detected features within the FOV of the medical imaging device 5804. In some exemplifications, the surgical hub 5802 can be configured to

calculate the distance between various points selected by a user on an interactive display that is paired with the surgical hub 5802.

[1049] The imaging system 5800 configured to detect and measure sizes according to a calibration scale 5808 affixed to surgical instruments 5806 provides the ability to accurately measure sizes and distances during video-assisted procedures. This can make it easier for surgeons to precisely perform video-assisted procedures by compensating for the optically distortive effects inherent in such procedures.

#### User Feedback Methods

[1050] The present disclosure provides user feedback techniques. In one aspect, the present disclosure provides a display of images through a medical imaging device (e.g., laparoscope, endoscope, thoracoscope, and the like). A medical imaging device comprises an optical component and an image sensor. The optical component may comprise a lens and a light source, for example. The image sensor may be implemented as a charge coupled device (CCD) or complementary oxide semiconductor (CMOS). The image sensor provides image data to electronic components in the surgical hub. The data representing the images may be transmitted by wired or wireless communication to display instrument status, feedback data, imaging data, and highlight tissue irregularities and underlining structures. In another aspect, the present disclosure provides wired or wireless communication techniques for communicating user feedback from a device (e.g., instrument, robot, or tool) to the surgical hub. In another aspect, the present disclosure provides identification and usage recording and enabling. Finally, in another aspect, the surgical hub may have a direct interface control between the device and the surgical hub.

#### Through Laparoscope Monitor Display of Data

[1051] In various aspects, the present disclosure provides through laparoscope monitor display of data. The through laparoscope monitor display of data may comprise displaying a current instrument alignment to adjacent previous operations, cooperation between local instrument displays and paired laparoscope display, and display of instrument specific data needed for efficient use of an end-effector portion of a surgical instrument. Each of these techniques is described hereinbelow.

#### Display of Current Instrument Alignment to Adjacent Previous Operations

[1052] In one aspect, the present disclosure provides alignment guidance display elements that provide the user information about the location of a previous firing or actuation and allow them to align the next instrument use to the proper position without the need for seeing the instrument directly. In another aspect, the first device and second device are separate; the first device is within the sterile field and the second is used from outside the sterile field.

[1053] During a colorectal transection using a double-stapling technique it is difficult to align the location of an anvil trocar of a circular stapler with the center of an overlapping staple line. During the procedure, the anvil trocar of the circular stapler is inserted in the rectum below the staple line and a laparoscope is inserted in the peritoneal cavity above the staple line. Because the staple line seals off the colon, there is no light of sight to align the anvil trocar

using the laparoscope to optically align the anvil trocar insertion location relative to the center of the staple line overlap.

**[1054]** One solution provides a non-contact sensor located on the anvil trocar of the circular stapler and a target located at the distal end of the laparoscope. Another solution provides a non-contact sensor located at the distal end of the laparoscope and a target located on the anvil trocar of the circular stapler.

**[1055]** A surgical hub computer processor receives signals from the non-contact sensor and displays a centering tool on a screen indicating the alignment of the anvil trocar of the circular stapler and the overlap portion at the center of staple line. The screen displays a first image of the target staple line with a radius around the staple line overlap portion and a second image of the projected anvil trocar location. The anvil trocar and the overlap portion at the center of staple line are aligned when the first and second images overlap.

**[1056]** In one aspect, the present disclosure provides a surgical hub for aligning a surgical instrument. The surgical hub comprises a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to receive image data from an image sensor, generate a first image based on the image data, display the first image on a monitor coupled to the processor, receive a signal from a non-contact sensor, generate a second image based on the position of the surgical device, and display the second image on the monitor. The first image data represents a center of a staple line seal. The first image represents a target corresponding to the center of the staple line. The signal is indicative of a position of a surgical device relative to the center of the staple line. The second image represents the position of the surgical device along a projected path of the surgical device toward the center of the staple line.

**[1057]** In one aspect, the center of the staple line is a double-staple overlap portion zone. In another aspect, the image sensor receives an image from a laparoscope. In another aspect, the surgical device is a circular stapler comprising an anvil trocar and the non-contact sensor is configured to detect the location of the anvil trocar relative to the center of the staple line seal. In another aspect, the non-contact sensor is an inductive sensor. In another aspect, the non-contact sensor is a capacitive sensor.

**[1058]** In various aspects, the present disclosure provides a control circuit to align the surgical instrument as described above. In various aspects, the present disclosure provides a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to align the surgical instrument as described above.

**[1059]** This technique provides better alignment of a surgical instrument such as a circular stapler about the overlap portion of the staple line to produce a better seal and cut after the circular stapler is fired.

**[1060]** In one aspect, the present disclosure provides a system for displaying the current instrument alignment relative to prior adjacent operations. The instrument alignment information may be displayed on a monitor or any suitable electronic device suitable for the visual presentation of data whether located locally on the instrument or remotely from the instrument through the modular communication hub. The system may display the current alignment of a circular staple cartridge to an overlapping staple line, display the current alignment of a circular staple cartridge relative to a prior linear staple line, and/or show the existing

staple line of the linear transection and an alignment circle indicating an appropriately centered circular staple cartridge. Each of these techniques is described hereinbelow.

**[1061]** In one aspect, the present disclosure provides alignment guidance display elements that provide the user information about the location of a previous firing or actuation of a surgical instrument (e.g., surgical stapler) and allows the user to align the next instrument use (e.g., firing or actuation of the surgical stapler) to the proper position without the need for seeing the instrument directly. In another aspect, the present disclosure provides a first device and a second device that is separate from the first device. The first device is located within a sterile field and the second is located outside the sterile field. The techniques described herein may be applied to surgical staplers, ultrasonic instruments, electrosurgical instruments, combination ultrasonic/electrosurgical instruments, and/or combination surgical stapler/electrosurgical instruments.

**[1062]** FIG. 106 illustrates a diagram 6000 of a surgical instrument 6002 centered on a staple line 6003 using the benefit of centering tools and techniques described in connection with FIGS. 23-33, according to one aspect of the present disclosure. As used in the following description of FIGS. 107-117 a staple line may include multiple rows of staggered staples and typically includes two or three rows of staggered staples, without limitation. The staple line may be a double staple line 6004 formed using a double-stapling technique as described in connection with FIGS. 107-111 or may be a linear staple line 6052 formed using a linear transection technique as described in connection with FIGS. 112-117. The centering tools and techniques described herein can be used to align the instrument 6002 located in one part of the anatomy with either the staple line 6003 or with another instrument located in another part of the anatomy without the benefit of a line of sight. The centering tools and techniques include displaying the current alignment of the instrument 6002 adjacent to previous operations. The centering tool is useful, for example, during laparoscopic-assisted rectal surgery that employ a double-stapling technique, also referred to as an overlapping stapling technique. In the illustrated example, during a laparoscopic-assisted rectal surgical procedure, a circular stapler 6002 is positioned in the rectum 6006 of a patient within the pelvic cavity 6008 and a laparoscope is positioned in the peritoneal cavity.

**[1063]** During the laparoscopic-assisted rectal surgery, the colon is transected and sealed by the staple line 6003 having a length "1." The double-stapling technique uses the circular stapler 6002 to create an end-to-end anastomosis and is currently used widely in laparoscopic-assisted rectal surgery. For a successful formation of an anastomosis using a circular stapler 6002, the anvil trocar 6010 of the circular stapler 6002 should be aligned with the center "1/2" of the staple line 6003 transection before puncturing through the center "1/2" of the staple line 6003 and/or fully clamping on the tissue before firing the circular stapler 6002 to cut out the staple overlap portion 6012 and forming the anastomosis. Misalignment of the anvil trocar 6010 to the center of the staple line 6003 transection may result in a high rate of anastomotic failures. This technique may be applied to ultrasonic instruments, electrosurgical instruments, combination ultrasonic/electrosurgical instruments, and/or combination surgical stapler/electrosurgical instruments. Several

techniques are now described for aligning the anvil trocar **6010** of the circular stapler **6002** to the center “ $\frac{1}{2}$ ” of the staple line **6003**.

[1064] In one aspect, as described in FIGS. 107-109 and with reference also to FIGS. 1-11 to show interaction with an interactive surgical system **100** environment including a surgical hub **106**, **206**, the present disclosure provides an apparatus and method for detecting the overlapping portion of the double staple line **6004** in a laparoscopic-assisted rectal surgery colorectal transection using a double stapling technique. The overlapping portion of the double staple line **6004** is detected and the current location of the anvil trocar **6010** of the circular stapler **6002** is displayed on a surgical hub display **215** coupled to the surgical hub **206**. The surgical hub display **215** displays the alignment of a circular stapler **6002** cartridge relative to the overlapping portion of the double staple line **6004**, which is located at the center of the double staple line **6004**. The surgical hub display **215** displays a circular image centered around the overlapping double staple line **6004** region to ensure that the overlapping portion of the double staple line **6004** is contained within the knife of the circular stapler **6002** and therefore removed following the circular firing. Using the display, the surgeon aligns the anvil trocar **6010** with the center of the double staple line **6004** before puncturing through the center of the double staple line **6004** and/or fully clamping on the tissue before firing the circular stapler **6002** to cut out the staple overlap portion **6012** and form the anastomosis.

[1065] FIGS. 107-109 illustrate a process of aligning an anvil trocar **6010** of a circular stapler **6022** to a staple overlap portion **6012** of a double staple line **6004** created by a double-stapling technique, according to one aspect of the present disclosure. The staple overlap portion **6012** is centered on the double staple line **6004** formed by a double-stapling technique. The circular stapler **6002** is inserted into the colon **6020** below the double staple line **6004** and a laparoscope **6014** is inserted through the abdomen above the double staple line **6004**. A laparoscope **6014** and a non-contact sensor **6022** are used to determine an anvil trocar **6010** location relative to the staple overlap portion **6012** of the double staple line **6004**. The laparoscope **6014** includes an image sensor to generate an image of the double staple line **6004**. The image sensor image is transmitted to the surgical hub **206** via the imaging module **238**. The sensor **6022** generates a signal **6024** that detects the metal staples using inductive or capacitive metal sensing technology. The signal **6024** varies based on the position of the anvil trocar **6010** relative to the staple overlap portion **6004**. A centering tool **6030** presents an image **6038** of the double staple line **6004** and a target alignment ring **6032** circumscribing the image **6038** of the double staple line **6004** centered about an image **6040** of the staple overlap portion **6012** on the surgical hub display **215**. The centering tool **6030** also presents a projected cut path **6034** of an anvil knife of the circular stapler **6002**. The alignment process includes displaying an image **6038** of the double staple line **6004** and a target alignment ring **6032** circumscribing the image **6038** of the double staple line **6004** centered on the image **6040** of the staple overlap portion **6012** to be cut out by the circular knife of the circular stapler **6002**. Also displayed is an image of a crosshair **6036** (X) relative to the image **6040** of the staple overlap portion **6012**.

[1066] FIG. 107 illustrates an anvil trocar **6010** of a circular stapler **6002** that is not aligned with a staple overlap

portion **6012** of a double staple line **6004** created by a double-stapling technique. The double staple line **6004** has a length “1” and the staple overlap portion **6012** is located midway along the double staple line **6004** at “ $\frac{1}{2}$ .” As shown in FIG. 107, the circular stapler **6002** is inserted into a section of the colon **6020** and is positioned just below the double staple line **6004** transection. A laparoscope **6014** is positioned above the double staple line **6004** transection and feeds an image of the double staple line **6004** and staple overlap portion **6012** within the field of view **6016** of the laparoscope **6014** to the surgical hub display **215**. The position of the anvil trocar **6010** relative to the staple overlap portion **6012** is detected by a sensor **6022** located on the circular stapler **6002**. The sensor **6022** also provides the position of the anvil trocar **6010** relative to the staple overlap portion **6012** to the surgical hub display **215**.

[1067] As shown in FIG. 107, the projected path **6018** of the anvil trocar **6010** is shown along a broken line to a position marked by an X. As shown in FIG. 107, the projected path **6018** of the anvil trocar **6010** is not aligned with the staple overlap portion **6012**. Puncturing the anvil trocar **6010** through the double staple line **6004** at a point off the staple overlap portion **6012** could lead to an anastomotic failure. Using the anvil trocar **6010** centering tool **6030** described in FIG. 109, the surgeon can align the anvil trocar **6010** with the staple overlap portion **6012** using the images displayed by the centering tool **6030**. For example, in one implementation, the sensor **6022** is an inductive sensor. Since the staple overlap portion **6012** contains more metal than the rest of the lateral portions of the double staple line **6004**, the signal **6024** is maximum when the sensor **6022** is aligned with and proximate to the staple overlap portion **6012**. The sensor **6022** provides a signal to the surgical hub **206** that indicates the location of the anvil trocar **6010** relative to the staple overlap portion **6012**. The output signal is converted to a visualization of the location of the anvil trocar **6010** relative to the staple overlap portion **6012** that is displayed on the surgical hub display **215**.

[1068] As shown in FIG. 108, the anvil trocar **6010** is aligned with the staple overlap portion **6012** at the center of the double staple line **6004** created by a double-stapling technique. The surgeon can now puncture the anvil trocar **6010** through the staple overlap portion **6012** of the double staple line **6004** and/or fully clamp on the tissue before firing the circular stapler **6002** to cut out the staple overlap portion **6012** and form an anastomosis.

[1069] FIG. 109 illustrates a centering tool **6030** displayed on a surgical hub display **215**, the centering tool providing a display of a staple overlap portion **6012** of a double staple line **6004** created by a double-stapling technique, where the anvil trocar **6010** is not aligned with the staple overlap portion **6012** of the double staple line **6004** as shown in FIG. 107. The centering tool **6030** presents an image **6038** on the surgical hub display **215** of the double staple line **6004** and an image **6040** of the staple overlap portion **6012** received from the laparoscope **6014**. A target alignment ring **6032** centered about the image **6040** of the staple overlap portion **6012** circumscribes the image **6038** of the double staple line **6004** to ensure that the staple overlap portion **6012** is located within the circumference of the projected cut path **6034** of the circular stapler **6002** knife when the projected cut path **6034** is aligned to the target alignment ring **6032**. The crosshair **6036** (X) represents the location of the anvil trocar **6010** relative to the staple overlap portion **6012**. The

crosshair **6036 (X)** indicates the point through the double staple line **6004** where the anvil trocar **6010** would puncture if it were advanced from its current location.

[1070] As shown in FIG. 109, the anvil trocar **6010** is not aligned with the desired puncture through location designated by the image **6040** of the staple overlap portion **6012**. To align the anvil trocar **6010** with the staple overlap portion **6012** the surgeon manipulates the circular stapler **6002** until the projected cut path **6034** overlaps the target alignment ring **6032** and the crosshair **6036 (X)** is centered on the image **6040** of the staple overlap portion **6012**. Once alignment is complete, the surgeon punctures the anvil trocar **6010** through the staple overlap portion **6012** of the double staple line **6004** and/or fully clamps on the tissue before firing the circular stapler **6002** to cut out the staple overlap portion **6012** and form the anastomosis.

[1071] As discussed above, the sensor **6022** is configured to detect the position of the anvil trocar **6010** relative to the staple overlap portion **6012**. Accordingly, the location of the crosshair **6036 (X)** presented on the surgical hub display **215** is determined by the surgical stapler sensor **6022**. In another aspect, the sensor **6022** may be located on the laparoscope **6014**, where the sensor **6022** is configured to detect the tip of the anvil trocar **6010**. In other aspects, the sensor **6022** may be located either on the circular stapler **6022** or the laparoscope **6014**, or both, to determine the location of the anvil trocar **6010** relative to the staple overlap portion **6012** and provide the information to the surgical hub display **215** via the surgical hub **206**.

[1072] FIGS. 110 and 111 illustrate a before image **6042** and an after image **6043** of a centering tool **6030**, according to one aspect of the present disclosure. FIG. 110 illustrates an image of a projected cut path **6034** of an anvil trocar **6010** and circular knife before alignment with the target alignment ring **6032** circumscribing the image **6038** of the double staple line **6004** over the image **6040** of the staple overlap portion **6040** presented on a surgical hub display **215**. FIG. 111 illustrates an image of a projected cut path **6034** of an anvil trocar **6010** and circular knife after alignment with the target alignment ring **6032** circumscribing the image **6038** of the double staple line **6004** over the image **6040** of the staple overlap portion **6040** presented on a surgical hub display **215**. The current location of the anvil trocar **6010** is marked by the crosshair **6036 (X)**, which as shown in FIG. 110, is positioned below and to the left of center of the image **6040** of the staple overlap portion **6040**. As shown in FIG. 111, as the surgeon moves the anvil trocar **6010** of the along the projected path **6046**, the projected cut path **6034** aligns with the target alignment ring **6032**. The target alignment ring **6032** may be displayed as a greyed out alignment circle overlaid over the current position of the anvil trocar **6010** relative to the center of the double staple line **6004**, for example. The image may include indication marks to assist the alignment process by indication which direction to move the anvil trocar **6010**. The target alignment ring **6032** may be shown in bold, change color or may be highlighted when it is located within a predetermined distance of center within acceptable limits.

[1073] In another aspect, the sensor **6022** may be configured to detect the beginning and end of a linear staple line in a colorectal transection and to provide the position of the current location of the anvil trocar **6010** of the circular stapler **6002**. In another aspect, the present disclosure provides a surgical hub display **215** to present the circular

stapler **6002** centered on the linear staple line, which would create even dog ears, and to provide the current position of the anvil trocar **6010** to allow the surgeon to center or align the anvil trocar **6010** as desired before puncturing and/or fully clamping on tissue prior to firing the circular stapler **6002**.

[1074] In another aspect, as described in FIGS. 112-114 and with reference also to FIGS. 1-11 to show interaction with an interactive surgical system **100** environment including a surgical hub **106, 206**, in a laparoscopic-assisted rectal surgery colorectal transection using a linear stapling technique, the beginning and end of the linear staple line **6052** is detected and the current location of the anvil trocar **6010** of the circular stapler **6002** is displayed on a surgical hub display **215** coupled to the surgical hub **206**. The surgical hub display **215** displays a circular image centered on the double staple line **6004**, which would create even dog ears and the current position of the anvil trocar **6002** is displayed to allow the surgeon to center or align the anvil trocar **6010** before puncturing through the linear staple line **6052** and/or fully clamping on the tissue before firing the circular stapler **6002** to cut out the center **6050** of the linear staple line **6052** to form an anastomosis.

[1075] FIGS. 112-114 illustrate a process of aligning an anvil trocar **6010** of a circular stapler **6022** to a center **6050** of a linear staple line **6052** created by a linear stapling technique, according to one aspect of the present disclosure. FIGS. 112 and 113 illustrate a laparoscope **6014** and a sensor **6022** located on the circular stapler **6022** to determine the location of the anvil trocar **6010** relative to the center **6050** of the linear staple line **6052**. The anvil trocar **6010** and the sensor **6022** is inserted into the colon **6020** below the linear staple line **6052** and the laparoscope **6014** is inserted through the abdomen above the linear staple line **6052**.

[1076] FIG. 112 illustrates the anvil trocar **6010** out of alignment with the center **6050** of the linear staple line **6052** and FIG. 113 illustrates the anvil trocar **6010** in alignment with the center **6050** of the linear staple line **6052**. The sensor **6022** is used to detect the center **6050** of the linear staple line **6052** to align the anvil trocar **6010** with the center of the staple line **6052**. In one aspect, the center **6050** of the linear staple line **6052** may be located by moving the circular stapler **6002** until one end of the linear staple line **6052** is detected. An end may be detected when there are no more staples in the path of the sensor **6022**. Once one of the ends is reached, the circular stapler **6002** is moved along the linear staple line **6053** until the opposite end is detected and the length "1" of the linear staple line **6052** is determined by measurement or by counting individual staples by the sensor **6022**. Once the length of the linear staple line **6052** is determined, the center **6050** of the linear staple line **6052** can be determined by dividing the length by two " $\frac{1}{2}$ ."

[1077] FIG. 114 illustrates a centering tool **6054** displayed on a surgical hub display **215**, the centering tool providing a display of a linear staple line **6052**, where the anvil trocar **6010** is not aligned with the staple overlap portion **6012** of the double staple line **6004** as shown in FIG. 112. The surgical hub display **215** presents a standard reticle field of view **6056** of the laparoscopic field of view **6016** of the linear staple line **6052** and a portion of the colon **6020**. The surgical hub display **215** also presents a target ring **6062** circumscribing the image center of the linear staple line and a projected cut path **6064** of the anvil trocar and circular knife. The crosshair **6066 (X)** represents the location of the

anvil trocar **6010** relative to the center **6050** of the linear staple line **6052**. The crosshair **6036** (X) indicates the point through the linear staple line **6052** where the anvil trocar **6010** would puncture if it were advanced from its current location.

[1078] As shown in FIG. 114, the anvil trocar **6010** is not aligned with the desired puncture through location designated by the offset between the target ring **6062** and the projected cut path **6064**. To align the anvil trocar **6010** with the center **6050** of the linear staple line **6052** the surgeon manipulates the circular stapler **6002** until the projected cut path **6064** overlaps the target alignment ring **6062** and the crosshair **6066** (X) is centered on the image **6040** of the staple overlap portion **6012**. Once alignment is complete, the surgeon punctures the anvil trocar **6010** through the center **6050** of the linear staple line **6052** and/or fully clamps on the tissue before firing the circular stapler **6002** to cut out the staple overlap portion **6012** and forming the anastomosis.

[1079] In one aspect, the present disclosure provides an apparatus and method for displaying an image of an linear staple line **6052** using a linear transection technique and an alignment ring or bullseye positioned as if the anvil trocar **6010** of the circular stapler **6022** were centered appropriately along the linear staple line **6052**. The apparatus displays a greyed out alignment ring overlaid over the current position of the anvil trocar **6010** relative to the center **6050** of the linear staple line **6052**. The image may include indication marks to assist the alignment process by indication which direction to move the anvil trocar **6010**. The alignment ring may be bold, change color or highlight when it is located within a predetermined distance of centered.

[1080] With reference now to FIGS. 112-115, FIG. 115 is an image **6080** of a standard reticle field view **6080** of a linear staple line **6052** transection of a surgical as viewed through a laparoscope **6014** displayed on the surgical hub display **215**, according to one aspect of the present disclosure. In a standard reticle view **6080**, it is difficult to see the linear staple line **6052** in the standard reticle field of view **6056**. Further, there are no alignment aids to assist with alignment and introduction of the anvil trocar **6010** to the center **6050** of the linear staple line. This view does not show an alignment circle or alignment mark to indicate if the circular stapler is centered appropriately and does not show the projected trocar path. In this view it also difficult to see the staples because there is no contrast with the background image.

[1081] With reference now to FIGS. 112-116, FIG. 116 is an image **6082** of a laser-assisted reticle field of view **6072** of the surgical site shown in FIG. 115 before the anvil trocar **6010** and circular knife of the circular stapler **6002** are aligned to the center **6050** of the linear staple line **6052**, according to one aspect of the present disclosure. The laser-assisted reticle field of view **6072** provides an alignment mark or crosshair **6066** (X), currently positioned below and to the left of center of the linear staple line **6052** showing the projected path of the anvil trocar **6010** to assist positioning of the anvil trocar **6010**. In addition to the projected path marked by the crosshair **6066** (X) of the anvil trocar **6010**, the image **6082** displays the staples of the linear staple line **6052** in a contrast color to make them more visible against the background. The linear staple line **6052** is highlighted and a bullseye target **6070** is displayed over the center **6050** of the linear staple line **6052**. Outside of the laser-assisted reticle field of view **6072**, the image **6082**

displays a status warning box **6068**, a suggestion box **6074**, a target ring **6062**, and the current alignment position of the anvil trocar **6010** marked by the crosshair **6066** (X) relative to the center **6050** of the linear staple line **6052**. As shown in FIG. 116, the status warning box **6068** indicates that the trocar is "MISALIGNED" and the suggestion box **6074** states "Adjust trocar to center staple line."

[1082] With reference now to FIGS. 112-117, FIG. 117 is an image **6084** of a laser-assisted reticle field of view **6072** of the surgical site shown in FIG. 116 after the anvil trocar **6010** and circular knife of the circular stapler **6002** are aligned to the center **6050** of the linear staple line **6052**, according to one aspect of the present disclosure. The laser-assisted reticle field of view **6072** provides an alignment mark or crosshair **6066** (X), currently positioned below and to the left of center of the linear staple line **6052** showing the projected path of the anvil trocar **6010** to assist positioning of the anvil trocar **6010**. In addition to the projected path marked by the crosshair **6066** (X) of the anvil trocar **6010**, the image **6082** displays the staples of the linear staple line **6052** in a contrast color to make them more visible against the background. The linear staple line **6052** is highlighted and a bullseye target **6070** is displayed over the center **6050** of the linear staple line **6052**. Outside of the laser-assisted reticle field of view **6072**, the image **6082** displays a status warning box **6068**, a suggestion box **6074**, a target ring **6062**, and the current alignment position of the anvil trocar **6010** marked by the crosshair **6066** (X) relative to the center **6050** of the linear staple line **6052**. As shown in FIG. 116, the status warning box **6068** indicates that the trocar is "MISALIGNED" and the suggestion box **6074** states "Adjust trocar to center staple line."

[1083] FIG. 117 is a laser assisted view of the surgical site shown in FIG. 116 after the anvil trocar **6010** and circular knife are aligned to the center of the staple line **6052**. In this view, inside the field of view **6072** of the laser-assisted reticle, the alignment mark crosshair **6066** (X) is positioned over the center of the staple line **6052** and the highlighted bullseye target to indicate alignment of the trocar to the center of the staple line. Outside the field of view **6072** of the laser-assisted reticle, the status warning box indicates that the trocar is "ALIGNED" and the suggestion is "Proceed trocar introduction."

[1084] FIG. 118 illustrates a non-contact inductive sensor **6090** implementation of the non-contact sensor **6022** to determine an anvil trocar **6010** location relative to the center of a staple line transection (the staple overlap portion **6012** of the double staple line **6004** shown in FIGS. 107-108 or the center **6050** of the linear staple line **6052** shown in FIGS. 112-113, for example), according to one aspect of the present disclosure. The non-contact inductive sensor **6090** includes an oscillator **6092** that drives an inductive coil **6094** to generate an electromagnetic field **6096**. As a metal target **6098**, such as a metal staple, is introduced into the electromagnetic field **6096**, eddy currents **6100** induced in the target **6098** oppose the electromagnetic field **6096** and the reluctance shifts and the amplitude of the oscillator voltage **6102** drops. An amplifier **6104** amplifies the oscillator voltage **6102** amplitude as it changes.

[1085] With reference now to FIGS. 1-11 to show interaction with an interactive surgical system **100** environment including a surgical hub **106**, **206** and also to FIGS. 106-117, the inductive sensor **6090** is a non-contact electronic sensor. It can be used for positioning and detecting metal objects

such as the metal staples in the staple lines **6003**, **6004**, **6052** described above. The sensing range of the inductive sensor **6090** is dependent on the type of metal being detected. Because the inductive sensor **6090** is a non-contact sensor, it can detect metal objects across a stapled tissue barrier. The inductive sensor **6090** can be located either on the circular stapler **6002** to detect staples in the staple lines **6003**, **6004**, **6052**, detect the location of the distal end of the laparoscope **6014**, or it may be located on the laparoscope **6014** to detect the location of the anvil trocar **6010**. A processor or control circuit located either in the circular stapler **6002**, laparoscope **6014**, or coupled to the surgical hub **206** receives signals from the inductive sensors **6090** and can be employed to display the centering tool on the surgical hub display **215** to determine the location of the anvil trocar **6010** relative to either staple overlap portion **6012** of a double staple line **6004** or the center **6050** of a linear staple line **6052**.

[1086] In one aspect, the distal end of the laparoscope **6014** may be detected by the inductive sensor **6090** located on the circular stapler **6002**. The inductive sensor **6090** may detect a metal target **6098** positioned on the distal end of the laparoscope **6014**. Once the laparoscope **6014** is aligned with the center **6050** of the linear staple line **6052** or the staple overlap portion **6012** of the double staple line **6004**, a signal from the inductive sensor **6090** is transmitted to circuits that convert the signals from the inductive sensor **6090** to present an image of the relative alignment of the laparoscope **6014** with the anvil trocar **6010** of the circular stapler **6002**.

[1087] FIGS. **119A** and **119B** illustrate one aspect of a non-contact capacitive sensor **6110** implementation of the non-contact sensor **6022** to determine an anvil trocar **6010** location relative to the center of a staple line transection (the staple overlap portion **6012** of the double staple line **6004** shown in FIGS. **107-108** or the center **6050** of the linear staple line **6052** shown in FIGS. **112-113**, for example), according to one aspect of the present disclosure. FIG. **119A** shows the non-contact capacitive sensor **6110** without a nearby metal target and FIG. **119B** shows the non-contact capacitive sensor **6110** near a metal target **6112**. The non-contact capacitive sensor **6110** includes capacitor plates **6114**, **6116** housed in a sensing head and establishes field lines **6118** when energized by an oscillator waveform to define a sensing zone. FIG. **119A** shows the field lines **6118** when no target is present proximal to the capacitor plates **6114**, **6116**. FIG. **119B** shows a ferrous or nonferrous metal target **6120** in the sensing zone. As the metal target **6120** enters the sensing zone, the capacitance increases causing the natural frequency to shift towards the oscillation frequency causing amplitude gain. Because the capacitive sensor **6110** is a non-contact sensor, it can detect metal objects across a stapled tissue barrier. The capacitive sensor **6110** can be located either on the circular stapler **6002** to detect the staple lines **6004**, **6052** or the location of the distal end of the laparoscope **6014** or the capacitive sensor **6110** may be located on the laparoscope **6014** to detect the location of the anvil trocar **6010**. A processor or control circuit located either in the circular stapler **6002**, the laparoscope **6014**, or coupled to the surgical hub **206** receives signals from the capacitive sensor **6110** to present an image of the relative alignment of the laparoscope **6014** with the anvil trocar **6010** of the circular stapler **6002**.

[1088] FIG. **120** is a logic flow diagram **6130** of a process depicting a control program or a logic configuration for aligning a surgical instrument, according to one aspect of the present disclosure. With reference to FIGS. **1-11** to show interaction with an interactive surgical system **100** environment including a surgical hub **106**, **206** and also to FIGS. **106-119**, the surgical hub **206** comprises a processor **244** and a memory **249** coupled to the processor **244**. The memory **249** stores instructions executable by the processor **244** to receive **6132** image data from a laparoscope image sensor, generate **6134** a first image based on the image data, display **6136** the first image on a surgical hub display **215** coupled to the processor **244**, receive **6138** a signal from a non-contact sensor **6022**, the signal indicative of a position of a surgical device, generate a second image based on the signal indicative of the position of the surgical device, e.g., the anvil trocar **6010** and display **6140** the second image on the surgical hub display **215**. The first image data represents a center **6044**, **6050** of a staple line **6004**, **6052** seal. The first image represents a target corresponding to the center **6044**, **6050** of the staple line **6004**, **6052** seal. The signal is indicative of a position of a surgical device, e.g., an anvil trocar **6010**, relative to the center **6044**, **6050** of the staple line **6004**, **6052** seal. The second image represents the position of the surgical device, e.g., an anvil trocar **6010**, along a projected path **6018** of the surgical device, e.g., an anvil trocar **6010**, toward the center **6044**, **6050** of the staple line **6004**, **6052** seal.

[1089] In one aspect, the center **6044** of the double staple line **6004** seal defines a staple overlap portion **6012**. In another aspect, an image sensor receives an image from a medical imaging device. In another aspect, the surgical device is a circular stapler **6002** comprising an anvil trocar **6010** and the non-contact sensor **6022** is configured to detect the location of the anvil trocar **6010** relative to the center **6044** of the double staple line **6004** seal. In another aspect, the non-contact sensor **6022** is an inductive sensor **6090**. In another aspect, the non-contact sensor **6022** is a capacitive sensor **6110**. In one aspect, the staple line may be a linear staple line **6052** formed using a linear transection technique.

#### Cooperation Between Local Instrument Displays and Paired Imaging Device Display

[1090] In one aspect, the present disclosure provides an instrument including a local display, a hub having an operating room (OR), or operating theater, display separate from the instrument display. When the instrument is linked to the surgical hub, the secondary display on the device reconfigures to display different information than when it is independent of the surgical hub connection. In another aspect, some portion of the information on the secondary display of the instrument is then displayed on the primary display of the surgical hub. In another aspect, image fusion allowing the overlay of the status of a device, the integration landmarks being used to interlock several images and at least one guidance feature are provided on the surgical hub and/or instrument display. Techniques for overlaying or augmenting images and/or text from multiple image/text sources to present composite images on a single display are described hereinbelow in connection with FIGS. **129-137** and FIGS. **147-151**.

[1091] In another aspect, the present disclosure provides cooperation between local instrument displays and a paired laparoscope display. In one aspect, the behavior of a local

display of an instrument changes when it senses the connectable presence of a global display coupled to the surgical hub. In another aspect, the present disclosure provides 3600 composite top visual field of view of a surgical site to avoid collateral structures. Each of these techniques is described hereinbelow.

[1092] During a surgical procedure, the surgical site is displayed on a remote “primary” surgical hub display. During a surgical procedure, surgical devices track and record surgical data and variables (e.g., surgical parameters) that are stored in the instrument (see FIGS. 12-19 for instrument architectures comprising processors, memory, control circuits, storage, etc.). The surgical parameters include force-to-fire (FTF), force-to-close (FTC), firing progress, tissue gap, power level, impedance, tissue compression stability (creep), and the like. Using conventional techniques during the procedure the surgeon needs to watch two separate displays. Providing image/text overlay is thus advantageous because during the procedure the surgeon can watch a single display presenting the overlaid image/text information.

[1093] One solution detects when the surgical device (e.g., instrument) is connected to the surgical hub and then display a composite image on the primary display that includes a field of view of the surgical site received from a first instrument (e.g., medical imaging device such as, e.g., laparoscope, endoscope, thoracoscope, and the like) augmented by surgical data and variables received from a second instrument (e.g., a surgical stapler) to provide pertinent images and data on the primary display.

[1094] During a surgical procedure the surgical site is displayed as a narrow field of view of a medical imaging device on the primary surgical hub display. Items outside the current field of view, collateral structures, cannot be viewed without moving the medical imaging device.

[1095] One solution provides a narrow field of view of the surgical site in a first window of the display augmented by a wide field of view of the surgical site in a separate window of the display. This provides a composite over head field of view mapped using two or more imaging arrays to provide an augmented image of multiple perspective views of the surgical site.

[1096] In one aspect, the present disclosure provides a surgical hub, comprising a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to detect a surgical device connection to the surgical hub, transmit a control signal to the detected surgical device to transmit to the surgical hub surgical parameter data associated with the detected device, receive the surgical parameter data, receive image data from an image sensor, and display, on a display coupled to the surgical hub, an image received from the image sensor in conjunction with the surgical parameter data received from the surgical device.

[1097] In another aspect, the present disclosure provides a surgical hub, comprising a processor and a memory coupled to the processor. The memory stores instructions executable by the processor to receive first image data from a first image sensor, receive second image data from a second image sensor, and display, on a display coupled to the surgical hub, a first image corresponding to the first field of view and a second image corresponding to the second field of view. The first image data represents a first field of view and the second image data represents a second field of view.

[1098] In one aspect, the first field of view is a narrow angle field of view and the second field of view is a wide angle field of view. In another aspect, the memory stores instructions executable by the processor to augment the first image with the second image on the display. In another aspect, the memory stores instructions executable by the processor to fuse the first image and the second image into a third image and display a fused image on the display. In another aspect, the fused image data comprises status information associated with a surgical device, an image data integration landmark to interlock a plurality of images, and at least one guidance parameter. In another aspect, the first image sensor is the same as the same image sensor and wherein the first image data is captured as a first time and the second image data is captured at a second time.

[1099] In another aspect, the memory stores instructions executable by the processor to receive third image data from a third image sensor, wherein the third image data represents a third field of view, generate composite image data comprising the second and third image data, display the first image in a first window of the display, wherein the first image corresponds to the first image data, and display a third image in a second window of the display, wherein the third image corresponds to the composite image data.

[1100] In another aspect, the memory stores instructions executable by the processor to receive third image data from a third image sensor, wherein the third image data represents a third field of view, fuse the second and third image data to generate fused image data, display the first image in a first window of the display, wherein the first image corresponds to the first image data, and display a third image in a second window of the display, wherein the third image corresponds to the fused image data.

[1101] In various aspects, the present disclosure provides a control circuit to perform the functions described above. In various aspects, the present disclosure provides a non-transitory computer readable medium storing computer readable instructions, which when executed, causes a machine to perform the functions described above.

[1102] By displaying endoscope images augmented with surgical device images on one primary surgical hub display, enables the surgeon to focus on one display to obtain a field of view of the surgical site augmented with surgical device data associated with the surgical procedure such as force-to-fire, force-to-close, firing progress, tissue gap, power level, impedance, tissue compression stability (creep), and the like.

[1103] Displaying a narrow field of view image in a first window of a display and a composite image of several other perspectives such as wider fields of view enables the surgeon to view a magnified image of the surgical site simultaneously with wider fields of view of the surgical site without moving the scope.

[1104] In one aspect, the present disclosure provides both global and local display of a device, e.g., a surgical instrument, coupled to the surgical hub. The device displays all of its relevant menus and displays on a local display until it senses a connection to the surgical hub at which point a sub-set of the information is displayed only on the monitor through the surgical hub and that information is either mirrored on the device display or is no longer accessible on the device detonated screen. This technique frees up the device display to show different information or display larger font information on the surgical hub display.



[1105] In one aspect, the present disclosure provides an instrument having a local display, a surgical hub having an operating theater (e.g., operating room or OR) display that is separate from the instrument display. When the instrument is linked to the surgical hub, the instrument local display becomes a secondary display and the instrument reconfigures to display different information than when it is operating independent of the surgical hub connection. In another aspect, some portion of the information on the secondary display is then displayed on the primary display in the operating theater through the surgical hub.

[1106] FIG. 121 illustrates a primary display 6200 of the surgical hub 206 comprising a global display 6202 and a local instrument display 6204, according to one aspect of the present disclosure. With continued reference to FIGS. 1-11 to show interaction with an interactive surgical system 100 environment including a surgical hub 106, 206 and FIGS. 12-21 for surgical hub connected instruments together with FIG. 121, the local instrument display 6204 behavior is displayed when the instrument 235 senses the connectable presence of a global display 6202 through the surgical hub 206. The global display 6202 shows a field of view 6206 of a surgical site 6208, as viewed through a medical imaging device such as, for example, a laparoscope/endoscope 219 coupled to an imaging module 238, at the center of the surgical hub display 215, referred to herein also as a monitor, for example. The end effector 6218 portion of the connected instrument 235 is shown in the field of view 6206 of the surgical site 6208 in the global display 6202. The images shown on the display 237 located on an instrument 235 coupled to the surgical hub 206 is shown, or mirrored, on the local instrument display 6204 located in the lower right corner of the monitor 6200 as shown in FIG. 121, for example. During operation, all relevant instrument and information and menus are displayed on the display 237 located on the instrument 235 until the instrument 235 senses a connection of the instrument 235 to the surgical hub 206 at which point all or some sub-set of the information presented on the instrument display 237 is displayed only on the local instrument display 6204 portion of the surgical hub display 6200 through the surgical hub 206. The information displayed on the local instrument display 6204 may be mirrored on the display 237 located on the instrument 235 or may be no longer accessible on the instrument display 237 detuned screen. This technique frees up the instrument 235 to show different information or to show larger font information on the surgical hub display 6200. Several techniques for overlaying or augmenting images and/or text from multiple image/text sources to present composite images on a single display are described hereinbelow in connection with FIGS. 129-137 and FIGS. 147-151.

[1107] The surgical hub display 6200 provides perioperative visualization of the surgical site 6208. Advanced imaging identifies and visually highlights 6222 critical structures such as the ureter 6220 (or nerves, etc.) and also tracks instrument proximity displays 6210 and shown on the left side of the display 6200. In the illustrated example, the instrument proximity displays 6210 show instrument specific settings. For example the top instrument proximity display 6212 shows settings for a monopolar instrument, the middle instrument proximity display 6214 shows settings for a bipolar instrument, and the bottom instrument proximity display 6212 shows settings for an ultrasonic instrument.

[1108] In another aspect, independent secondary displays or dedicated local displays can be linked to the surgical hub 206 to provide both an interaction portal via a touchscreen display and/or a secondary screen that can display any number of surgical hub 206 tracked data feeds to provide a clear non-confusing status. The secondary screen may display force to fire (FTF), tissue gap, power level, impedance, tissue compression stability (creep), etc., while the primary screen may display only key variables to keep the feed free of clutter. The interactive display may be used to move the display of specific information to the primary display to a desired location, size, color, etc. In the illustrated example, the secondary screen displays the instrument proximity displays 6210 on the left side of the display 6200 and the local instrument display 6204 on the bottom right side of the display 6200. The local instrument display 6204 presented on the surgical hub display 6200 displays an icon of the end effector 6218, such as the icon of a staple cartridge 6224 currently in use, the size 6226 of the staple cartridge 6224 (e.g., 60 mm), and an icon of the current position of the knife 6228 of the end effector.

[1109] In another aspect, the display 237 located on the instrument 235 displays the wireless or wired attachment of the instrument 235 to the surgical hub 206 and the instrument's communication/recording on the surgical hub 206. A setting may be provided on the instrument 235 to enable the user to select mirroring or extending the display to both monitoring devices. The instrument controls may be used to interact with the surgical hub display of the information being sourced on the instrument. As previously discussed, the instrument 235 may comprise wireless communication circuits to communicate wirelessly with the surgical hub 206.

[1110] In another aspect, a first instrument coupled to the surgical hub 206 can pair to a screen of a second instrument coupled to the surgical hub 206 allowing both instruments to display some hybrid combination of information from the two devices of both becoming mirrors of portions of the primary display. In yet another aspect, the primary display 6200 of the surgical hub 206 provides a 3600 composite top visual view of the surgical site 6208 to avoid collateral structures. For example, a secondary display of the end-effector surgical stapler may be provided within the primary display 6200 of the surgical hub 206 or on another display in order to provide better perspective around the areas within a current the field of view 6206. These aspects are described hereinbelow in connection with FIGS. 122-124.

[1111] FIGS. 122-124 illustrate a composite overhead views of an end-effector 6234 portion of a surgical stapler mapped using two or more imaging arrays or one array and time to provide multiple perspective views of the end-effector 6234 to enable the composite imaging of an overhead field of view. The techniques described herein may be applied to ultrasonic instruments, electrosurgical instruments, combination ultrasonic/electrosurgical instruments, and/or combination surgical stapler/electrosurgical instruments. Several techniques for overlaying or augmenting images and/or text from multiple image/text sources to present composite images on a single display are described hereinbelow in connection with FIGS. 129-137 and FIGS. 147-151.

[1112] FIG. 122 illustrates a primary display 6200 of the surgical hub 206, according to one aspect of the present disclosure. A primary window 6230 is located at the center

of the screen shows a magnified or exploded narrow angle view of a surgical field of view **6232**. The primary window **6230** located in the center of the screen shows a magnified or narrow angle view of an end-effector **6234** of the surgical stapler grasping a vessel **6236**. The primary window **6230** displays knitted images to produce a composite image that enables visualization of structures adjacent to the surgical field of view **6232**. A second window **6240** is shown in the lower left corner of the primary display **6200**. The second window **6240** displays a knitted image in a wide angle view at standard focus of the image shown in the primary window **6230** in an overhead view. The overhead view provided in the second window **6240** enables the viewer to easily see items that are out of the narrow field surgical field of view **6232** without moving the laparoscope, or other imaging device **239** coupled to the imaging module **238** of the surgical hub **206**. A third window **6242** is shown in the lower right corner of the primary display **6200** shows an icon **6244** representative of the staple cartridge of the end-effector **6234** (e.g., a staple cartridge in this instance) and additional information such as “4 Row” indicating the number of staple rows **6246** and “35 mm” indicating the distance **6248** traversed by the knife along the length of the staple cartridge. Below the third window **6242** is displayed an icon **6258** of a frame of the current state of a clamp stabilization sequence **6250** (FIG. **123**) that indicates clamp stabilization.

[1113] FIG. **123** illustrates a clamp stabilization sequence **6250** over a five second period, according to one aspect of the present disclosure. The clamp stabilization sequence **6250** is shown over a five second period with intermittent displays **6252**, **6254**, **6256**, **6258**, **6260** spaced apart at one second intervals **6268** in addition to providing the real time **6266** (e.g., 09:35:10), which may be a pseudo real time to preserve anonymity of the patient. The intermittent displays **6252**, **6254**, **6256**, **6258**, **6260** show elapsed by filling in the circle until the clamp stabilization period is complete. At that point, the last display **6260** is shown in solid color. Clamp stabilization after the end effector **6234** clamps the vessel **6236** enables the formation of a better seal.

[1114] FIG. **124** illustrates a diagram **6270** of four separate wide angle view images **6272**, **6274**, **6276**, **6278** of a surgical site at four separate times during the procedure, according to one aspect of the present disclosure. The sequence of images shows the creation of an overhead composite image in wide and narrow focus over time. A first image **6272** is a wide angle view of the end-effector **6234** clamping the vessel **6236** taken at an earlier time to (e.g., 09:35:09). A second image **6274** is another wide angle view of the end-effector **6234** clamping the vessel **6236** taken at the present time  $t_1$  (e.g., 09:35:13). A third image **6276** is a composite image of an overhead view of the end-effector **6234** clamping the vessel **6236** taken at present time  $t_1$ . The third image **6276** is displayed in the second window **6240** of the primary display **6200** of the surgical hub **206** as shown in FIG. **122**. A fourth image **6278** is a narrow angle view of the end-effector **6234** clamping the vessel **6236** at present time  $t_1$  (e.g., 09:35:13). The fourth image **6278** is the narrow angle view of the surgical site shown in the primary window **6230** of the primary display **6200** of the surgical hub **206** as shown in FIG. **122**.

#### Display of Instrument Specific Data Needed for Efficient Use of the End-Effector

[1115] In one aspect, the present disclosure provides a surgical hub display of instrument specific data needed for

efficient use of a surgical instrument, such as a surgical stapler. The techniques described herein may be applied to ultrasonic instruments, electrosurgical instruments, combination ultrasonic/electrosurgical instruments, and/or combination surgical stapler/electrosurgical instruments. In one aspect, a clamp time indicator based on tissue properties is shown on the display. In another aspect, a **3600** composite top visual view is shown on the display to avoid collateral structures as shown and described in connection with FIGS. **121-124** is incorporated herein by reference and, for conciseness and clarity of disclosure, the description of FIGS. **121-124** will not be repeated here.

[1116] In one aspect, the present disclosure provides a display of tissue creep to provide the user with in-tissue compression/tissue stability data and to guide the user making an appropriate choice of when to conduct the next instrument action. In one aspect, an algorithm calculates a constant advancement of a progressive time based feedback system related to the viscoelastic response of tissue. These and other aspects are described hereinbelow.

[1117] FIG. **125** is a graph **6280** of tissue creep clamp stabilization curves **6282**, **6284** for two tissue types, according to one aspect of the present disclosure. The clamp stabilization curves **6284**, **6284** are plotted as force-to-close (FTC) as a function of time, where FTC (N) is displayed along the vertical axis and Time,  $t$ , (Sec) is displayed along the horizontal axis. The FTC is the amount of force exerted to close the clamp arm on the tissue. The first clamp stabilization curve **6282** represents stomach tissue and the second clamp stabilization curve **6284** represents lung tissue. In one aspect, the FTC along the vertical axis is scaled from 0-180 N, and the horizontal axis is scaled from 0-5 Sec. As shown, the FTC as a different profile over a five second clamp stabilization period (e.g., as shown in FIG. **123**).

[1118] With reference to the first clamp stabilization curve **6282**, as the stomach tissue is clamped by the end-effector **6234**, the force-to-close (FTC) applied by the end-effector **6234** increases from 0 N to a peak force-to-close of ~180 N after ~1 Sec. While the end-effector **6234** remains clamped on the stomach tissue, the force-to-close decays and stabilizes to ~150 N overtime due to tissue creep.

[1119] Similarly, with reference to the second clamp stabilization curve **6284**, as the lung tissue is clamped by the end-effector **6234**, the force-to-close applied by the end-effector **6234** increases from 0 N to a peak force-to-close of ~90 N after just less than ~1 Sec. While the end-effector **6234** remains clamped on the lung tissue, the force-to-close decays and stabilizes to ~60 N over time due to tissue creep.

[1120] The end-effector **6234** clamp stabilization is monitored as described above in connection with FIGS. **122-124** and is displayed every second corresponding the sampling times  $t_1$ ,  $t_2$ ,  $t_3$ ,  $t_4$ ,  $t_5$  of the force-to-close to provide user feedback regarding the state of the clamped tissue. FIG. **125** shows an example of monitoring tissue stabilization for the lung tissue by sampling the force-to-close every second over a 5 seconds period. At each sample time  $t_1$ ,  $t_2$ ,  $t_3$ ,  $t_4$ ,  $t_5$ , the instrument **235** or the surgical hub **206** calculates a corresponding vector tangent **6288**, **6292**, **6294**, **6298**, **6302** to the second clamp stabilization curve **6284**. The vector tangent **6288**, **6292**, **6294**, **6298**, **6302** is monitored until its slope drops below a threshold to indicate that the tissue creep is complete and the tissue is ready to sealed and cut. As shown in FIG. **125**, the lung tissue is ready to be sealed and cut after ~5 Sec. clamp stabilization period, where a solid gray circle

is shown at sample time **6300**. As shown, the vector tangent **6302** is less than a predetermined threshold.

[1121] The equation of a vector tangent **6288**, **6292**, **6294**, **6298**, **6302** to the clamp stabilization curve **6284** may be calculated using differential calculus techniques, for example. In one aspect, at a given point on the clamp stabilization curve **6284**, the gradient of the curve **6284** is equal to the gradient of the tangent to the curve **6284**. The derivative (or gradient function) describes the gradient of the curve **6284** at any point on the curve **6284**. Similarly, it also describes the gradient of a tangent to the curve **6284** at any point on the curve **6284**. The normal to the curve **6284** is a line perpendicular to the tangent to the curve **6284** at any given point. To determine the equation of a tangent to a curve find the derivative using the rules of differentiation. Substitute the x coordinate (independent variable) of the given point into the derivative to calculate the gradient of the tangent. Substitute the gradient of the tangent and the coordinates of the given point into an appropriate form of the straight line equation. Make the y coordinate (dependent variable) the subject of the formula.

[1122] FIG. **126** is a graph **6310** of time dependent proportionate fill of a clamp force stabilization curve, according to one aspect of the present disclosure. The graph **6310** includes clamp stabilization curves **6312**, **6314**, **6316** for standard thick stomach tissue, thin stomach tissue, and standard lung tissue. The vertical axis represents FTC (N) scaled from 0-240 N and the horizontal axis represents Time, t, (Sec) scaled from 0-15 Sec. As shown, the standard thick stomach tissue curve **6316** is the default force decay stability curve. All three clamp stabilization curves **6312**, **6314**, **6316** FTC profiles reach a maximum force shortly after clamping on the tissue and then the FTC decreases over time until it eventually stabilizes due to the viscoelastic response of the tissue. As shown the standard lung tissue clamp stabilization curve **6312** stabilizes after a period of ~5 Sec., the thin stomach tissue clamp stabilization curve **6314** stabilizes after a period of ~10 Sec., and the thick stomach tissue clamp stabilization curve **6316** stabilizes after a period of ~15 Sec.

[1123] FIG. **127** is a graph **6320** of the role of tissue creep in the clamp force stabilization curve **6322**, according to one aspect of the present disclosure. The vertical axis represents force-to-close FTC (N) and the horizontal axis represents Time, t, (Sec) in seconds. Vector tangent angles  $d\theta_1$ ,  $d\theta_2$ , . . .  $d\theta_n$  are measured at each force-to-close sampling (to,  $t_1$ ,  $t_2$ ,  $t_3$ ,  $t_4$ , etc.) times. The vector tangent angle  $d\theta$  is used to determine when the tissue has reached the creep termination threshold, which indicates that the tissue has reached creep stability.

[1124] FIGS. **128A** and **128B** illustrate two graphs **6330**, **6340** for determining when the clamped tissue has reached creep stability, according to one aspect of the present disclosure. The graph **6330** in FIG. **128A** illustrates a curve **6332** that represents a vector tangent angle  $d\theta$  as a function of time. The vector tangent angle  $d\theta$  is calculated as discussed in FIG. **127**. The horizontal line **6334** is the tissue creep termination threshold. The tissue creep is deemed to be stable at the intersection **6336** of the vector tangent angle  $d\theta$  curve **6332** and the tissue creep termination threshold **6334**. The graph **6340** in FIG. **128B** illustrates a  $\Delta$ FTC curve **6342** that represents  $\Delta$ FTC as a function of time. The  $\Delta$ FTC curve **6342** illustrates the threshold **6344** to 100% complete

tissue creep stability meter. The tissue creep is deemed to be stable at the intersection **6346** of the  $\Delta$ FTC curve **6342** and the threshold **6344**.

#### Communication Techniques

[1125] With reference to FIGS. **1-11** to show interaction with an interactive surgical system **100** environment including a surgical hub **106**, **206**, and in particular, FIGS. **9-10**, in various aspects, the present disclosure provides communications techniques for exchanging information between an instrument **235**, or other modules, and the surgical hub **206**. In one aspect, the communications techniques include image fusion to place instrument status and analysis over a laparoscope image, such as a screen overlay of data, within and around the perimeter of an image presented on a surgical hub display **215**, **217**. In another aspect, the communication techniques include combining an intermediate short range wireless, e.g., Bluetooth, signal with the image, and in another aspect, the communication techniques include applying security and identification of requested pairing. In yet another aspect, the communication techniques include an independent interactive headset worn by a surgeon that links to the hub with audio and visual information that avoids the need for overlays, but allows customization of displayed information around periphery of view. Each of these communication techniques is discussed hereinbelow.

#### Screen Overlay of Data within and Around the Perimeter of the Displayed Image

[1126] In one aspect, the present disclosure provides image fusion allowing the overlay of the status of a device, the integration landmarks being used to interlock several images, and at least one guidance feature. In another aspect, the present disclosure provides a technique for screen overlay of data within and around the perimeter of displayed image. Radiographic integration may be employed for live internal sensing and pre-procedure overlay. Image fusion of one source may be superimposed over another. Image fusion may be employed to place instrument status and analysis on a medical imaging device (e.g., laparoscope, endoscope, thoracoscope, etc.) image. Image fusion allows the overlay of the status of a device or instrument, integration landmarks to interlock several images, and at least one guidance feature.

[1127] FIG. **129** illustrates an example of an augmented video image **6350** comprising a pre-operative video image **6352** augmented with data **6354**, **6356**, **6358** identifying displayed elements. An augmented reality vision system may be employed in surgical procedures to implement a method for augmenting data onto a pre-operative image **6352**. The method includes generating a pre-operative image **6352** of an anatomical section of a patient and generating an augmented video image of a surgical site within the patient. The augmented video image **6350** includes an image of at least a portion of a surgical tool **6354** operated by a user **6456**. The method further includes processing the pre-operative image **6352** to generate data about the anatomical section of the patient. The data includes a label **6358** for the anatomical section and a peripheral margin of at least a portion of the anatomical section. The peripheral margin is configured to guide a surgeon to a cutting location relative to the anatomical section, embedding the data and an identity of the user **6356** within the pre-operative image **6350** to

display an augmented video image **6350** to the user about the anatomical section of the patient. The method further includes sensing a loading condition on the surgical tool **6354**, generating a feedback signal based on the sensed loading condition, and updating, in real time, the data and a location of the identity of the user operating the surgical tool **6354** embedded within the augmented video image **6350** in response to a change in a location of the surgical tool **6354** within the augmented video image **6350**. Further examples are disclosed in U.S. Pat. No. 9,123,155, titled APPARATUS AND METHOD FOR USING AUGMENTED REALITY VISION SYSTEM IN SURGICAL PROCEDURES, which issued on Sep. 1, 2015, which is herein incorporated by reference in its entirety.

[1128] In another aspect, radiographic integration techniques may be employed to overlay the pre-operative image **6352** with data obtained through live internal sensing or pre-procedure techniques. Radiographic integration may include marker and landmark identification using surgical landmarks, radiographic markers placed in or outside the patient, identification of radio-opaque staples, clips or other tissue-fixed items. Digital radiography techniques may be employed to generate digital images for overlaying with a pre-operative image **6352**. Digital radiography is a form of X-ray imaging that employs a digital image capture device with digital X-ray sensors instead of traditional photographic film. Digital radiography techniques provide immediate image preview and availability for overlaying with the pre-operative image **6352**. In addition, special image processing techniques can be applied to the digital X-ray images to enhance the overall display quality of the image.

[1129] Digital radiography techniques employ image detectors that include flat panel detectors (FPDs), which are classified in two main categories indirect FPDs and direct FPDs. Indirect FPDs include amorphous silicon (a-Si) combined with a scintillator in the detector's outer layer, which is made from cesium iodide (CsI) or gadolinium oxy-sulfide (Gd<sub>2</sub>O<sub>2</sub>S), converts X-rays to light. The light is channeled through the a-Si photodiode layer where it is converted to a digital output signal. The digital signal is then read out by thin film transistors (TFTs) or fiber-coupled charge coupled devices (CCDs). Direct FPDs include amorphous selenium (a-Se) FPDs that convert X-ray photons directly into charge. The outer layer of a flat panel in this design is typically a high-voltage bias electrode. X-ray photons create electron-hole pairs in a-Se, and the transit of these electrons and holes depends on the potential of the bias voltage charge. As the holes are replaced with electrons, the resultant charge pattern in the selenium layer is read out by a TFT array, active matrix array, electrometer probes or micro plasma line addressing. Other direct digital detectors are based on CMOS and CCD technology. Phosphor detectors also may be employed to record the X-ray energy during exposure and is scanned by a laser diode to excite the stored energy which is released and read out by a digital image capture array of a CCD.

[1130] FIG. 130 is a logic flow diagram **6360** of a process depicting a control program or a logic configuration to display images, according to one aspect of the present disclosure. With reference also to FIGS. 1-11 to show interaction with an interactive surgical system **100** environment including a surgical hub **106**, **206**, the present disclosure provides, in one aspect, a surgical hub **206**, comprising a processor **244** and a memory **249** coupled to the processor

**244**. The memory **249** stores instructions executable by the processor **244** to receive **6362** first image data from a first image sensor, receive **6364** second image data from a second image sensor, and display **6366**, on a display **217** coupled to the surgical hub **206**, a first image corresponding to the first field of view and a second image corresponding to the second field of view. The first image data represents a first field of view and the second image data represents a second field of view.

[1131] In one aspect, the first field of view is a narrow angle field of view and the second field of view is a wide angle field of view. In another aspect, the memory **249** stores instructions executable by the processor **244** to augment the first image with the second image on the display. In another aspect, the memory **249** stores instructions executable by the processor **244** to fuse the first image and the second image into a third image and display a fused image on the display **217**. In another aspect, the fused image data comprises status information associated with a surgical device **235**, an image data integration landmark to interlock a plurality of images, and at least one guidance parameter. In another aspect, the first image sensor is the same as the same image sensor and wherein the first image data is captured as a first time and the second image data is captured at a second time.

[1132] In another aspect, the memory **249** stores instructions executable by the processor **244** to receive third image data from a third image sensor, wherein the third image data represents a third field of view, generate composite image data comprising the second and third image data, display the first image in a first window of the display, wherein the first image corresponds to the first image data, and display a third image in a second window of the display **215**, wherein the third image corresponds to the composite image data.

[1133] In another aspect, the memory **249** stores instructions executable by the processor **244** to receive third image data from a third image sensor, wherein the third image data represents a third field of view, fuse the second and third image data to generate fused image data, display the first image in a first window of the display **217**, wherein the first image corresponds to the first image data, and display a third image in a second window of the display **217**, wherein the third image corresponds to the fused image data.

#### Intermediate Short Range Wireless (e.g., Bluetooth) Signal Combiner

[1134] An intermediate short range wireless, e.g., Bluetooth, signal combiner may comprise a wireless heads-up display adapter placed into the communication path of the monitor to a laparoscope console allowing the surgical hub to overlay data onto the screen. Security and identification of requested pairing may augment the communication techniques.

[1135] FIG. 131 illustrates a communication system **6370** comprising an intermediate signal combiner **6372** positioned in the communication path between an imaging module **238** and a surgical hub display **217**, according to one aspect of the present disclosure. The signal combiner **6372** receives image data from an imaging module **238** in the form of short range wireless or wired signals. The signal combiner **6372** also receives audio and image data from a headset **6374** and combines the image data from the imaging module **238** with the audio and image data from the headset **6374**. The surgical hub **206** receives the combined data from the combiner **6372** and overlays the data provided to the display

217, where the overlaid data is displayed. The signal combiner 6372 may communicate with the surgical hub 206 via wired or wireless signals. The headset 6374 receives image data from an imaging device 6376 coupled to the headset 6374 and receives audio data from an audio device 6378 coupled to the headset 6374. The imaging device 6376 may be a digital video camera and the audio device 6378 may be a microphone. In one aspect, the signal combiner 6372 may be an intermediate short range wireless, e.g., Bluetooth, signal combiner. The signal combiner 6374 may comprise a wireless heads-up display adapter to couple to the headset 6374 placed into the communication path of the display 217 to a console allowing the surgical hub 206 to overlay data onto the screen of the display 217. Security and identification of requested pairing may augment the communication techniques. The imaging module 238 may be coupled to a variety of imaging devices such as an endoscope 239, laparoscope, etc., for example.

#### Independent Interactive Headset

[1136] FIG. 132 illustrates an independent interactive headset 6380 worn by a surgeon 6382 to communicate data to the surgical hub, according to one aspect of the present disclosure. Peripheral information of the independent interactive headset 6380 does not include active video. Rather, the peripheral information includes only device settings, or signals that do not have same demands of refresh rates. Interaction may augment the surgeon's 6382 information based on linkage with preoperative computerized tomography (CT) or other data linked in the surgical hub 206. The independent interactive headset 6380 can identify structure—ask whether instrument is touching a nerve, vessel, or adhesion, for example. The independent interactive headset 6380 may include pre-operative scan data, an optical view, tissue interrogation properties acquired throughout procedure, and/or processing in the surgical hub 206 used to provide an answer. The surgeon 6382 can dictate notes to the independent interactive headset 6380 to be saved with patient data in the hub storage 248 for later use in report or in follow up.

[1137] In one aspect, the independent interactive headset 6380 worn by the surgeon 6382 links to the surgical hub 206 with audio and visual information to avoid the need for overlays, and allows customization of displayed information around periphery of view. The independent interactive headset 6380 provides signals from devices (e.g., instruments), answers queries about device settings, or positional information linked with video to identify quadrant or position. The independent interactive headset 6380 has audio control and audio feedback from the headset 6380. The independent interactive headset 6380 is still able to interact with all other systems in the operating theater (e.g., operating room), and have feedback and interaction available wherever the surgeon 6382 is viewing.

#### Identification And Usage Recording

[1138] In one aspect, the present disclosure provides a display of the authenticity of reloads, modular components, or loading units. FIG. 133 illustrates a method 6390 for controlling the usage of a device 6392. A device 6392 is connected to an energy source 6394. The device 6392 includes a memory device 6396 that includes storage 6398 and communication 6400 devices. The storage 6398

includes data 6402 that may be locked data 6404 or unlocked data 6406. Additionally, the storage 6398 includes an error-detecting code 6408 such as a cyclic redundancy check (CRC) value and a sterilization indicator 6410. The energy source 6394 includes a reader 6412, display 6414, a processor 6416, and a data port 6418 that couples the energy source 6394 to a network 6420. The network 6420 is coupled to a central server 6422, which is coupled to a central database 6424. The network 6420 also is coupled to a reprocessing facility 6426. The reprocessing facility 6426 includes a reprocessing data reader/writer 6428 and a sterilizing device 6430.

[1139] The method comprises connecting the device to an energy source 6394. Data is read from a memory device 6396 incorporated in the device 6392. The data including one or more of a unique identifier (UID), a usage value, an activation value, a reprocessing value, or a sterilization indicator. The usage value is incremented when the device 6392 is connected to the energy source 6394. The activation value is incremented when the device 6392 is activated permitting energy to flow from the energy source 6394 to an energy consuming component of the device 6392. Usage of the device 6392 may be prevented if: the UID is on a list of prohibited UIDs, the usage value is not lower than a usage limitation value, the reprocessing value is equal to a reprocessing limitation value, the activation value is equal to an activation limitation value, and/or the sterilization indicator does not indicate that the device has been sterilized since its previous usage. Further examples are disclosed in U.S. Patent Application Publication No. 2015/0317899, titled SYSTEM AND METHOD FOR USING RFID TAGS TO DETERMINE STERILIZATION OF DEVICES, which published on Nov. 5, 2015, which is herein incorporated by reference in its entirety.

[1140] FIG. 134 provides a surgical system 6500 in accordance with the present disclosure and includes a surgical instrument 6502 that is in communication with a console 6522 or a portable device 6526 through a local area network 6518 or a cloud network 6520 via a wired or wireless connection. In various aspects, the console 6522 and the portable device 6526 may be any suitable computing device. The surgical instrument 6502 includes a handle 6504, an adapter 6508, and a loading unit 6514. The adapter 6508 releasably couples to the handle 6504 and the loading unit 6514 releasably couples to the adapter 6508 such that the adapter 6508 transmits a force from a drive shaft to the loading unit 6514. The adapter 6508 or the loading unit 6514 may include a force gauge (not explicitly shown) disposed therein to measure a force exerted on the loading unit 6514. The loading unit 6514 includes an end effector 6530 having a first jaw 6532 and a second jaw 6534. The loading unit 6514 may be an in-situ loaded or multi-firing loading unit (MFLU) that allows a clinician to fire a plurality of fasteners multiple times without requiring the loading unit 6514 to be removed from a surgical site to reload the loading unit 6514.

[1141] The first and second jaws 6532, 6534 are configured to clamp tissue therebetween, fire fasteners through the clamped tissue, and sever the clamped tissue. The first jaw 6532 may be configured to fire at least one fastener a plurality of times, or may be configured to include a replaceable multi-fire fastener cartridge including a plurality of fasteners (e.g., staples, clips, etc.) that may be fired more than one time prior to being replaced. The second jaw 6534 may

include an anvil that deforms or otherwise secures the fasteners about tissue as the fasteners are ejected from the multi-fire fastener cartridge.

[1142] The handle 6504 includes a motor that is coupled to the drive shaft to affect rotation of the drive shaft. The handle 6504 includes a control interface to selectively activate the motor. The control interface may include buttons, switches, levers, sliders, touchscreen, and any other suitable input mechanisms or user interfaces, which can be engaged by a clinician to activate the motor.

[1143] The control interface of the handle 6504 is in communication with a controller 6528 of the handle 6504 to selectively activate the motor to affect rotation of the drive shafts. The controller 6528 is disposed within the handle 6504 and is configured to receive input from the control interface and adapter data from the adapter 6508 or loading unit data from the loading unit 6514. The controller 6528 analyzes the input from the control interface and the data received from the adapter 6508 and/or loading unit 6514 to selectively activate the motor. The handle 6504 may also include a display that is viewable by a clinician during use of the handle 6504. The display is configured to display portions of the adapter or loading unit data before, during, or after firing of the instrument 6502.

[1144] The adapter 6508 includes an adapter identification device 6510 disposed therein and the loading unit 6514 includes a loading unit identification device 6516 disposed therein. The adapter identification device 6510 is in communication with the controller 6528, and the loading unit identification device 6516 is in communication with the controller 6528. It will be appreciated that the loading unit identification device 6516 may be in communication with the adapter identification device 6510, which relays or passes communication from the loading unit identification device 6516 to the controller 6528.

[1145] The adapter 6508 may also include a plurality of sensors 6512 (one shown) disposed thereabout to detect various conditions of the adapter 6508 or of the environment (e.g., if the adapter 6508 is connected to a loading unit, if the adapter 6508 is connected to a handle, if the drive shafts are rotating, the torque of the drive shafts, the strain of the drive shafts, the temperature within the adapter 6508, a number of firings of the adapter 6508, a peak force of the adapter 6508 during firing, a total amount of force applied to the adapter 6508, a peak retraction force of the adapter 6508, a number of pauses of the adapter 6508 during firing, etc.). The plurality of sensors 6512 provides an input to the adapter identification device 6510 in the form of data signals. The data signals of the plurality of sensors 6512 may be stored within, or be used to update the adapter data stored within, the adapter identification device 6510. The data signals of the plurality of sensors 6512 may be analog or digital. The plurality of sensors 6512 may include a force gauge to measure a force exerted on the loading unit 6514 during firing.

[1146] The handle 6504 and the adapter 6508 are configured to interconnect the adapter identification device 6510 and the loading unit identification device 6516 with the controller 6528 via an electrical interface. The electrical interface may be a direct electrical interface (i.e., include electrical contacts that engage one another to transmit energy and signals therebetween). Additionally or alternatively, the electrical interface may be a non-contact electrical interface to wirelessly transmit energy and signals therebe-

tween (e.g., inductively transfer). It is also contemplated that the adapter identification device 6510 and the controller 6528 may be in wireless communication with one another via a wireless connection separate from the electrical interface.

[1147] The handle 6504 includes a transmitter 6506 that is configured to transmit instrument data from the controller 6528 to other components of the system 6500 (e.g., the LAN 6518, the cloud 6520, the console 6522, or the portable device 6526). The transmitter 6506 also may receive data (e.g., cartridge data, loading unit data, or adapter data) from the other components of the system 6500. For example, the controller 6528 may transmit instrument data including a serial number of an attached adapter (e.g., adapter 6508) attached to the handle 6504, a serial number of a loading unit (e.g., loading unit 6514) attached to the adapter, and a serial number of a multi-fire fastener cartridge (e.g., multi-fire fastener cartridge), loaded into the loading unit, to the console 6528. Thereafter, the console 6522 may transmit data (e.g., cartridge data, loading unit data, or adapter data) associated with the attached cartridge, loading unit, and adapter, respectively, back to the controller 6528. The controller 6528 can display messages on the local instrument display or transmit the message, via transmitter 6506, to the console 6522 or the portable device 6526 to display the message on the display 6524 or portable device screen, respectively.

#### Multi-Functional Surgical Control System and Switching Interface for Verbal Control of Imaging Device

[1148] FIG. 135 illustrates a verbal AESOP camera positioning system. Further examples are disclosed in U.S. Pat. No. 7,097,640, titled MULTI-FUNCTIONAL SURGICAL CONTROL SYSTEM AND SWITCHING INTERFACE, which issued on Aug. 29, 2006, which is herein incorporated by reference in its entirety. FIG. 135 shows a surgical system 6550 that may be coupled to surgical hub 206, described in connection with FIGS. 1-11. The system 6550 allows a surgeon to operate a number of different surgical devices 6552, 6554, 6556, and 6558 from a single input device 6560. Providing a single input device reduces the complexity of operating the various devices and improves the efficiency of a surgical procedure performed by a surgeon. The system 6550 may be adapted and configured to operate a positioning system for an imaging device such as a camera or endoscope using verbal commands.

[1149] The surgical device 6552 may be a robotic arm which can hold and move a surgical instrument. The arm 6552 may be a device such as that sold by Computer Motion, Inc. of Goleta, Calif. under the trademark AESOP, which is an acronym for Automated Endoscopic System for Optimal Positioning. The arm 6552 is commonly used to hold and move an endoscope within a patient. The system 6550 allows the surgeon to control the operation of the robotic arm 6552 through the input device 6560.

[1150] The surgical device 6554 may be an electrocautery device. Electrocautery devices typically have a bi-polar tip which carries a current that heats and denatures tissue. The device is typically coupled to an on-off switch to actuate the device and heat the tissue. The electrocautery device may also receive control signals to vary its power output. The system 6550 allows the surgeon to control the operation of the electrocautery device through the input device 6560.

[1151] The surgical device **6556** may be a laser. The laser **6556** may be actuated through an on-off switch. Additionally, the power of the laser **6556** may be controlled by control signals. The system **6550** allows the surgeon to control the operation of the laser **6556** through the input device **6560**.

[1152] The device **6558** may be an operating table. The operating table **6558** may contain motors and mechanisms which adjust the position of the table. The present invention allows the surgeon to control the position of the table **6558** through the input device **6560**. Although four surgical devices **6552**, **6554**, **6556**, and **6558** are described, it is to be understood that other functions within the operating room may be controlled through the input device **6560**. By way of example, the system **6560** may allow the surgeon to control the lighting and temperature of the operating room through the input device **6560**.

[1153] The input device **6560** may be a foot pedal which has a plurality of buttons **6562**, **6564**, **6565**, **6566**, and **6568** that can be depressed by the surgeon. Each button is typically associated with a specific control command of a surgical device. For example, when the input device **6560** is controlling the robotic arm **6552**, depressing the button **6562** may move the arm in one direction and depressing the button **6566** may move the arm in an opposite direction. Likewise, when the electrocautery device **6554** or the laser **6556** is coupled to the input device **6560**, depressing the button **6568** may energize the devices, and so forth and so on. Although a foot pedal is shown and described, it is to be understood that the input device **6560** may be a hand controller, a speech interface which accepts voice commands from the surgeon, a cantilever pedal or other input devices which may be well known in the art of surgical device control. Using the speech interface, the surgeon is able to position a camera or endoscope connected to the robotic arm **6552** using verbal commands. The imaging device, such as a camera or endoscope, may be coupled to the robotic arm **6552** positioning system that be controlled through the system **6550** using verbal commands.

[1154] The system **6550** has a switching interface **6570** which couples the input device **6560** to the surgical devices **6552**, **6554**, **6556**, and **6558**. The interface **6570** has an input channel **6572** which is connected to the input device **6560** by a bus **6574**. The interface **6570** also has a plurality of output channels **6576**, **6578**, **6580**, and **6582** that are coupled to the surgical devices by busses **6584**, **6586**, **6588**, **6590**, **6624**, **6626**, **6628** and which may have adapters or controllers disposed in electrical communication therewith and therebetween. Such adapters and controllers will be discussed in more detail hereinbelow.

[1155] Because each device **6552**, **6554**, **6556**, **6558** may require specifically configured control signals for proper operation, adapters **6620**, **6622** or a controller **6618** may be placed intermediate and in electrical communication with a specific output channel and a specific surgical device. In the case of the robotic arm system **6552**, no adapter is necessary and as such, the robotic arm system **6552** may be in direct connection with a specific output channel. The interface **6570** couples the input channel **6572** to one of the output channels **6576**, **6578**, **6580**, and **6582**.

[1156] The interface **6570** has a select channel **6592** which can switch the input channel **6572** to a different output channel **6576**, **6578**, **6580**, or **6582** so that the input device **6560** can control any of the surgical devices. The interface

**6570** may be a multiplexor circuit constructed as an integrated circuit and placed on an ASIC. Alternatively, the interface **6570** may be a plurality of solenoid actuated relays coupled to the select channel by a logic circuit. The interface **6570** switches to a specific output channel in response to an input signal or switching signal applied on the select channel **6592**.

[1157] As depicted in FIG. 135, there may be several inputs to the select channel **6592**. Such inputs originate from the foot pedal **6560**, the speech interface **6600** and the CPU **6602**. The interface **6570** may have a multiplexing unit such that only one switching signal may be received at the select channel **6592** at any one time, thus ensuring no substantial hardware conflicts. The prioritization of the input devices may be configured so the foot pedal has highest priority followed by the voice interface and the CPU. This is intended for example as the prioritization scheme may be employed to ensure the most efficient system. As such other prioritization schemes may be employed. The select channel **6592** may sequentially connect the input channel to one of the output channels each time a switching signal is provided to the select channel **6592**. Alternatively, the select channel **6592** may be addressable so that the interface **6570** connects the input channel to a specific output channel when an address is provided to the select channel **6592**. Such addressing is known in the art of electrical switches.

[1158] The select channel **6592** may be connected by line **6594** to a dedicated button **6596** on the foot pedal **6560**. The surgeon can switch surgical devices by depressing the button **6596**. Alternatively, the select channel **6592** may be coupled by line **6598** to a speech interface **6600** which allows the surgeon to switch surgical devices with voice commands.

[1159] The system **6550** may have a central processing unit (CPU) **6602** which receives input signals from the input device **6560** through the interface **6570** and a bus **6585**. The CPU **6602** receives the input signals, and can ensure that no improper commands are being input at the controller. If this occurs, the CPU **6602** may respond accordingly, either by sending a different switching signal to select channel **6592**, or by alerting the surgeon via a video monitor or speaker.

[1160] The CPU **6602** can also provide output commands for the select channel **6592** on the bus **6608** and receives input commands from the speech interface **6600** on the same bi-directional bus **6608**. The CPU **6602** may be coupled to a monitor **6610** and/or a speaker **6612** by buses **6614** and **6616**, respectively. The monitor **6610** may provide a visual indication of which surgical device is coupled to the input device **6560**. The monitor may also provide a menu of commands which can be selected by the surgeon either through the speech interface **6600** or button **6596**. Alternatively, the surgeon could switch to a surgical device by selecting a command through a graphic user interface. The monitor **6610** may also provide information regarding improper control signals sent to a specific surgical device **6552**, **6554**, **6556**, **6558** and recognized by the CPU **6602**. Each device **6552**, **6554**, **6556**, **6558** has a specific appropriate operating range, which is well known to the skilled artisan. As such, the CPU **6602** may be programmed to recognize when the requested operation from the input device **6560** is inappropriate and will then alert the surgeon either visually via the monitor **6610** or audibly via the speaker **6612**. The speaker **6612** may also provide an audio indication of which surgical device is coupled to the input device **6560**.

[1161] The system 6550 may include a controller 6618 which receives the input signals from the input device 6560 and provides corresponding output signals to control the operating table 6558. Likewise, the system may have adapters 6620, 6622 which provide an interface between the input device 6560 and the specific surgical instruments connected to the system.

[1162] In operation, the interface 6570 initially couples the input device 6560 to one of the surgical devices. The surgeon can control a different surgical device by generating an input command that is provided to the select channel 6592. The input command switches the interface 6570 so that the input device 6560 is coupled to a different output channel and corresponding surgical device or adapter. What is thus provided is an interface 6570 that allows a surgeon to select, operate and control a plurality of different surgical devices through a common input device 6560.

[1163] FIG. 136 illustrates a multi-functional surgical control system 6650 and switching interface for virtual operating room integration. A virtual control system for controlling surgical equipment in an operating room while a surgeon performs a surgical procedure on a patient, comprising: a virtual control device including an image of a control device located on a surface and a sensor for interrogating contact interaction of an object with the image on the surface, the virtual control device delivering an interaction signal indicative of the contact interaction of the object with the image; and a system controller connected to receive the interaction signal from the virtual control device and to deliver a control signal to the surgical equipment in response to the interaction signal to control the surgical equipment in response to the contact interaction of the object with the image. Further examples are disclosed in U.S. Pat. No. 7,317,955, titled VIRTUAL OPERATING ROOM INTEGRATION, which issued on Jan. 8, 2008, which is herein incorporated by reference in its entirety.

[1164] As shown in FIG. 136, communication links 6674 are established between the system controller 6676 and the various components and functions of the virtual control system 6650. The communication links 6674 are preferably optical paths, but the communication links may also be formed by radio frequency transmission and reception paths, hardwired electrical connections, or combinations of optical, radio frequency and hardwired connection paths as may be appropriate for the type of components and functions obtained by those components. The arrows at the ends of the links 6674 represent the direction of primary information flow.

[1165] The communication links 6674 with the surgical equipment 6652, a virtual control panel 6556, a virtual foot switch 6654 and patient monitoring equipment 6660 are bidirectional, meaning that the information flows in both directions through the links 6674 connecting those components and functions. For example, the system controller 6676 supplies signals which are used to create a control panel image from the virtual control panel 6656 and a foot switch image from the virtual foot switch 6654. The virtual control panel 6656 and the virtual foot switch 6654 supply information to the system controller 6676 describing the physical interaction of the surgeon's finger and foot relative to a projected control panel image and the projected foot switch image. The system controller 6676 responds to the information describing the physical interaction with the projected image, and supplies control signals to the surgical

equipment 6652 and patient monitoring equipment 6660 to control functionality of those components in response to the physical interaction information. The control, status and functionality information describing the surgical equipment 6652 and patient monitoring equipment 6660 flows to the system controller 6676, and after that information is interpreted by the system controller 6676, it is delivered to a system display 6670, a monitor 6666, and/or a heads up display 6668 for presentation.

[1166] The communication links 6674 between the system controller 6676 and the system display 6670, the heads up display 6668, the monitor 6666, a tag printer 6658 and output devices 6664 are all uni-directional, meaning that the information flows from the system controller 6676 to those components and functions. In a similar manner, the communication links 6674 between the system controller 6676 and a scanner 6672 and the input devices 6662 are also unidirectional, but the information flows from the components 6662, 6672 to the system controller 6676. In certain circumstances, certain control and status information may flow between the system controller 6676 and the components 6658, 6660, 6662, 6664, 6666, 6668, 6670, 6672 in order to control the functionality of the those components.

[1167] Each communication link 6674 preferably has a unique identity so that the system controller 6676 can individually communicate with each of the components of the virtual control system 6650. The unique identity of each communication link is preferable when some or all of the communication links 6674 are through the same medium, as would be the case of optical and radio frequency communications. The unique identity of each communication link 6674 assures that the system controller 6676 has the ability to exercise individual control over each of the components and functions on a very rapid and almost simultaneous manner. The unique identity of each communication link 6674 can be achieved by using different frequencies for each communication link 6674 or by using unique address and identification codes associated with the communications transferred over each communication link 6674.

[1168] In one aspect, the present disclosure provides illustrates a surgical communication and control headset that interfaces with the surgical hub 206 described in connection with FIGS. 1-11. Further examples are disclosed in U.S. Patent Application Publication No. 2009/0046146, titled SURGICAL COMMUNICATION AND CONTROL SYSTEM, which published on Feb. 19, 2009, which is herein incorporated by reference in its entirety. FIG. 137 illustrates a diagram 6680 of a beam source and combined beam detector system utilized as a device control mechanism in an operating theater. The system 6680 is configured and wired to allow for device control with the overlay generated on the primary procedural display. The footswitch shows a method to allow the user to click on command icons that would appear on the screen while the beam source is used to aim at the particular desired command icon to be clicked. The control system graphic user interface (GUI) and device control processor communicate and parameters are changed using the system. The system 6680 includes a display 6684 coupled to a beam detecting sensor 6682 and a head mounted source 6686. The beam detecting sensor 6682 is in communication with a control system GUI overlay processor and beam source processor 6688. The surgeon operates a footswitch 6692 or other adjunctive switch, which provides a signal to a device control interface unit 6694.



[1169] The system **6680** will provide a means for a sterile clinician to control procedural devices in an easy and quick, yet hands free and centralized fashion. The ability to maximize the efficiency of the operation and minimize the time a patient is under anesthesia is important to the best patient outcomes. It is common for surgeons, cardiologists or radiologists to verbally request adjustments be made to certain medical devices and electronic equipment used in the procedure outside the sterile field. It is typical that he or she must rely on another staff member to make the adjustments he or she needs to settings on devices such as cameras, bovie, surgical beds, shavers, insufflators, injectors, to name a few. In many circumstances, having to command a staff member to make a change to a setting can slow down a procedure because the non-sterile staff member is busy with another task. The sterile physician cannot adjust non-sterile equipment without compromising sterility, so he or she must often wait for the non-sterile staff member to make the requested adjustment to a certain device before resuming the procedure.

[1170] The system **6680** allows a user to use a beam source and beam detector to regenerate a pointer overlay coupled with a GUI and a concurrent switching method (i.e., a foot switch, etc.) to allow the clinician to click through commands on the primary display. In one aspect, a GUI could appear on the procedural video display when activated, such as when the user tilts his or her head twice to awaken it or steps on a foot switch provided with the system. Or it is possible that a right head tilt wakes up the system, and a left head tilt simply activates the beam source. When the overlay (called device control GUI overlay) appears on the screen it shows button icons representing various surgical devices and the user can use the beam source, in this case a laser beam, to aim at the button icons. Once the laser is over the proper button icon, a foot switch, or other simultaneous switch method can be activated, effectively acting like a mouse click on a computer. For example a user can “wake up” the system, causing a the device control GUI overlay to pop up that lists button icons on the screen, each one labeled as a corresponding procedural medical device. The user can point the laser at the correct box or device and click a foot pedal (or some other concurrent control-like voice control, waistband button, etc.) to make a selection, much like clicking a mouse on a computer. The sterile physician can then select “insufflator, for example” The subsequent screen shows arrow icons that can be clicked for various settings for the device that need to be adjusted (pressure, rate, etc.). In one iteration, the user can then can point the laser at the up arrow and click the foot pedal repeatedly until the desired setting is attained.

[1171] In one aspect, components of the system **6680** could be coupled with existing robotic endoscope holders to “steer” a rigid surgical endoscopic camera by sending movement commands to the robotic endoscope holding arm (provided separately, i.e., AESOP by Computer Motion). The endoscope is normally held by an assistant nurse or resident physician. There are robotic and mechanical scope holders currently on the market and some have even had been introduced with voice control. However, voice control systems have often proven cumbersome, slow and inaccurate. This aspect would employ a series of software and hardware components to allow the overlay to appear as a crosshair on the primary procedural video screen. The user could point the beam source at any part of the quadrant and

click a simultaneous switch, such as a foot pedal, to send movement commands to the existing robotic arm, which, when coupled with the secondary trigger (i.e., a foot switch, waist band switch, etc.) would send a command to adjust the arm in minute increments in the direction of the beam source. It could be directed by holding down the secondary trigger until the desired camera angle and position is achieved and then released. This same concept could be employed for surgical bed adjustments by having the overlay resemble the controls of a surgical bed. The surgical bed is commonly adjusted during surgery to allow better access to the anatomy. Using the combination of the beam source, in this case a laser, a beam detecting sensor such as a camera, a control system GUI overlay processing unit and beam source processor, and a device control interface unit, virtually any medical device could be controlled through this system.

[1172] Control codes would be programmed into the device control interface unit, and most devices can be connected using an RS-232 interface, which is a standard for serial binary data signals connecting between a DTE (Data Terminal Equipment) and a DCE (Data Circuit-terminating Equipment). The present invention while described with reference to application in the medical field can be expanded/modified for use in other fields. Another use of this invention could be in helping those who are without use of their hands due to injury or handicap or for professions where the hands are occupied and hands free interface is desired.

Surgical Hub with Direct Interface Control with  
Secondary Surgeon Display Units Designed to be  
within the Sterile Field and Accessible for Input  
and Display by the Surgeon

[1173] In one aspect, the surgical hub **206** provides a secondary user interface that enables display and control of surgical hub **206** functions from within the sterile field. The secondary display could be used to change display locations, what information is displayed where, pass off control of specific functions or devices.

[1174] During a surgical procedure, the surgeon may not have a user interface device accessible for interactive input by the surgeon and display within the sterile field. Thus, the surgeon cannot interface with the user interface device and the surgical hub from within the sterile field and cannot control other surgical devices through the surgical hub from within the sterile field.

[1175] One solution provides a display unit designed to be used within the sterile field and accessible for input and display by the surgeon to allow the surgeon to have interactive input control from the sterile field to control other surgical devices coupled to the surgical hub. The display unit is sterile and located within the sterile field to allow the surgeons to interface with the display unit and the surgical hub to directly interface and configure instruments as necessary without leaving the sterile field. The display unit is a master device and may be used for display, control, interchanges of tool control, allowing feeds from other surgical hubs without the surgeon leaving the sterile field.

[1176] In one aspect, the present disclosure provides a control unit, comprising an interactive touchscreen display, an interface configured to couple the interactive touchscreen display to a surgical hub, a processor, and a memory coupled to the processor. The memory stores instructions executable

by the processor to receive input commands from the interactive touchscreen display located inside a sterile field and transmits the input commands to a surgical hub to control devices coupled to the surgical hub located outside the sterile field.

[1177] In another aspect, the present disclosure provides a control unit, comprising an interactive touchscreen display, an interface configured to couple the interactive touchscreen display to a surgical hub, and a control circuit configured to receive input commands from the interactive touchscreen display located inside a sterile field and transmit the input commands to a surgical hub to control devices coupled to the surgical hub located outside the sterile field.

[1178] In another aspect, the present disclosure provides a non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to receive input commands from an interactive touchscreen display located inside a sterile field and transmit the input commands to a surgical hub through an interface configured to couple the interactive touchscreen display to the surgical hub to control devices coupled to the surgical hub located outside the sterile field.

[1179] Providing a display unit designed to be used within the sterile field and accessible for input and display by the surgeon provides the surgeon interactive input control from the sterile field to control other surgical devices coupled to the surgical hub.

[1180] This display unit within the sterile field is sterile and allows the surgeons to interface with it and the surgical hub. This gives the surgeon control of the instruments coupled to the surgical hub and allows the surgeon to directly interface and configure the instruments as necessary without leaving the sterile field. The display unit is a master device and may be used for display, control, interchanges of tool control, allowing feeds from other surgical hubs without the surgeon leaving the sterile field.

[1181] In various aspects, the present disclosure provides a secondary user interface to enable display and control of surgical hub functions from within a sterile field. This control could be a display device like an I-pad, e.g., a portable interactive touchscreen display device configured to be introduced into the operating theater in a sterile manner. It could be paired like any other device or it could be location sensitive. The display device would be allowed to function in this manner whenever the display device is placed over a specific location of the draped abdomen of the patient during a surgical procedure. In other aspects, the present disclosure provides a smart retractor and a smart sticker. These and other aspects are described hereinbelow.

[1182] In one aspect, the present disclosure provides a secondary user interface to enable display and control of surgical hub functions from within the sterile field. In another aspect, the secondary display could be used to change display locations, determine what information and where the information is displayed, and pass off control of specific functions or devices.

[1183] There are four types of secondary surgeon displays in two categories. One type of secondary surgeon display units is designed to be used within the sterile field and accessible for input and display by the surgeon within the sterile field interactive control displays. Sterile field interactive control displays may be shared or common sterile field input control displays.

[1184] A sterile field display may be mounted on the operating table, on a stand, or merely laying on the abdomen or chest of the patient. The sterile field display is sterile and allows the surgeons to interface with the sterile field display and the surgical hub. This gives the surgeon control of the system and allows them to directly interface and configure the sterile field display as necessary. The sterile field display may be configured as a master device and may be used for display, control, interchanges of tool control, allowing feeds from other surgical hubs, etc.

[1185] In one aspect, the sterile field display may be employed to re-configure the wireless activation devices within the operating theater (OR) and their paired energy device if a surgeon hands the device to another. FIGS. 138A-138E illustrate various types of sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 according to various aspects of the present disclosure. Each of the disclosed sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 comprise at least one touchscreen 6701, 6704/6706, 6709, 6713, 6716 input/output device layered on the top of an electronic visual display of an information processing system. The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 may include batteries as a power source. Some include a cable 6710 to connect to a separate power source or to recharge the batteries. A user can give input or control the information processing system through simple or multi-touch gestures by touching the touchscreen 6701, 6704/6706, 6709, 6713, 6716 with a stylus, one or more fingers, or a surgical tool. The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 may be used to re-configure wireless activation devices within the operating theater and a paired energy device if a surgeon hands the device to another surgeon. The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 may be used to accept consult feeds from another operating theater where it would then configure a portion of the operating theater screens or all of them to mirror the other operating theater so the surgeon is able to see what is needed to help. The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 are configured to communicate with the surgical hub 206. Accordingly, the description of the surgical hub 206 discussed in connection with FIGS. 1-11 is incorporated in this section by reference.

[1186] FIG. 138A illustrates a single zone sterile field control and data input console 6700, according to one aspect of the present disclosure. The single zone console 6700 is configured for use in a single zone within a sterile field. Once deployed in a sterile field, the single zone console 6700 can receive touchscreen inputs from a user in the sterile field. The touchscreen 6701 enables the user to interact directly with what is displayed, rather than using a mouse, touchpad, or other such devices (other than a stylus or surgical tool). The single zone console 6700 includes wireless communication circuits to communicate wirelessly to the surgical hub 206.

[1187] FIG. 138B illustrates a multi zone sterile field control and data input console 6702, according to one aspect of the present disclosure. The multi zone console 6702 comprises a first touchscreen 6704 to receive an input from a first zone of a sterile field and a second touchscreen 6706 to receive an input from a second zone of a sterile field. The multi zone console 6702 is configured to receive inputs from multiple users in a sterile field. The multi zone console 6702

includes wireless communication circuits to communicate wirelessly to the surgical hub 206. Accordingly, the multi zone sterile field control and data input console 6702 comprises an interactive touchscreen display with multiple input and output zones.

[1188] FIG. 138C illustrates a tethered sterile field control and data input console 6708, according to one aspect of the present disclosure. The tethered console 6708 includes a cable 6710 to connect the tethered console 6708 to the surgical hub 206 via a wired connection. The cable 6710 enables the tethered console 6708 to communicate over a wired link in addition to a wireless link. The cable 6710 also enables the tethered console 6708 to connect to a power source for powering the console 6708 and/or recharging the batteries in the console 6708.

[1189] FIG. 138D illustrates a battery operated sterile field control and data input console 6712, according to one aspect of the present disclosure. The sterile field console 6712 is battery operated and includes wireless communication circuits to communicate wirelessly with the surgical hub 206. In particular, in one aspect, the sterile field console 6712 is configured to communicate with any of the modules coupled to the hub 206 such as the generator module 240. Through the sterile field console 6712, the surgeon can adjust the power output level of a generator using the touchscreen 6713 interface. One example is described below in connection with FIG. 138E.

[1190] FIG. 138E illustrates a battery operated sterile field control and data input console 6714, according to one aspect of the present disclosure. The sterile field console 6714 includes a user interface displayed on the touchscreen of a generator. The surgeon can thus control the output of the generator by touching the up/down arrow icons 6718A, 6718B that increase/decrease the power output of the generator module 240. Additional icons 6719 enable access to the generator module settings 6174, volume 6178 using the +/- icons, among other features directly from the sterile field console 6714. The sterile field console 6714 may be employed to adjust the settings or reconfigure other wireless activations devices or modules coupled to the hub 206 within the operating theater and their paired energy device when the surgeon hands the sterile field console 6714 to another.

[1191] FIGS. 139A-139B illustrate a sterile field console 6700 in use in a sterile field during a surgical procedure, according to one aspect of the present disclosure. FIG. 139A shows the sterile field console 6714 positioned in the sterile field near two surgeons engaged in an operation. In FIG. 139B, one of the surgeons is shown tapping the touchscreen 6701 of the sterile field console with a surgical tool 6722 to adjust the output of a modular device coupled to the surgical hub 206, reconfigure the modular device, or an energy device paired with the modular device coupled to the surgical hub 206.

[1192] In another aspect, the sterile field display may be employed to accept consult feeds from another operating room (OR), such as another operating theater or surgical hub 206, where it would then configure a portion of the OR screens or all of them to mirror the other ORs so the surgeon could see what is needed to help. FIG. 140 illustrates a process 6750 for accepting consult feeds from another operating room, according to one aspect of the present disclosure. The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 shown in FIGS. 138A-138E,

139A-139B may be used as an interact-able scalable secondary display allowing the surgeon to overlay other feeds or images from laser Doppler image scanning arrays or other image sources. The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 may be used to call up a pre-operative scan or image to review. Laser Doppler techniques are described in U.S. Provisional Patent Application No. 62/611,341, filed Dec. 28, 2017, and titled INTERACTIVE SURGICAL PLATFORM, which is incorporated herein by reference in its entirety.

[1193] It is recognized that the tissue penetration depth of light is dependent on the wavelength of the light used. Thus, the wavelength of the laser source light may be chosen to detect particle motion (such a blood cells) at a specific range of tissue depth. A laser Doppler employs means for detecting moving particles such as blood cells based at a variety of tissue depths based on the laser light wavelength. A laser source may be directed to a surface of a surgical site. A blood vessel (such as a vein or artery) may be disposed within the tissue at some depth  $\delta$  from the tissue surface. Red laser light (having a wavelength in the range of about 635 nm to about 660 nm) may penetrate the tissue to a depth of about 1 mm. Green laser light (having a wavelength in the range of about 520 nm to about 532 nm) may penetrate the tissue to a depth of about 2-3 mm. Blue laser light (having a wavelength in the range of about 405 nm to about 445 nm) may penetrate the tissue to a depth of about 4 mm or greater. A blood vessel may be located at a depth of about 2-3 mm below the tissue surface. Red laser light will not penetrate to this depth and thus will not detect blood cells flowing within this vessel. However, both green and blue laser light can penetrate this depth. Therefore, scattered green and blue laser light from the blood cells will result in an observed Doppler shift in both the green and blue.

[1194] In some aspects, a tissue may be probed by red, green, and blue laser illumination in a sequential manner and the effect of such illumination may be detected by a CMOS imaging sensor over time. It may be recognized that sequential illumination of the tissue by laser illumination at differing wavelengths may permit a Doppler analysis at varying tissue depths over time. Although red, green, and blue laser sources may be used to illuminate the surgical site, it may be recognized that other wavelengths outside of visible light (such as in the infrared or ultraviolet regions) may be used to illuminate the surgical site for Doppler analysis. The imaging sensor information may be provided to the sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714.

[1195] The sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 provide access to past recorded data. In one operating theater designated as OR1, the sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 may be configured as "consultants" and to erase all data when the consultation is complete. In another operating theater designated as OR3 (operating room 3), the sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714 may be configured as a "consultees" and are configured to record all data received from operating theater OR1 (operating room 1) sterile field control and data input consoles 6700, 6702, 6708, 6712, 6714. These configurations are summarized in TABLE 2 below:

TABLE 2

Sterile Field Control And Data Input Console In OR1	Sterile Field Control And Data Input Console In OR3
Access to past recorded data	
OR1 Consultant	OR 3 Consultee
Erase data when done	Record all data

[1196] In one implementation of the process 6750, operating theater OR1 receives a consult request from OR3. Data is transferred to the OR1 sterile field control and data input console 6700, for example. The data is temporarily stored 6754. The data is backed up in time and the OR1 view 6756 of the temporary data begins on the OR1 sterile field control and data input console 6700 touchscreen 6701. When the view is complete, the data is erased 6758 and control returns 6760 to OR1. The data is then erased 6762 from the OR1 sterile field control and data input console 6700 memory.

[1197] In yet another aspect, the sterile field display may be employed as an interactable scalable secondary display allowing the surgeon to overlay other feeds or images like laser Doppler scanning arrays. In yet another aspect, the sterile field display may be employed to call up a pre-operative scan or image to review. Once vessel path and depth and device trajectory are estimated, the surgeon employs a sterile field interactable scalable secondary display allowing the surgeon to overlay other feeds or images.

[1198] FIG. 141 is a diagram 6770 that illustrates a technique for estimating vessel path, depth, and device trajectory. Prior to dissecting a vessel 6772, 6774 located below the surface of the tissue 6775 using a standard approach, the surgeon estimates the path and depth of the vessel 6772, 6774 and a trajectory 6776 of a surgical device 6778 will take to reach the vessel 6772, 6774. It is often difficult to estimate the path and depth 6776 of a vessel 6772, 6774 located below the surface of the tissue 6775 because the surgeon cannot accurately visualize the location of the vessel 6772, 6774 path and depth 6776.

[1199] FIGS. 142A-142D illustrate multiple real time views of images of a virtual anatomical detail for dissection including perspective views (FIGS. 142A, 142C) and side views (FIGS. 142B, 142D). The images are displayed on a sterile field display of tablet computer or sterile field control and data input console employed as an interactable scalable secondary display allowing the surgeon to overlay other feeds or images, according to one aspect of the present disclosure. The images of the virtual anatomy enable the surgeon to more accurately predict the path and depth of a vessel 6772, 6774 located below the surface of the tissue 6775 as shown in FIG. 141 and the best trajectory 6776 of the surgical device 6778.

[1200] FIG. 142A is a perspective view of a virtual anatomy 6780 displayed on a tablet computer or sterile field control and data input console. FIG. 142B is a side view of the virtual anatomy 6780 shown in FIG. 142A, according to one aspect of the present disclosure. With reference to FIGS. 142A-142B, in one aspect, the surgeon uses a smart surgical device 6778 and a tablet computer to visualize the virtual anatomy 6780 in real time and in multiple views. The three dimensional perspective view includes a portion of tissue 6775 in which the vessels 6772, 6774 are located below surface. The portion of tissue is overlaid with a grid 6786 to enable the surgeon to visualize a scale and gauge the path

and depth of the vessels 6772, 6774 at target locations 6782, 6784 each marked by an X. The grid 6786 also assists the surgeon determine the best trajectory 6776 of the surgical device 6778. As illustrated, the vessels 6772, 6774 have an unusual vessel path.

[1201] FIG. 142C illustrates a perspective view of the virtual anatomy 6780 for dissection, according to one aspect of the present disclosure. FIG. 142D is a side view of the virtual anatomy 6780 for dissection, according to one aspect of the present disclosure. With reference to FIGS. 142C-142D, using the tablet computer, the surgeon can zoom and pan 3600 to obtain an optimal view of the virtual anatomy 6780 for dissection. The surgeon then determines the best path or trajectory 6776 to insert the surgical device 6778 (e.g., a dissector in this example). The surgeon may view the anatomy in a three-dimensional perspective view or any one of six views. See for example the side view of the virtual anatomy in FIG. 142D and the insertion of the surgical device 6778 (e.g., the dissector).

[1202] In another aspect, a sterile field control and data input console may allow live chatting between different departments, such as, for example, with the oncology or pathology department, to discuss margins or other particulars associated with imaging. The sterile field control and data input console may allow the pathology department to tell the surgeon about relationships of the margins within a specimen and show them to the surgeon in real time using the sterile field console.

[1203] In another aspect, a sterile field control and data input console may be used to change the focus and field of view of its own image or control that of any of the other monitors coupled to the surgical hub.

[1204] In another aspect, a sterile field control and data input console may be used to display the status of any of the equipment or modules coupled to the surgical hub 206. Knowledge of which device coupled to the surgical hub 206 is being used may be obtained via information such as the device is not on the instrument pad or on-device sensors. Based on this information, the sterile field control and data input console may change display, configurations, switch power to drive one device, and not another, one cord from capital to instrument pad and multiple cords from there. Device diagnostics may obtain knowledge that the device is inactive or not being used. Device diagnostics may be based on information such as the device is not on the instrument pad or based on-device sensors.

[1205] In another aspect, a sterile field control and data input console may be used as a learning tool. The console may display checklists, procedure steps, and/or sequence of steps. A timer/clock may be displayed to measure time to complete steps and/or procedures. The console may display room sound pressure level as indicator for activity, stress, etc.

[1206] FIGS. 143A-143B illustrate a touchscreen display 6890 that may be used within the sterile field, according to one aspect of the present disclosure. Using the touchscreen display 6890, a surgeon can manipulate images 6892 displayed on the touchscreen display 6890 using a variety of gestures such as, for example, drag and drop, scroll, zoom, rotate, tap, double tap, flick, drag, swipe, pinch open, pinch close, touch and hold, two-finger scroll, among others.

[1207] FIG. 143A illustrates an image 6892 of a surgical site displayed on a touchscreen display 6890 in portrait mode. FIG. 143B shows the touchscreen display 6890

rotated **6894** to landscape mode and the surgeon uses his index finger **6896** to scroll the image **6892** in the direction of the arrows. FIG. **143C** shows the surgeon using his index finger **6896** and thumb **6898** to pinch open the image **6892** in the direction of the arrows **6899** to zoom in. FIG. **143D** shows the surgeon using his index finger **6896** and thumb **6898** to pinch close the image **6892** in the direction of the arrows **6897** to zoom out. FIG. **143E** shows the touchscreen display **6890** rotated in two directions indicated by arrows **6894**, **6896** to enable the surgeon to view the image **6892** in different orientations.

[**1208**] Outside the sterile field, control and static displays are used that are different from the control and static displays used inside the sterile field. The control and static displays located outside the sterile field provide interactive and static displays for operating theater (OR) and device control. The control and static displays located outside the sterile field may include secondary static displays and secondary touchscreens for input and output.

[**1209**] Secondary static non-sterile displays **107**, **109**, **119** (FIG. **2**) for used outside the sterile field include monitors placed on the wall of the operating theater, on a rolling stand, or on capital equipment. A static display is presented with a feed from the control device to which they are attached and merely displays what is presented to it.

[**1210**] Secondary touch input screens located outside the sterile field may be part of the visualization system **108** (FIG. **2**), part of the surgical hub **108** (FIG. **2**), or may be fixed placement touch monitors on the walls or rolling stands. One difference between secondary touch input screens and static displays is that a user can interact with a secondary touch input screen by changing what is displayed on that specific monitor or others. For capital equipment applications, it could be the interface to control the setting of the connected capital equipment. The secondary touch input screens and the static displays outside the sterile field can be used to preload the surgeon's preferences (instrumentation settings and modes, lighting, procedure and preferred steps and sequence, music, etc.)

[**1211**] Secondary surgeon displays may include personal input displays with a personal input device that functions similarly to the common sterile field input display device but it is controlled by a specific surgeon. Personal secondary displays may be implemented in many form factors such as, for example, a watch, a small display pad, interface glasses, etc. A personal secondary display may include control capabilities of a common display device and since it is located on or controlled by a specific surgeon, the personal secondary display would be keyed to him/her specifically and would indicate that to others and itself. Generally speaking, a personal secondary display would normally not be useful to exchanging paired devices because they are not accessible to more than one surgeon. Nevertheless, a personal secondary display could be used to grant permission for release of a device.

[**1212**] A personal secondary display may be used to provide dedicated data to one of several surgical personnel that wants to monitor something that the others typically would not want to monitor. In addition, a personal secondary display may be used as the command module. Further, a personal secondary display may be held by the chief surgeon in the operating theater and would give the surgeon the control to override any of the other inputs from anyone else. A personal secondary display may be coupled to a short

range wireless, e.g., Bluetooth, microphone and earpiece allowing the surgeon to have discrete conversations or calls or the personal secondary display may be used to broadcast to all the others in the operating theater or other department.

[**1213**] FIG. **144** illustrates a surgical site **6900** employing a smart surgical retractor **6902** comprising a direct interface control to a surgical hub **206** (FIGS. **1-11**), according to one aspect of the present disclosure. The smart surgical retractor **6902** helps the surgeon and operating room professionals hold an incision or wound open during surgical procedures. The smart surgical retractor **6902** aids in holding back underlying organs or tissues, allowing doctors/nurses better visibility and access to the exposed area. With reference also to FIGS. **1-11**, the smart surgical retractor **6902** may comprise an input display **6904** operated by the smart surgical retractor **6902**. The smart surgical retractor **6902** may comprise a wireless communication device to communicate with a device connected to a generator module **240** coupled to the surgical hub **206**. Using the input display **6904** of the smart surgical retractor **6902**, the surgeon can adjust power level or mode of the generator module **240** to cut and/or coagulate tissue. If using automatic on/off for energy delivery on closure of an end effector on the tissue, the status of automatic on/off may be indicated by a light, screen, or other device located on the smart retractor **6902** housing. Power being used may be changed and displayed.

[**1214**] In one aspect, the smart surgical retractor **6902** can sense or know what device/instrument **235** the surgeon is using, either through the surgical hub **206** or RFID or other device placed on the device/instrument **235** or the smart surgical retractor **6902**, and provide an appropriate display. Alarm and alerts may be activated when conditions require. Other features include displaying the temperature of the ultrasonic blade, nerve monitoring, light source **6906** or fluorescence. The light source **6906** may be employed to illuminate the surgical field of view **6908** and to charge photocells **6918** on single use sticker display that stick onto the smart retractor **6902** (see FIG. **145**, for example). In another aspect, the smart surgical retractor **6902** may include an augmented reality projected on the patient's anatomy (e.g., like a vein viewer).

[**1215**] FIG. **145** illustrates a surgical site **6910** with a smart flexible sticker display **6912** attached to the body/skin **6914** of a patient, according to one aspect of the present disclosure. As shown, the smart flexible sticker display **6912** is applied to the body/skin **6914** of a patient between the area exposed by the surgical retractors **6916**. In one aspect, the smart flexible sticker display **6912** may be powered by light, an on board battery, or a ground pad. The flexible sticker display **6912** may communicate via short range wireless (e.g., Bluetooth) to a device, may provide readouts, lock power, or change power. The smart flexible sticker display **6912** also comprises photocells **6918** to power the smart flexible sticker display **6912** using ambient light energy. The flexible sticker display **6912** includes a display of a control panel **6920** user interface to enable the surgeon to control devices **235** or other modules coupled to the surgical hub **206** (FIGS. **1-11**).

[**1216**] FIG. **146** is a logic flow diagram **6920** of a process depicting a control program or a logic configuration to communicate from inside a sterile field to a device located outside the sterile field, according to one aspect of the present disclosure. In one aspect, a control unit comprises an interactive touchscreen display, an interface configured to

couple the interactive touchscreen display to a surgical hub, a processor, and a memory coupled to the processor. The memory stores instructions executable by the processor to receive **6922** input commands from the interactive touchscreen display located inside a sterile field and transmits **6924** the input commands to a surgical hub to control devices coupled to the surgical hub located outside the sterile field.

[1217] FIG. 147 illustrates a system for performing surgery. The system comprises a control box which includes internal circuitry; a surgical instrument including a distal element and techniques for sensing a position or condition of said distal element; techniques associated with said surgical instrument for transmitting said sensed position or condition to said internal circuitry of said control box; and for transmitting said sensed position or condition from said internal circuitry of said control box to a video monitor for display thereon, wherein said sensed position or condition is displayed on said video monitor as an icon or symbol, further comprising a voltage source for generating a voltage contained entirely within said surgical instrument. Further examples are disclosed in U.S. Pat. No. 5,503,320, titled SURGICAL APPARATUS WITH INDICATOR, which issued on Apr. 2, 1996, which is herein incorporated by reference in its entirety.

[1218] FIG. 147 shows schematically a system whereby data is transmitted to a video monitor for display, such data relating to the position and/or condition of one or more surgical instruments. As shown in FIG. 147, a laparoscopic surgical procedure is being performed wherein a plurality of trocar sleeves **6930** are inserted through a body wall **6931** to provide access to a body cavity **6932**. A laparoscope **6933** is inserted through one of the trocar sleeves **6930** to provide illumination (light cable **6934** is shown leading toward a light source, not pictured) to the surgical site and to obtain an image thereof. A camera adapter **6935** is attached at the proximal end of laparoscope **6933** and image cable **6936** extends therefrom to a control box **6937** discussed in more detail below. Image cable inputs to image receiving port **416** on control box **6937**.

[1219] Additional surgical instruments **6939**, **6940** are inserted through additional trocar sleeves **6900** which extend through body wall **6931**. In FIG. 147, instrument **6939** schematically illustrates an endoscopic stapling device, e.g., an Endo GIA\* instrument manufactured by the assignee of this application, and instrument **6940** schematically illustrates a hand instrument, e.g., an Endo Grasp\* device also manufactured by the present assignee. Additional and/or alternative instruments may also be utilized according to the present invention; the illustrated instruments are merely exemplary of surgical instruments which may be utilized according to the present invention.

[1220] Instruments **6939**, **6940** include adapters **6941**, **6942** associated with their respective handle portions. The adapters electronically communicate with conductive mechanisms (not pictured). These mechanisms, which include electrically conductive contact members electrically connected by wires, cables and the like, are associated with the distal elements of the respective instruments, e.g., the anvil **6943** and cartridge **6944** of the Endo GIA\* instrument, the jaws **6945**, **6946** of the Endo Grasp\* device, and the like. The mechanisms are adapted to interrupt an electronic circuit when the distal elements are in a first position or condition and to complete the electronic circuit when the

distal elements are in a second position or condition. A voltage source for the electronic circuit may be provided in the surgical instrument, e.g., in the form of a battery, or supplied from control box **6937** through cables **6947**, **6948**.

[1221] Control box **6937** includes a plurality of jacks **6949** which are adapted to receive cables **6947**, **6948** and the like. Control box **6937** further includes an outgoing adapter **6950** which is adapted to cooperate with a cable **6951** for transmitting the laparoscopic image obtained by the laparoscope **6933** together with data concerning surgical instruments **6939**, **6940** to video monitor **6952**. Circuitry within control box **6937** is provided for converting the presence of an interrupted circuit, e.g., for the electronics within cable **6947** and the mechanism associated with the distal elements of instrument **6939**, to an icon or symbol for display on video monitor **6952**. Similarly, the circuitry within control box **6937** is adapted to provide a second icon or symbol to video monitor **6952** when a completed circuit exists for cable **6947** and the associated mechanism.

[1222] Illustrative icons/symbols **6953**, **6954** are shown on video monitor **6952**. Icon **6953** shows a surgical staple and could be used to communicate to the surgeon that the cartridge **6944** and anvil **6943** of instrument **6939** are properly positioned to form staples in tissue **6955**. Icon **6953** could take another form when the cartridge **6944** and anvil **6943** are not properly positioned for forming staples, thereby interrupting the circuit. Icon **6954** shows a hand instrument with jaws spread apart, thereby communicating to the surgeon that the jaws **6945**, **6946** of instrument **6940** are open. Icon **6954** could take another form when jaws **6945**, **6946** are closed, thereby completing the circuit.

[1223] FIG. 148 illustrates a second layer of information overlaying a first layer of information. The second layer of information includes a symbolic representation of the knife overlapping the detected position of the knife in the DLU depicted in the first layer of information. Further examples are disclosed in U.S. Pat. No. 9,283,054, titled SURGICAL APPARATUS WITH INDICATOR, which issued on Mar. 15, 2016, which is herein incorporated by reference in its entirety.

[1224] Referring to FIG. 148, the second layer of information **6963** can overlay at least a portion of the first layer of information **6962** on the display **6960**. Furthermore, the touch screen **6961** can allow a user to manipulate the second layer of information **6963** relative to the video feedback in the underlying first layer of information **6962** on the display **6960**. For example, a user can operate the touch screen **6961** to select, manipulate, reformat, resize, and/or otherwise modify the information displayed in the second layer of information **6963**. In certain aspects, the user can use the touch screen **6961** to manipulate the second layer of information **6963** relative to the surgical instrument **6964** depicted in the first layer of information **6962** on the display **6960**. A user can select a menu, category and/or classification of the control panel **6967** thereof, for example, and the second layer of information **6963** and/or the control panel **6967** can be adjusted to reflect the user's selection. In various aspects, a user may select a category from the instrument feedback category **6969** that corresponds to a specific feature or features of the surgical instrument **6964** depicted in the first layer of information **6962**. Feedback corresponding to the user-selected category can move, locate itself, and/or "snap" to a position on the display **6960** relative to the specific feature or features of the surgical

instrument 6964. For example, the selected feedback can move to a position near and/or overlapping the specific feature or features of the surgical instrument 6964 depicted in the first layer of information 6962.

[1225] The instrument feedback menu 6969 can include a plurality of feedback categories, and can relate to the feedback data measured and/or detected by the surgical instrument 6964 during a surgical procedure. As described herein, the surgical instrument 6964 can detect and/or measure the position 6970 of a moveable jaw between an open orientation and a closed orientation, the thickness 6973 of clamped tissue, the clamping force 6976 on the clamped tissue, the articulation 6974 of the DLU 6965, and/or the position 6971, velocity 6972, and/or force 6975 of the firing element, for example. Furthermore, the feedback controller in signal communication with the surgical instrument 6964 can provide the sensed feedback to the display 6960, which can display the feedback in the second layer of information 6963. As described herein, the selection, placement, and/or form of the feedback data displayed in the second layer of information 6963 can be modified based on the user's input to the touch screen 6961, for example.

[1226] When the knife of the DLU 6965 is blocked from view by the end effector jaws 6966 and/or tissue T, for example, the operator can track and/or approximate the position of the knife in the DLU 6964 based on the changing value of the feedback data and/or the shifting position of the feedback data relative to the DLU 6965 depicted in the underlying first layer of information 6962.

[1227] In various aspects, the display menu 6977 of the control panel 6967 can relate to a plurality of categories, such as unit systems 6978 and/or data modes 6979, for example. In certain aspects, a user can select the unit systems category 6978 to switch between unit systems, such as between metric and U.S. customary units, for example. Additionally, a user can select the data mode category 6979 to switch between types of numerical representations of the feedback data and/or types of graphical representations of the feedback data, for example. The numerical representations of the feedback data can be displayed as numerical values and/or percentages, for example. Furthermore, the graphical representations of the feedback data can be displayed as a function of time and/or distance, for example. As described herein, a user can select the instrument controller menu 6980 from the control panel 6967 to input directives for the surgical instrument 6964, which can be implemented via the instrument controller and/or the microcontroller, for example. A user can minimize or collapse the control panel 6967 by selecting the minimize/maximize icon 6968, and can maximize or un-collapse the control panel 6967 by re-selecting the minimize/maximize icon 6968.

[1228] FIG. 149 depicts a perspective view of a surgeon using a surgical instrument that includes a handle assembly housing and a wireless circuit board during a surgical procedure, with the surgeon wearing a set of safety glasses. The wireless circuit board transmits a signal to a set of safety glasses worn by a surgeon using the surgical instrument during a procedure. The signal is received by a wireless port on the safety glasses. One or more lighting devices on a front lens of the safety glasses change color, fade, or glow in response to the received signal to indicate information to the surgeon about the status of the surgical instrument. The lighting devices are disposable on peripheral edges of the front lens to not distract the direct line of vision of the

surgeon. Further examples are disclosed in U.S. Pat. No. 9,011,427, titled SURGICAL INSTRUMENT WITH SAFETY GLASSES, which issued on Apr. 21, 2015, which is herein incorporated by reference in its entirety.

[1229] FIG. 149 shows a version of safety glasses 6991 that may be worn by a surgeon 6992 during a surgical procedure while using a medical device. In use, a wireless communications board housed in a surgical instrument 6993 may communicate with a wireless port 6994 on safety glasses 6991. Exemplary surgical instrument 6993 is a battery-operated device, though instrument 6993 could be powered by a cable or otherwise. Instrument 6993 includes an end effector. Particularly, wireless communications board 6995 transmits one or more wireless signals indicated by arrows (B, C) to wireless port 6994 of safety glasses 6991. Safety glasses 6991 receive the signal, analyze the received signal, and display indicated status information received by the signal on lenses 6996 to a user, such as surgeon 6992, wearing safety glasses 6991. Additionally or alternatively, wireless communications board 6995 transmits a wireless signal to surgical monitor 6997 such that surgical monitor 6997 may display received indicated status information to surgeon 6992, as described above.

[1230] Aversion of the safety glasses 6991 may include lighting device on peripheral edges of the safety glasses 6991. A lighting device provides peripheral-vision sensory feedback of instrument 6993, with which the safety glasses 6991 communicate to a user wearing the safety glasses 6991. The lighting device may be, for example, a light-emitted diode ("LED"), a series of LEDs, or any other suitable lighting device known to those of ordinary skill in the art and apparent in view of the teachings herein.

[1231] LEDs may be located at edges or sides of a front lens of the safety glasses 6991 so not to distract from a user's center of vision while still being positioned within the user's field of view such that the user does not need to look away from the surgical site to see the lighting device. Displayed lights may pulse and/or change color to communicate to the wearer of the safety glasses 6991 various aspects of information retrieved from instrument 6993, such as system status information or tissue sensing information (i.e., whether the end effector has sufficiently severed and sealed tissue). Feedback from housed wireless communications board 6995 may cause a lighting device to activate, blink, or change color to indicate information about the use of instrument 6993 to a user. For example, a device may incorporate a feedback mechanism based on one or more sensed tissue parameters. In this case, a change in the device output(s) based on this feedback in synch with a tone change may submit a signal through wireless communications board 6995 to the safety glasses 6991 to trigger activation of the lighting device. Such described means of activation of the lighting device should not be considered limiting as other means of indicating status information of instrument 6993 to the user via the safety glasses 6991 are contemplated. Further, the safety glasses 6991 may be single-use or reusable eyewear. Button-cell power supplies such as button-cell batteries may be used to power wireless receivers and LEDs of versions of safety glasses 6991, which may also include a housed wireless board and tri-color LEDs. Such button-cell power supplies may provide a low-cost means of providing sensory feedback of information about instrument 6993 when in use to surgeon 6992 wearing safety glasses 6991.

[1232] FIG. 150 is a schematic diagram of a feedback control system for controlling a surgical instrument. The surgical instrument includes a housing and an elongated shaft that extends distally from the housing and defines a first longitudinal axis. The surgical instrument also includes a firing rod disposed in the elongated shaft and a drive mechanism disposed at least partially within the housing. The drive mechanism mechanically cooperates with the firing rod to move the firing rod. A motion sensor senses a change in the electric field (e.g., capacitance, impedance, or admittance) between the firing rod and the elongated shaft. The measurement unit determines a parameter of the motion of the firing rod, such as the position, speed, and direction of the firing rod, based on the sensed change in the electric field. A controller uses the measured parameter of the motion of the firing rod to control the drive mechanism. Further examples are disclosed in U.S. Pat. No. 8,960,520, titled METHOD AND APPARATUS FOR DETERMINING PARAMETERS OF LINEAR MOTION IN A SURGICAL INSTRUMENT, which issued on Feb. 24, 2015, which is herein incorporated by reference in its entirety.

[1233] With reference to FIG. 150, aspects of the present disclosure may include a feedback control system 6150. The system 6150 includes a feedback controller 6152. The surgical instrument 6154 is connected to the feedback controller 6152 via a data port, which may be either wired (e.g., FireWire®, USB, Serial RS232, Serial RS485, USART, Ethernet, etc.) or wireless (e.g., Bluetooth®, ANT3®, KNX®, Z-Wave X10®, Wireless USB®, Wi-Fi®, IrDA®, nanoNET®, TinyOS®, ZigBee®, 802.11 IEEE, and other radio, infrared, UHF, VHF communications and the like). The feedback controller 6152 is configured to store the data transmitted to it by the surgical instrument 6154 as well as process and analyze the data. The feedback controller 6152 is also connected to other devices, such as a video display 6154, a video processor 6156 and a computing device 6158 (e.g., a personal computer, a PDA, a smartphone, a storage device, etc.). The video processor 6156 is used for processing output data generated by the feedback controller 6152 for output on the video display 6154. The computing device 6158 is used for additional processing of the feedback data. In one aspect, the results of the sensor feedback analysis performed by a microcontroller may be stored internally for later retrieval by the computing device 6158.

[1234] FIG. 151 illustrates a feedback controller 6152 including an on-screen display (OSD) module and a heads-up-display (HUD) module. The modules process the output of a microcontroller for display on various displays. More specifically, the OSD module overlays text and/or graphical information from the feedback controller 6152 over other video images received from the surgical site via cameras disposed therein. The modified video signal having overlaid text is transmitted to the video display allowing the user to visualize useful feedback information from the surgical instrument 6154 and/or feedback controller 6152 while still observing the surgical site. The feedback controller 6152 includes a data port 6160 coupled to a microcontroller which allows the feedback controller 6152 to be connected to the computing device 6158 (FIG. 150). The data port 6160 may provide for wired and/or wireless communication with the computing device 6158 providing for an interface between the computing device 6158 and the feedback controller 6152 for retrieval of stored feedback data, configuration of oper-

ating parameters of the feedback controller 6152 and upgrade of firmware and/or other software of the feedback controller 6152.

[1235] The feedback controller 6152 includes a housing 6162 and a plurality of input and output ports, such as a video input 6164, a video output 6166, and a HUD display output 6168. The feedback controller 6152 also includes a screen for displaying status information concerning the feedback controller 6152. Further examples are disclosed in U.S. Pat. No. 8,960,520, titled METHOD AND APPARATUS FOR DETERMINING PARAMETERS OF LINEAR MOTION IN A SURGICAL INSTRUMENT, which issued on Feb. 24, 2015, which is herein incorporated by reference in its entirety.

#### Visualization System

[1236] During a surgical procedure, a surgeon may be required to manipulate tissues to effect a desired medical outcome. The actions of the surgeon are limited by what is visually observable in the surgical site. Thus, the surgeon may not be aware, for example, of the disposition of vascular structures that underlie the tissues being manipulated during the procedure. Since the surgeon is unable to visualize the vasculature beneath a surgical site, the surgeon may accidentally sever one or more critical blood vessels during the procedure. The solution is a surgical visualization system that can acquire imaging data of the surgical site for presentation to a surgeon, in which the presentation can include information related to the presence and depth of vascular structures located beneath the surface of a surgical site.

[1237] In one aspect, the surgical hub 106 incorporates a visualization system 108 to acquire imaging data during a surgical procedure. The visualization system 108 may include one or more illumination sources and one or more light sensors. The one or more illumination sources and one or more light sensors may be incorporated together into a single device or may comprise one or more separate devices. The one or more illumination sources may be directed to illuminate portions of the surgical field. The one or more light sensors may receive light reflected or refracted from the surgical field including light reflected or refracted from tissue and/or surgical instruments. The following description includes all of the hardware and software processing techniques disclosed above and in those applications incorporated herein by reference as presented above.

[1238] In some aspects, the visualization system 108 may be integrated into a surgical system 100 as disclosed above and depicted in FIGS. 1 and 2. In addition to the visualization system 108, the surgical system 100 may include one or more hand-held intelligent instruments 112, a multi-functional robotic system 110, one or more visualization systems 108, and a centralized surgical hub system 106, among other components. The centralized surgical hub system 106 may control several functions as disclosed above and also depicted in FIG. 3. In one non-limiting example, such functions may include supplying and controlling power to any number of powered surgical devices. In another non-limiting example, such functions may include controlling fluid supplied to and evacuated from the surgical site. The centralized surgical hub system 106 may also be configured to manage and analyze data received from any of the surgical system components as well as communicate data and other information among and between the components of the surgical system. The centralized surgical hub system 106 may also be



in data communication with a cloud computing system **104** as disclosed above and depicted, for example, in FIG. 1.

[1239] In some non-limiting examples, imaging data generated by the visualization system **108** may be analyzed by on-board computational components of the visualization system **108**, and analysis results may be communicated to the centralized surgical hub **106**. In alternative non-limiting examples, the imaging data generated by the visualization system **108** may be communicated directly to the centralized surgical hub **106** where the data may be analyzed by computational components in the hub system **106**. The centralized surgical hub **106** may communicate the image analysis results to any one or more of the other components of the surgical system. In some other non-limiting examples, the centralized surgical hub may communicate the image data and/or the image analysis results to the cloud computing system **104**.

[1240] FIGS. 152A-D and FIGS. 153A-F depict various aspects of one example of a visualization system **2108** that may be incorporated into a surgical system. The visualization system **2108** may include an imaging control unit **2002** and a hand unit **2020**. The imaging control unit **2002** may include one or more illumination sources, a power supply for the one or more illumination sources, one or more types of data communication interfaces (including USB, Ethernet, or wireless interfaces **2004**), and one or more video outputs **2006**. The imaging control unit **2002** may further include an interface, such as a USB interface **2010**, configured to transmit integrated video and image capture data to a USB enabled device. The imaging control unit **2002** may also include one or more computational components including, without limitation, a processor unit, a transitory memory unit, a non-transitory memory unit, an image processing unit, a bus structure to form data links among the computational components, and any interface (e.g. input and/or output) devices necessary to receive information from and transmit information to components not included in the imaging control unit. The non-transitory memory may further contain instructions that when executed by the processor unit, may perform any number of manipulations of data that may be received from the hand unit **2020** and/or computational devices not included in the imaging control unit.

[1241] The illumination sources may include a white light source **2012** and one or more laser light sources. The imaging control unit **2002** may include one or more optical and/or electrical interfaces for optical and/or electrical communication with the hand unit **2020**. The one or more laser light sources may include, as non-limiting examples, any one or more of a red laser light source, a green laser light source, a blue laser light source, an infrared laser light source, and an ultraviolet laser light source. In some non-limiting examples, the red laser light source may source illumination having a peak wavelength that may range between 635 nm and 660 nm, inclusive. Non-limiting examples of a red laser peak wavelength may include about 635 nm, about 640 nm, about 645 nm, about 650 nm, about 655 nm, about 660 nm, or any value or range of values therebetween. In some non-limiting examples, the green laser light source may source illumination having a peak wavelength that may range between 520 nm and 532 nm, inclusive. Non-limiting examples of a green laser peak wavelength may include about 520 nm, about 522 nm, about 524 nm, about 526 nm, about 528 nm, about 530 nm, about

532 nm, or any value or range of values therebetween. In some non-limiting examples, the blue laser light source may source illumination having a peak wavelength that may range between 405 nm and 445 nm, inclusive. Non-limiting examples of a blue laser peak wavelength may include about 405 nm, about 410 nm, about 415 nm, about 420 nm, about 425 nm, about 430 nm, about 435 nm, about 440 nm, about 445 nm, or any value or range of values therebetween. In some non-limiting examples, the infrared laser light source may source illumination having a peak wavelength that may range between 750 nm and 3000 nm, inclusive. Non-limiting examples of an infrared laser peak wavelength may include about 750 nm, about 1000 nm, about 1250 nm, about 1500 nm, about 1750 nm, about 2000 nm, about 2250 nm, about 2500 nm, about 2750 nm, 3000 nm, or any value or range of values therebetween. In some non-limiting examples, the ultraviolet laser light source may source illumination having a peak wavelength that may range between 200 nm and 360 nm, inclusive. Non-limiting examples of an ultraviolet laser peak wavelength may include about 200 nm, about 220 nm, about 240 nm, about 260 nm, about 280 nm, about 300 nm, about 320 nm, about 340 nm, about 360 nm, or any value or range of values therebetween.

[1242] In one non-limiting aspect, the hand unit **2020** may include a body **2021**, a camera scope cable **2015** attached to the body **2021**, and an elongated camera probe **2024**. The body **2021** of the hand unit **2020** may include hand unit control buttons **2022** or other controls to permit a health professional using the hand unit **2020** to control the operations of the hand unit **2020** or other components of the imaging control unit **2002**, including, for example, the light sources. The camera scope cable **2015** may include one or more electrical conductors and one or more optical fibers. The camera scope cable **2015** may terminate with a camera head connector **2008** at a proximal end in which the camera head connector **2008** is configured to mate with the one or more optical and/or electrical interfaces of the imaging control unit **2002**. The electrical conductors may supply power to the hand unit **2020**, including the body **2021** and the elongated camera probe **2024**, and/or to any electrical components internal to the hand unit **2020** including the body **2021** and/or elongated camera probe **2024**. The electrical conductors may also serve to provide bi-directional data communication between any one or more components the hand unit **2020** and the imaging control unit **2002**. The one or more optical fibers may conduct illumination from the one or more illumination sources in the imaging control unit **2002** through the hand unit body **2021** and to a distal end of the elongated camera probe **2024**. In some non-limiting aspects, the one or more optical fibers may also conduct light reflected or refracted from the surgical site to one or more optical sensors disposed in the elongated camera probe **2024**, the hand unit body **2021**, and/or the imaging control unit **2002**.

[1243] FIG. 152B (a top plan view) depicts in more detail some aspects of a hand unit **2020** of the visualization system **2108**. The hand unit body **2021** may be constructed of a plastic material. The hand unit control buttons **2022** or other controls may have a rubber overmolding to protect the controls while permitting them to be manipulated by the surgeon. The camera scope cable **2015** may have optical fibers integrated with electrical conductors, and the camera scope cable **2015** may have a protective and flexible overcoating such as PVC. In some non-limiting examples, the

camera scope cable **2015** may be about 10 ft. long to permit ease of use during a surgical procedure. The length of the camera scope cable **2015** may range from about 5 ft. to about 15 ft. Non-limiting examples of a length of the camera scope cable **2015** may be about 5 ft., about 6 ft., about 7 ft., about 8 ft., about 9 ft., about 10 ft., about 11 ft., about 12 ft., about 13 ft., about 14 ft., about 15 ft., or any length or range of lengths therebetween. The elongated camera probe **2024** may be fabricated from a rigid material such as stainless steel. In some aspects, the elongated camera probe **2024** may be joined with the hand unit body **2021** via a rotatable collar **2026**. The rotatable collar **2026** may permit the elongated camera probe **2024** to be rotated with respect to the hand unit body **2021**. In some aspects, the elongated camera probe **2024** may terminate at a distal end with a plastic window **2028** sealed with epoxy.

[1244] The side plan view of the hand unit, depicted in FIG. 152C illustrates that a light or image sensor **2030** maybe disposed at a distal end **2032a** of the elongated camera probe or within the hand unit body **2032b**. In some alternative aspects, the light or image sensor **2030** may be disposed with additional optical elements in the imaging control unit **2002**. FIG. 152C further depicts an example of a light sensor **2030** comprising a CMOS image sensor **2034** disposed within a mount **2036** having a radius of about 4 mm. FIG. 152D illustrates aspects of the CMOS image sensor **2034**, depicting the active area **2038** of the image sensor. Although the CMOS image sensor in FIG. 152C is depicted to be disposed within a mount **2036** having a radius of about 4 mm, it may be recognized that such a sensor and mount combination may be of any useful size to be disposed within the elongated camera probe **2024**, the hand unit body **2021**, or in the image control unit **2002**. Some non-limiting examples of such alternative mounts may include a 5.5 mm mount **2136a**, a 4 mm mount **2136b**, a 2.7 mm mount **2136c**, and a 2 mm mount **2136d**. It may be recognized that the image sensor may also comprise a CCD image sensor. The CMOS or CCD sensor may comprise an array of individual light sensing elements (pixels).

[1245] FIGS. 153A-153F depict various aspects of some examples of illumination sources and their control that may be incorporated into the visualization system **2108**.

[1246] FIG. 153A illustrates an aspect of a laser illumination system having a plurality of laser bundles emitting a plurality of wavelengths of electromagnetic energy. As can be seen in the figure, the illumination system **2700** may comprise a red laser bundle **2720**, a green laser bundle **2730**, and a blue laser bundle **2740** that are all optically coupled together though fiber optics **2755**. As can be seen in the figure, each of the laser bundles may have a corresponding light sensing element or electromagnetic sensor **2725**, **2735**, **2745** respectively, for sensing the output of the specific laser bundle or wavelength.

[1247] Additional disclosures regarding the laser illumination system depicted in FIG. 153A for use in a surgical visualization system **2108** may be found in U.S. Patent Application Publication No. 2014/0268860, titled CONTROLLING THE INTEGRAL LIGHT ENERGY OF A LASER PULSE filed on Mar. 15, 2014, which issued on Oct. 3, 2017 as U.S. Pat. No. 9,777,913, the contents thereof being incorporated by reference herein in its entirety and for all purposes.

[1248] FIG. 153B illustrates the operational cycles of a sensor used in rolling readout mode. It will be appreciated

that the x direction corresponds to time and the diagonal lines **2202** indicate the activity of an internal pointer that reads out each frame of data, one line at time. The same pointer is responsible for resetting each row of pixels for the next exposure period. The net integration time for each row **2219a-c** is equivalent, but they are staggered in time with respect to one another due to the rolling reset and read process. Therefore, for any scenario in which adjacent frames are required to represent different constitutions of light, the only option for having each row be consistent is to pulse the light between the readout cycles **2230a-c**. More specifically, the maximum available period corresponds to the sum of the blanking time plus any time during which optical black or optically blind (OB) rows (**2218**, **2220**) are serviced at the start or end of the frame.

[1249] FIG. 153B illustrates the operational cycles of a sensor used in rolling readout mode or during the sensor readout **2200**. The frame readout may start at and may be represented by vertical line **2210**. The read out period is represented by the diagonal or slanted line **2202**. The sensor may be read out on a row by row basis, the top of the downwards slanted edge being the sensor top row **2212** and the bottom of the downwards slanted edge being the sensor bottom row **2214**. The time between the last row readout and the next readout cycle may be called the blanking time **2216a-d**. It may be understood that the blanking time **2216a-d** may be the same between success readout cycles or it may differ between success readout cycles. It should be noted that some of the sensor pixel rows might be covered with a light shield (e.g., a metal coating or any other substantially black layer of another material type). These covered pixel rows may be referred to as optical black rows **2218** and **2220**. Optical black rows **2218** and **2220** may be used as input for correction algorithms.

[1250] As shown in FIG. 153B, these optical black rows **2218** and **2220** may be located on the top of the pixel array or at the bottom of the pixel array or at the top and the bottom of the pixel array. In some aspects, it may be desirable to control the amount of electromagnetic radiation, e.g., light, that is exposed to a pixel, thereby integrated or accumulated by the pixel. It will be appreciated that photons are elementary particles of electromagnetic radiation. Photons are integrated, absorbed, or accumulated by each pixel and converted into an electrical charge or current. In some aspects, an electronic shutter or rolling shutter may be used to start the integration time (**2219a-c**) by resetting the pixel. The light will then integrate until the next readout phase. In some aspects, the position of the electronic shutter can be moved between two readout cycles **2202** in order to control the pixel saturation for a given amount of light. In some alternative aspects lacking an electronic shutter, the integration time **2219a-c** of the incoming light may start during a first readout cycle **2202** and may end at the next readout cycle **2202**, which also defines the start of the next integration. In some alternative aspects, the amount of light accumulated by each pixel may be controlled by a time during which light is pulsed **2230a-d** during the blanking times **2216a-d**. This ensures that all rows see the same light issued from the same light pulse **2230a-c**. In other words, each row will start its integration in a first dark environment **2231**, which may be at the optical black back row **2220** of read out frame (m) for a maximum light pulse width, and will then receive a light strobe and will end its integration in a second dark environment **2232**, which may be at the optical black

front row **2218** of the next succeeding read out frame (m+1) for a maximum light pulse width. Thus, the image generated from the light pulse **2230a-c** will be solely available during frame (m+1) readout without any interference with frames (m) and (m+2).

[1251] It should be noted that the condition to have a light pulse **2230a-c** to be read out only in one frame and not interfere with neighboring frames is to have the given light pulse **2230a-c** firing during the blanking time **2216**. Because the optical black rows **2218**, **2220** are insensitive to light, the optical black back rows **2220** time of frame (m) and the optical black front rows **2218** time of frame (m+1) can be added to the blanking time **2216** to determine the maximum range of the firing time of the light pulse **2230**.

[1252] In some aspects, FIG. **153B** depicts an example of a timing diagram for sequential frame captures by a conventional CMOS sensor. Such a CMOS sensor may incorporate a Bayer pattern of color filters, as depicted in FIG. **153C**. It is recognized that the Bayer pattern provides for greater luminance detail than chrominance. It may further be recognized that the sensor has a reduced spatial resolution since a total of 4 adjacent pixels are required to produce the color information for the aggregate spatial portion of the image. In an alternative approach, the color image may be constructed by rapidly strobing the visualized area at high speed with a variety of optical sources (either laser or light-emitting diodes) having different central optical wavelengths.

[1253] The optical strobing system may be under the control of the camera system, and may include a specially designed CMOS sensor with high speed readout. The principal benefit is that the sensor can accomplish the same spatial resolution with significantly fewer pixels compared with conventional Bayer or 3-sensor cameras. Therefore, the physical space occupied by the pixel array may be reduced. The actual pulse periods (**2230a-c**) may differ within the repeating pattern, as illustrated in FIG. **153B**. This is useful for, e.g., apportioning greater time to the components that require the greater light energy or those having the weaker sources. As long as the average captured frame rate is an integer multiple of the requisite final system frame rate, the data may simply be buffered in the signal processing chain as appropriate.

[1254] The facility to reduce the CMOS sensor chip-area to the extent allowed by combining all of these methods is particularly attractive for small diameter (~3-10 mm) endoscopy. In particular, it allows for endoscope designs in which the sensor is located in the space-constrained distal end, thereby greatly reducing the complexity and cost of the optical section, while providing high definition video. A consequence of this approach is that to reconstruct each final, full color image, requires that data be fused from three separate snapshots in time. Any motion within the scene, relative to the optical frame of reference of the endoscope, will generally degrade the perceived resolution, since the edges of objects appear at slightly different locations within each captured component. In this disclosure, a means of diminishing this issue is described which exploits the fact that spatial resolution is much more important for luminance information, than for chrominance.

[1255] The basis of the approach is that, instead of firing monochromatic light during each frame, combinations of the three wavelengths are used to provide all of the luminance information within a single image. The chrominance infor-

mation is derived from separate frames with, e.g., a repeating pattern such as Y-Cb-Y-Cr (FIG. **153D**). While it is possible to provide pure luminance data by a shrewd choice of pulse ratios, the same is not true of chrominance.

[1256] In one aspect, as illustrated in FIG. **153D**, an endoscopic system **2300a** may comprise a pixel array **2302a** having uniform pixels and the system **2300a** may be operated to receive Y (luminance pulse) **2304a**, Cb (ChromaBlue) **2306a** and Cr (ChromaRed) **2308a** pulses.

[1257] To complete a full color image requires that the two components of chrominance also be provided. However, the same algorithm that was applied for luminance cannot be directly applied for chrominance images since it is signed, as reflected in the fact that some of the RGB coefficients are negative. The solution to this is to add a degree of luminance of sufficient magnitude that all of the final pulse energies become positive. As long as the color fusion process in the ISP is aware of the composition of the chrominance frames, they can be decoded by subtracting the appropriate amount of luminance from a neighboring frame. The pulse energy proportions are given by:

$$Y=0.183\cdot R+0.614\cdot G+0.062\cdot B$$

$$Cb=\lambda\cdot Y-0.101\cdot R-0.339\cdot G+0.439\cdot B$$

$$Cr=\delta\cdot Y+0.439\cdot R-0.399\cdot G-0.040\cdot B$$

where

$$\lambda\geq 0.399/0.614=0.552$$

$$\delta\geq 0.399/0.614=0.650$$

[1258] It turns out that if the X factor is equal to 0.552; both the red and the green components are exactly cancelled, in which case the Cb information can be provided with pure blue light. Similarly, setting  $\delta=0.650$  cancels out the blue and green components for Cr which becomes pure red. This particular example is illustrated in FIG. **153E**, which also depicts  $\lambda$  and  $\delta$  as integer multiples of  $\frac{1}{2}^8$ . This is a convenient approximation for the digital frame reconstruction.

[1259] In the case of the Y-Cb-Y-Cr pulsing scheme, the image data is already in the YCbCr space following the color fusion. Therefore, in this case it makes sense to perform luminance and chrominance based operations up front, before converting back to linear RGB to perform the color correction etc.

[1260] The color fusion process is more straightforward than de-mosaic, which is necessitated by the Bayer pattern (see FIG. **153C**), since there is no spatial interpolation. It does require buffering of frames though in order to have all of the necessary information available for each pixel. In one general aspect, data for the Y-Cb-Y-Cr pattern may be pipelined to yield one full color image per two raw captured images. This is accomplished by using each chrominance sample twice. In FIG. **153F** the specific example of a 120 Hz frame capture rate providing 60 Hz final video is depicted.

[1261] Additional disclosures regarding the control of the laser components of an illumination system as depicted in FIGS. **153B-153F** for use in a surgical visualization system **108** may be found in U.S. Patent Application Publication No. 2014/0160318, titled YCBCR PULSED ILLUMINATION SCHEME IN A LIGHT DEFICIENT ENVIRONMENT, filed on Jul. 26, 2013, which issued on Dec. 6, 2016 as U.S. Pat. No. 9,516,239, and U.S. Patent Application

Publication No. 2014/0160319, titled CONTINUOUS VIDEO IN A LIGHT DEFICIENT ENVIRONMENT, filed on Jul. 26, 2013, which issued on Aug. 22, 2017 as U.S. Pat. No. 9,743,016, the contents thereof being incorporated by reference herein in their entirety and for all purposes.

#### Subsurface Vascular Imaging

**[1262]** During a surgical procedure, a surgeon may be required to manipulate tissues to effect a desired medical outcome. The actions of the surgeon are limited by what is visually observable in the surgical site. Thus, the surgeon may not be aware, for example, of the disposition of vascular structures that underlie the tissues being manipulated during the procedure.

**[1263]** Since the surgeon is unable to visualize the vasculature beneath a surgical site, the surgeon may accidentally sever one or more critical blood vessels during the procedure.

**[1264]** Therefore, it is desirable to have a surgical visualization system that can acquire imaging data of the surgical site for presentation to a surgeon in which the presentation can include information related to the presence of vascular structures located beneath the surface of a surgical site.

**[1265]** Some aspects of the present disclosure further provide for a control circuit configured to control the illumination of a surgical site using one or more illumination sources such as laser light sources and to receive imaging data from one or more image sensors. In some aspects, the present disclosure provides for a non-transitory computer readable medium storing computer readable instructions that, when executed, cause a device to detect a blood vessel in a tissue and determine its depth below the surface of the tissue.

**[1266]** In some aspects, a surgical image acquisition system may include a plurality of illumination sources wherein each illumination source is configured to emit light having a specified central wavelength, a light sensor configured to receive a portion of the light reflected from a tissue sample when illuminated by the one or more of the plurality of illumination sources, and a computing system. The computing system may be configured to: receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; determine a depth location of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources, and calculate visualization data regarding the structure and the depth location of the structure. In some aspects, the visualization data may have a data format that may be used by a display system, and the structure may comprise one or more vascular tissues.

#### Vascular Imaging Using NIR Spectroscopy

**[1267]** In one aspect, a surgical image acquisition system may include an independent color cascade of illumination sources comprising visible light and light outside of the visible range to image one or more tissues within a surgical site at different times and at different depths. The surgical image acquisition system may further detect or calculate characteristics of the light reflected and/or refracted from the surgical site. The characteristics of the light may be used to provide a composite image of the tissue within the surgical site as well as provide an analysis of underlying tissue not

directly visible at the surface of the surgical site. The surgical image acquisition system may determine tissue depth location without the need for separate measurement devices.

**[1268]** In one aspect, the characteristic of the light reflected and/or refracted from the surgical site may be an amount of absorbance of light at one or more wavelengths. Various chemical components of individual tissues may result in specific patterns of light absorption that are wavelength dependent.

**[1269]** In one aspect, the illumination sources may comprise a red laser source and a near infrared laser source, wherein the one or more tissues to be imaged may include vascular tissue such as veins or arteries. In some aspects, red laser sources (in the visible range) may be used to image some aspects of underlying vascular tissue based on spectroscopy in the visible red range. In some non-limiting examples, a red laser light source may source illumination having a peak wavelength that may range between 635 nm and 660 nm, inclusive. Non-limiting examples of a red laser peak wavelength may include about 635 nm, about 640 nm, about 645 nm, about 650 nm, about 655 nm, about 660 nm, or any value or range of values therebetween. In some other aspects, near infrared laser sources may be used to image underlying vascular tissue based on near infrared spectroscopy. In some non-limiting examples, a near infrared laser source may emit illumination have a wavelength that may range between 750-3000 nm, inclusive. Non-limiting examples of an infrared laser peak wavelength may include about 750 nm, about 1000 nm, about 1250 nm, about 1500 nm, about 1750 nm, about 2000 nm, about 2250 nm, about 2500 nm, about 2750 nm, 3000 nm, or any value or range of values therebetween. It may be recognized that underlying vascular tissue may be probed using a combination of red and infrared spectroscopy. In some examples, vascular tissue may be probed using a red laser source having a peak wavelength at about 660 nm and a near IR laser source having a peak wavelength at about 750 nm or at about 850 nm.

**[1270]** Near infrared spectroscopy (NIRS) is a non-invasive technique that allows determination of tissue oxygenation based on spectro-photometric quantitation of oxy- and deoxyhemoglobin within a tissue. In some aspects, NIRS can be used to image vascular tissue directly based on the difference in illumination absorbance between the vascular tissue and non-vascular tissue. Alternatively, vascular tissue can be indirectly visualized based on a difference of illumination absorbance of blood flow in the tissue before and after the application of physiological interventions, such as arterial and venous occlusions methods.

**[1271]** Instrumentation for near-IR (NIR) spectroscopy may be similar to instruments for the UV-visible and mid-IR ranges. Such spectroscopic instruments may include an illumination source, a detector, and a dispersive element to select a specific near-IR wavelength for illuminating the tissue sample. In some aspects, the source may comprise an incandescent light source or a quartz halogen light source. In some aspects, the detector may comprise semiconductor (for example, an InGaAs) photodiode or photo array. In some aspects, the dispersive element may comprise a prism or, more commonly, a diffraction grating. Fourier transform NIR instruments using an interferometer are also common, especially for wavelengths greater than about 1000 nm.

Depending on the sample, the spectrum can be measured in either reflection or transmission mode.

[1272] FIG. 154 depicts schematically one example of instrumentation 2400 similar to instruments for the UV-visible and mid-IR ranges for NIR spectroscopy. A light source 2402 may emit a broad spectral range of illumination 2404 that may impinge upon a dispersive element 2406 (such as a prism or a diffraction grating). The dispersive element 2406 may operate to select a narrow wavelength portion 2408 of the light emitted by the broad spectrum light source 2402, and the selected portion 2408 of the light may illuminate the tissue 2410. The light reflected from the tissue 2412 may be directed to a detector 2416 (for example, by means of a dichroic mirror 2414) and the intensity of the reflected light 2412 may be recorded. The wavelength of the light illuminating the tissue 2410 may be selected by the dispersive element 2406. In some aspects, the tissue 2410 may be illuminated only by a single narrow wavelength portion 2408 selected by the dispersive element 2406 from the light source 2402. In other aspects, the tissue 2410 may be scanned with a variety of narrow wavelength portions 2408 selected by the dispersive element 2406. In this manner, a spectroscopic analysis of the tissue 2410 may be obtained over a range of NIR wavelengths.

[1273] FIG. 155 depicts schematically one example of instrumentation 2430 for determining NIRS based on Fourier transform infrared imaging. In FIG. 155, a laser source emitting 2432 light in the near IR range 2434 illuminates a tissue sample 2440. The light reflected 2436 by the tissue 2440 is reflected 2442 by a mirror, such as a dichroic mirror 2444, to a beam splitter 2446. The beam splitter 2446 directs one portion of the light 2448 reflected 2436 by the tissue 2440 to a stationary mirror 2450 and one portion of the light 2452 reflected 2436 by the tissue 2440 a moving mirror 2454. The moving mirror 2454 may oscillate in position based on an affixed piezoelectric transducer activated by a sinusoidal voltage having a voltage frequency. The position of the moving mirror 2454 in space corresponds to the frequency of the sinusoidal activation voltage of the piezoelectric transducer. The light reflected from the moving mirror and the stationary mirror may be recombined 2458 at the beam splitter 2446 and directed to a detector 2456. Computational components may receive the signal output of the detector 2456 and perform a Fourier transform (in time) of the received signal. Because the wavelength of the light received from the moving mirror 2454 varies in time with respect to the wavelength of the light received from the stationary mirror 2450, the time-based Fourier transform of the recombined light corresponds to a wavelength-based Fourier transform of the recombined light 2458. In this manner, a wavelength-based spectrum of the light reflected from the tissue 2440 may be determined and spectral characteristics of the light reflected 2436 from the tissue 2440 may be obtained. Changes in the absorbance of the illumination in spectral components from the light reflected from the tissue 2440 may thus indicate the presence or absence of tissue having specific light absorbing properties (such as hemoglobin).

[1274] An alternative to near infrared light to determine hemoglobin oxygenation would be the use of monochromatic red light to determine the red light absorbance characteristics of hemoglobin. The absorbance characteristics of red light having a central wavelength of about 660 nm by the

hemoglobin may indicate if the hemoglobin is oxygenated (arterial blood) or deoxygenated (venous blood).

[1275] In some alternative surgical procedures, contrasting agents can be used to improve the data that is collected on oxygenation and tissue oxygen consumption. In one non-limiting example, NIRS techniques may be used in conjunction with a bolus injection of a near-IR contrast agent such as indocyanine green (ICG) which has a peak absorbance at about 800 nm. ICG has been used in some medical procedures to measure cerebral blood flow.

#### Vascular Imaging Using Laser Doppler Flowmetry

[1276] In one aspect, the characteristic of the light reflected and/or refracted from the surgical site may be a Doppler shift of the light wavelength from its illumination source.

[1277] Laser Doppler flowmetry may be used to visualize and characterized a flow of particles moving relative to an effectively stationary background. Thus, laser light scattered by moving particles, such as blood cells, may have a different wavelength than that of the original illuminating laser source. In contrast, laser light scattered by the effectively stationary background (for example, the vascular tissue) may have the same wavelength of that of the original illuminating laser source. The change in wavelength of the scattered light from the blood cells may reflect both the direction of the flow of the blood cells relative to the laser source as well as the blood cell velocity. FIGS. 156A-C illustrate the change in wavelength of light scattered from blood cells that may be moving away from (FIG. 156A) or towards (FIG. 156C) the laser light source.

[1278] In each of FIGS. 156A-C, the original illuminating light 2502 is depicted having a relative central wavelength of 0. It may be observed from FIG. 156A that light scattered from blood cells moving away from the laser source 2504 has a wavelength shifted by some amount 2506 to a greater wavelength relative to that of the laser source (and is thus red shifted). It may also be observed from FIG. 156C that light scattered from blood cells moving towards from the laser source 2508 has a wavelength shifted by some amount 2510 to a shorter wavelength relative to that of the laser source (and is thus blue shifted). The amount of wavelength shift (for example 2506 or 2510) may be dependent on the velocity of the motion of the blood cells. In some aspects, an amount of a red shift (2506) of some blood cells may be about the same as the amount of blue shift (2510) of some other blood cells. Alternatively, an amount of a red shift (2506) of some blood cells may differ from the amount of blue shift (2510) of some other blood cells. Thus, the velocity of the blood cells flowing away from the laser source as depicted in FIG. 156A may be less than the velocity of the blood cells flowing towards the laser source as depicted in FIG. 156C based on the relative magnitude of the wavelength shifts (2506 and 2510). In contrast, and as depicted in FIG. 156B, light scattered from tissue not moving relative to the laser light source (for example blood vessels 2512 or non-vascular tissue 2514) may not demonstrate any change in wavelength.

[1279] FIG. 157 depicts an aspect of instrumentation 2530 that may be used to detect a Doppler shift in laser light scattered from portions of a tissue 2540. Light 2534 originating from a laser 2532 may pass through a beam splitter 2544. Some portion of the laser light 2536 may be transmitted by the beam splitter 2544 and may illuminate tissue

**2540.** Another portion of the laser light may be reflected **2546** by the beam splitter **2544** to impinge on a detector **2550**. The light back-scattered **2542** by the tissue **2540** may be directed by the beam splitter **2544** and also impinge on the detector **2550**. The combination of the light **2534** originating from the laser **2532** with the light back-scattered **2542** by the tissue **2540** may result in an interference pattern detected by the detector **2550**. The interference pattern received by the detector **2550** may include interference fringes resulting from the combination of the light **2534** originating from the laser **2532** and the Doppler shifted (and thus wavelength shifted) light back-scattered **2452** from the tissue **2540**.

**[1280]** It may be recognized that back-scattered light **2542** from the tissue **2540** may also include back scattered light from boundary layers within the tissue **2540** and/or wavelength-specific light absorption by material within the tissue **2540**. As a result, the interference pattern observed at the detector **2550** may incorporate interference fringe features from these additional optical effects and may therefore confound the calculation of the Doppler shift unless properly analyzed.

**[1281]** FIG. **158** depicts some of these additional optical effects. It is well known that light traveling through a first optical medium having a first refractive index,  $n_1$ , may be reflected at an interface with a second optical medium having a second refractive index,  $n_2$ . The light transmitted through the second optical medium will have a transmission angle relative to the interface that differs from the angle of the incident light based on a difference between the refractive indices  $n_1$  and  $n_2$  (Snell's Law). FIG. **158** illustrates the effect of Snell's Law on light impinging on the surface of a multi-component tissue **2150**, as may be presented in a surgical field. The multi-component tissue **2150** may be composed of an outer tissue layer **2152** having a refractive index  $n_1$  and a buried tissue, such as a blood vessel having a vessel wall **2156**. The blood vessel wall **2156** may be characterized by a refractive index  $n_2$ . Blood may flow within the lumen of the blood vessel **2160**. In some aspects, it may be important during a surgical procedure to determine the position of the blood vessel **2160** below the surface **2154** of the outer tissue layer **2152** and to characterize the blood flow using Doppler shift techniques.

**[1282]** An incident laser light **2170a** may be used to probe for the blood vessel **2160** and may be directed on the top surface **2154** of the outer tissue layer **2152**. A portion **2172** of the incident laser light **2170a** may be reflected at the top surface **2154**. Another portion **2170b** of the incident laser light **2170a** may penetrate the outer tissue layer **2152**. The reflected portion **2172** at the top surface **2154** of the outer tissue layer **2152** has the same path length of the incident light **2170a**, and therefore has the same wavelength and phase of the incident light **2170a**. However, the portion **2170b** of light transmitted into the outer tissue layer **2152** will have a transmission angle that differs from the incidence angle of the light impinging on the tissue surface because the outer tissue layer **2152** has an index of refraction  $n_1$  that differs from the index of refraction of air.

**[1283]** If the portion of light transmitted through the outer tissue layer **2152** impinges on a second tissue surface **2158**, for example of the blood vessel wall **2156**, some portion **2174a,b** of light will be reflected back towards the source of the incident light **2170a**. The light thus reflected **2174a** at the interface between the outer tissue layer **2152** and the blood

vessel wall **2156** will have the same wavelength as the incident light **2170a**, but will be phase shifted due to the change in the light path length. Projecting the light reflected **2174a,b** from the interface between the outer tissue layer **2152** and the blood vessel wall **2156** along with the incident light on the sensor, will produce an interference pattern based on the phase difference between the two light sources.

**[1284]** Further, a portion of the incident light **2170c** may be transmitted through the blood vessel wall **2156** and penetrate into the blood vessel lumen **2160**. This portion of the incident light **2170c** may interact with the moving blood cells in the blood vessel lumen **2160** and may be reflected back **2176a-c** towards the source of the impinging light having a wavelength Doppler shifted according to the velocity of the blood cells, as disclosed above. The Doppler shifted light reflected **2176a-c** from the moving blood cells may be projected along with the incident light on the sensor, resulting in an interference pattern having a fringe pattern based on the wavelength difference between the two light sources.

**[1285]** In FIG. **158**, a light path **2178** is presented of light impinging on the red blood cells in the blood vessel lumen **2160** if there are no changes in refractive index between the emitted light and the light reflected by the moving blood cells. In this example, only a Doppler shift in the reflected light wavelength can be detected. However, the light reflected by the blood cells (**2176a-c**) may incorporate phase changes due to the variation in the tissue refractive indices in addition to the wavelength changes due to the Doppler Effect.

**[1286]** Thus, it may be understood that if the light sensor receives the incident light, the light reflected from one or more tissue interfaces (**2172**, and **2174a,b**) and the Doppler shifted light from the blood cells (**2176a-c**), the interference pattern thus produced on the light sensor may include the effects due to the Doppler shift (change in wavelength) as well as the effects due to the change in refractive index within the tissue (change in phase). As a result, a Doppler analysis of the light reflected by the tissue sample may produce erroneous results if the effects due to changes in the refractive index within the sample are not compensated for.

**[1287]** FIG. **159** illustrates an example of the effects on a Doppler analysis of light that impinge **2250** on a tissue sample to determine the depth and location of an underlying blood vessel. If there is no intervening tissue between the blood vessel and the tissue surface, the interference pattern detected at the sensor may be due primarily to the change in wavelength reflected from the moving blood cells. As a result, a spectrum **2252** derived from the interference pattern may generally reflect only the Doppler shift of the blood cells. However, if there is intervening tissue between the blood vessel and the tissue surface, the interference pattern detected at the sensor may be due to a combination of the change in wavelength reflected from the moving blood cells and the phase shift due to the refractive index of the intervening tissue. A spectrum **2254** derived from such an interference pattern, may result in the calculation of the Doppler shift that is confounded due to the additional phase change in the reflected light. In some aspects, if information regarding the characteristics (thickness and refractive index) of the intervening tissue is known, the resulting spectrum **2256** may be corrected to provide a more accurate calculation of the change in wavelength.

[1288] It is recognized that the tissue penetration depth of light is dependent on the wavelength of the light used. Thus, the wavelength of the laser source light may be chosen to detect particle motion (such as blood cells) at a specific range of tissue depth. FIGS. 160A-C depict schematically a means for detecting moving particles such as blood cells at a variety of tissue depths based on the laser light wavelength. As illustrated in FIG. 160A, a laser source 2340 may direct an incident beam of laser light 2342 onto a surface 2344 of a surgical site. A blood vessel 2346 (such as a vein or artery) may be disposed within the tissue 2348 at some depth  $\delta$  from the tissue surface. The penetration depth 2350 of a laser into a tissue 2348 may be dependent at least in part on the laser wavelength. Thus, laser light having a wavelength in the red range of about 635 nm to about 660 nm, may penetrate the tissue 2351a to a depth of about 1 mm. Laser light having a wavelength in the green range of about 520 nm to about 532 nm may penetrate the tissue 2351b to a depth of about 2-3 mm. Laser light having a wavelength in the blue range of about 405 nm to about 445 nm may penetrate the tissue 2351c to a depth of about 4 mm or greater. In the example depicted in FIGS. 160A-C, a blood vessel 2346 may be located at a depth  $\delta$  of about 2-3 mm below the tissue surface. Red laser light will not penetrate to this depth and thus will not detect blood cells flowing within this vessel. However, both green and blue laser light can penetrate this depth. Therefore, scattered green and blue laser light from the blood cells within the blood vessel 2346 may demonstrate a Doppler shift in wavelength.

[1289] FIG. 160B illustrates how a Doppler shift 2355 in the wavelength of reflected laser light may appear. The emitted light (or laser source light 2342) impinging on a tissue surface 2344 may have a central wavelength 2352. For example, light from a green laser may have a central wavelength 2352 within a range of about 520 nm to about 532 nm. The reflected green light may have a central wavelength 2354 shifted to a longer wavelength (red shifted) if the light was reflected from a particle such as a red blood cell that is moving away from the detector. The difference between the central wavelength 2352 of the emitted laser light and the central wavelength 2354 of the emitted laser light comprises the Doppler shift 2355.

[1290] As disclosed above with respect to FIGS. 158 and 159, laser light reflected from structures within a tissue 2348 may also show a phase shift in the reflected light due to changes in the index of refraction arising from changes in tissue structure or composition. The emitted light (or laser source light 2342) impinging on a tissue surface 2344 may have a first phase characteristic 2356. The reflected laser light may have a second phase characteristic 2358. It may be recognized that blue laser light that can penetrate tissue to a depth of about 4 mm or greater 2351c may encounter a greater variety of tissue structures than red laser light (about 1 mm 2351a) or green laser light (about 2-3 mm 2351b). Consequently, as illustrated in FIG. 160C, the phase shift 2358 of reflected blue laser light may be significant at least due to the depth of penetration.

[1291] FIG. 160D illustrates aspects of illuminating tissue by red 2360a, green 2360b and blue 2360c laser light in a sequential manner. In some aspects, a tissue may be probed by red 2360a, green 2360b and blue 2360c laser illumination in a sequential manner. In some alternative examples, one or more combinations of red 2360a, green 2360b, and blue 2360c laser light, as depicted in FIGS. 153D-153F and

disclosed above, may be used to illuminate the tissue according to a defined illumination sequence. 30D illustrates the effect of such illumination on a CMOS imaging sensor 2362a-d over time. Thus, at a first time  $t_1$ , the CMOS sensor 2362a may be illuminated by the red 2360a laser. At a second time  $t_2$  the CMOS sensor 2362b may be illuminated by the green 2360b laser. At a third time  $t_3$ , the CMOS sensor 2362c may be illuminated by the blue 2360c laser. The illumination cycle may then be repeated starting at a fourth time  $t_4$  in which the CMOS sensor 2362d may be illuminated by the red 2360a laser again. It may be recognized that sequential illumination of the tissue by laser illumination at differing wavelengths may permit a Doppler analysis at varying tissue depths over time. Although red 2360a, green 2360b and blue 2360c laser sources may be used to illuminate the surgical site, it may be recognized that other wavelengths outside of visible light (such as in the infrared or ultraviolet regions) may be used to illuminate the surgical site for Doppler analysis.

[1292] FIG. 161 illustrates an example of a use of Doppler imaging to detect the presence of blood vessels not otherwise viewable at a surgical site 2600. In FIG. 161, a surgeon may wish to excise a tumor 2602 found in the right superior posterior lobe 2604 of a lung. Because the lungs are highly vascular, care must be taken to identify only those blood vessels associated with the tumor and to seal only those vessels without compromising the blood flow to the non-affected portions of the lung. In FIG. 161, the surgeon has identified the margin 2606 of the tumor 2604. The surgeon may then cut an initial dissected area 2608 in the margin region 2606, and exposed blood vessels 2610 may be observed for cutting and sealing. The Doppler imaging detector 2620 may be used to locate and identify blood vessels not observable 2612 in the dissected area. An imaging system may receive data from the Doppler imaging detector 2620 for analysis and display of the data obtained from the surgical site 2600. In some aspects, the imaging system may include a display to illustrate the surgical site 2600 including a visible image of the surgical site 2600 along with an image overlay of the hidden blood vessels 2612 on the image of the surgical site 2600.

[1293] In the scenario disclosed above regarding FIG. 161, a surgeon wishes to sever blood vessels that supply oxygen and nutrients to a tumor while sparing blood vessels associated with non-cancerous tissue. Additionally, the blood vessels may be disposed at different depths in or around the surgical site 2600. The surgeon must therefore identify the position (depth) of the blood vessels as well as determine if they are appropriate for resection. FIG. 162 illustrates one method for identifying deep blood vessels based on a Doppler shift of light from blood cells flowing therethrough. As disclosed above, red laser light has a penetration depth of about 1 mm and green laser light has a penetration depth of about 2-3 mm. However, a blood vessel having a below-surface depth of 4 mm or more will be outside the penetration depths at these wavelengths. Blue laser light, however, can detect such blood vessels based on their blood flow.

[1294] FIG. 162 depicts the Doppler shift of laser light reflected from a blood vessel at a specific depth below a surgical site. The site may be illuminated by red laser light, green laser light, and blue laser light. The central wavelength 2630 of the illuminating light may be normalized to a relative central 3631. If the blood vessel lies at a depth of 4 or more mm below the surface of the surgical site, neither

the red laser light nor the green laser light will be reflected by the blood vessel. Consequently, the central wavelength 2632 of the reflected red light and the central wavelength 2634 of the reflected green light will not differ much from the central wavelength 2630 of the illuminating red light or green light, respectively. However, if the site is illuminated by blue laser light, the central wavelength 2638 of the reflected blue light 2636 will differ from the central wavelength 2630 of the illuminating blue light. In some instances, the amplitude of the reflected blue light 2636 may also be significantly reduced from the amplitude of the illuminating blue light. A surgeon may thus determine the presence of a deep lying blood vessel along with its approximate depth, and thereby avoiding the deep blood vessel during surface tissue dissection.

[1295] FIGS. 163 and 164 illustrates schematically the use of laser sources having differing central wavelengths (colors) for determining the approximate depth of a blood vessel beneath the surface of a surgical site. FIG. 163 depicts a first surgical site 2650 having a surface 2654 and a blood vessel 2656 disposed below the surface 2654. In one method, the blood vessel 2656 may be identified based on a Doppler shift of light impinging on the flow 2658 of blood cells within the blood vessel 2656. The surgical site 2650 may be illuminated by light from a number of lasers 2670, 2676, 2682, each laser being characterized by emitting light at one of several different central wavelengths. As noted above, illumination by a red laser 2670 can only penetrate tissue by about 1 mm. Thus, if the blood vessel 2656 was located at a depth of less than 1 mm 2672 below the surface 2654, the red laser illumination would be reflected 2674 and a Doppler shift of the reflected red illumination 2674 may be determined. Further, as noted above, illumination by a green laser 2676 can only penetrate tissue by about 2-3 mm. If the blood vessel 2656 was located at a depth of about 2-3 mm 2678 below the surface 2654, the green laser illumination would be reflected 2680 while the red laser illumination 2670 would not, and a Doppler shift of the reflected green illumination 2680 may be determined. However, as depicted in FIG. 163, the blood vessel 2656 is located at a depth of about 4 mm 2684 below the surface 2654. Therefore, neither the red laser illumination 2670 nor the green laser illumination 2676 would be reflected. Instead, only the blue laser illumination would be reflected 2686 and a Doppler shift of the reflected blue illumination 2686 may be determined.

[1296] In contrast to the blood vessel 2656 depicted in FIG. 163, the blood vessel 2656' depicted in FIG. 164 is located closer to the surface of the tissue at the surgical site. Blood vessel 2656' may also be distinguished from blood vessel 2656 in that blood vessel 2656' is illustrated to have a much thicker wall 2657. Thus, blood vessel 2656' may be an example of an artery while blood vessel 2656 may be an example of a vein because arterial walls are known to be thicker than venous walls. In some examples, arterial walls may have a thickness of about 1.3 mm. As disclosed above, red laser illumination 2670' can penetrate tissue to a depth of about 1 mm 2672'. Thus, even if a blood vessel 2656' is exposed at a surgical site (see 2610 at FIG. 161), red laser light that is reflected 2674' from the surface of the blood vessel 2656', may not be able to visualize blood flow 2658' within the blood vessel 2656' under a Doppler analysis due to the thickness of the blood vessel wall 2657. However, as disclosed above, green laser light impinging 2676' on the surface of a tissue may penetrate to a depth of about 2-3 mm

2678'. Further, blue laser light impinging 2682' on the surface of a tissue may penetrate to a depth of about 4 mm 2684'. Consequently, green laser light may be reflected 2680' from the blood cells flowing 2658' within the blood vessel 2656' and blue laser light may be reflected 2686' from the blood cells flowing 2658' within the blood vessel 2656'. As a result, a Doppler analysis of the reflected green light 2680' and reflected blue light 2686' may provide information regarding blood flow in near-surface blood vessel, especially the approximate depth of the blood vessel.

[1297] As disclosed above, the depth of blood vessels below the surgical site may be probed based on wavelength-dependent Doppler imaging. The amount of blood flow through such a blood vessel may also be determined by speckle contrast (interference) analysis. Doppler shift may indicate a moving particle with respect to a stationary light source. As disclosed above, the Doppler wavelength shift may be an indication of the velocity of the particle motion. Individual particles such as blood cells may not be separately observable. However, the velocity of each blood cell will produce a proportional Doppler shift. An interference pattern may be generated by the combination of the light back-scattered from multiple blood cells due to the differences in the Doppler shift of the back-scattered light from each of the blood cells. The interference pattern may be an indication of the number density of blood cells within a visualization frame. The interference pattern may be termed speckle contrast. Speckle contrast analysis may be calculated using a full frame 300x300 CMOS imaging array, and the speckle contrast may be directly related to the amount of moving particles (for example blood cells) interacting with the laser light over a given exposure period.

[1298] A CMOS image sensor may be coupled to a digital signal processor (DSP). Each pixel of the sensor may be multiplexed and digitized. The Doppler shift in the light may be analyzed by looking at the source laser light in comparison to the Doppler shifted light. A greater Doppler shift and speckle may be related to a greater number of blood cells and their velocity in the blood vessel.

[1299] FIG. 165 depicts an aspect of a composite visual display 2800 that may be presented to a surgeon during a surgical procedure. The composite visual display 2800 may be constructed by overlaying a white light image 2830 of the surgical site with a Doppler analysis image 2850.

[1300] In some aspects, the white light image 2830 may portray the surgical site 2832, one or more surgical incisions 2834, and the tissue 2836 readily visible within the surgical incision 2834. The white light image 2830 may be generated by illuminating 2840 the surgical site 2832 with a white light source 2838 and receiving the reflected white light 2842 by an optical detector. Although a white light source 2838 may be used to illuminate the surface of the surgical site, in one aspect, the surface of the surgical site may be visualized using appropriate combinations of red 2854, green 2856, and blue 2858 laser light as disclosed above with respect to FIGS. 153C-153F.

[1301] In some aspects, the Doppler analysis image 2850 may include blood vessel depth information along with blood flow information 2852 (from speckle analysis). As disclosed above, blood vessel depth and blood flow velocity may be obtained by illuminating the surgical site with laser light of multiple wavelengths, and determining the blood vessel depth and blood flow based on the known penetration depth of the light of a particular wavelength. In general, the



surgical site **2832** may be illuminated by light emitted by one or more lasers such as a red laser **2854**, a green laser **2856**, and a blue laser **2858**. A CMOS detector **2872** may receive the light reflected back (**2862**, **2866**, **2870**) from the surgical site **2832** and its surrounding tissue. The Doppler analysis image **2850** may be constructed **2874** based on an analysis of the multiple pixel data from the CMOS detector **2872**.

[**1302**] In one aspect, a red laser **2854** may emit red laser illumination **2860** on the surgical site **2832** and the reflected light **2862** may reveal surface or minimally subsurface structures. In one aspect, a green laser **2856** may emit green laser illumination **2864** on the surgical site **2832** and the reflected light **2866** may reveal deeper subsurface characteristics. In another aspect, a blue laser **2858** may emit blue laser illumination **2868** on the surgical site **2832** and the reflected light **2870** may reveal, for example, blood flow within deeper vascular structures. In addition, the speckle contrast analysis may present the surgeon with information regarding the amount and velocity of blood flow through the deeper vascular structures.

[**1303**] Although not depicted in FIG. **165**, it may be understood that the imaging system may also illuminate the surgical site with light outside of the visible range. Such light may include infrared light and ultraviolet light. In some aspects, sources of the infrared light or ultraviolet light may include broad-band wavelength sources (such as a tungsten source, a tungsten-halogen source, or a deuterium source). In some other aspects, the sources of the infrared or ultraviolet light may include narrow-band wavelength sources (IR diode lasers, UV gas lasers or dye lasers).

[**1304**] FIG. **166** is a flow chart **2900** of a method for determining a depth of a surface feature in a piece of tissue. An image acquisition system may illuminate **2910** a tissue with a first light beam having a first central frequency and receive **2912** a first reflected light from the tissue illuminated by the first light beam. The image acquisition system may then calculate **2914** a first Doppler shift based on the first light beam and the first reflected light. The image acquisition system may then illuminate **2916** the tissue with a second light beam having a second central frequency and receive **2918** a second reflected light from the tissue illuminated by the second light beam. The image acquisition system may then calculate **2920** a second Doppler shift based on the second light beam and the second reflected light. The image acquisition system may then calculate **2922** a depth of a tissue feature based at least in part on the first central wavelength, the first Doppler shift, the second central wavelength, and the second Doppler shift. In some aspects, the tissue features may include the presence of moving particles, such as blood cells moving within a blood vessel, and a direction and velocity of flow of the moving particles. It may be understood that the method may be extended to include illumination of the tissue by any one or more additional light beams. Further, the system may calculate an image comprising a combination of an image of the tissue surface and an image of the structure disposed within the tissue.

[**1305**] In some aspects, multiple visual displays may be used. For example, a 3D display may provide a composite image displaying the combined white light (or an appropriate combination of red, green, and blue laser light) and laser Doppler image. Additional displays may provide only the white light display or a displaying showing a composite white light display and an NIRS display to visualize only the

blood oxygenation response of the tissue. However, the NIRS display may not be required every cycle allowing for response of tissue.

#### Subsurface Tissue Characterization Using Multispectral OCT

[**1306**] During a surgical procedure, the surgeon may employ “smart” surgical devices for the manipulation of tissue. Such devices may be considered “smart” in that they include automated features to direct, control, and/or vary the actions of the devices based parameters relevant to their uses. The parameters may include the type and/or composition of the tissue being manipulated. If the type and/or composition of the tissue being manipulated is unknown, the actions of the smart devices may be inappropriate for the tissue being manipulated. As a result, tissues may be damaged or the manipulation of the tissue may be ineffective due to inappropriate settings of the smart device.

[**1307**] The surgeon may manually attempt to vary the parameters of the smart device in a trial-and-error manner, resulting in an inefficient and lengthy surgical procedure.

[**1308**] Therefore, it is desirable to have a surgical visualization system that can probe tissue structures underlying a surgical site to determine their structural and compositional characteristics, and to provide such data to smart surgical instruments being used in a surgical procedure.

[**1309**] Some aspects of the present disclosure further provide for a control circuit configured to control the illumination of a surgical site using one or more illumination sources such as laser light sources and to receive imaging data from one or more image sensors. In some aspects, the present disclosure provides for a non-transitory computer readable medium storing computer readable instructions that, when executed, cause a device to characterize structures below the surface at a surgical site and determine the depth of the structures below the surface of the tissue.

[**1310**] In some aspects, a surgical image acquisition system may comprise a plurality of illumination sources wherein each illumination source is configured to emit light having a specified central wavelength, a light sensor configured to receive a portion of the light reflected from a tissue sample when illuminated by the one or more of the plurality of illumination sources, and a computing system. The computing system may be configured to receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources, calculate structural data related to a characteristic of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the illumination sources, and transmit the structural data related to the characteristic of the structure to be received by a smart surgical device. In some aspects, the characteristic of the structure is a surface characteristic or a structure composition.

[**1311**] In one aspect, a surgical system may include multiple laser light sources and may receive laser light reflected from a tissue. The light reflected from the tissue may be used by the system to calculate surface characteristics of components disposed within the tissue. The characteristics of the components disposed within the tissue may include a composition of the components and/or a metric related to surface irregularities of the components.

[**1312**] In one aspect, the surgical system may transmit data related to the composition of the components and/or

metrics related to surface irregularities of the components to a second instrument to be used on the tissue to modify the control parameters of the second instrument.

[1313] In some aspects, the second device may be an advanced energy device and the modifications of the control parameters may include a clamp pressure, an operational power level, an operational frequency, and a transducer signal amplitude.

[1314] As disclosed above, blood vessels may be detected under the surface of a surgical site base on the Doppler shift in light reflected by the blood cells moving within the blood vessels.

[1315] Laser Doppler flowmetry may be used to visualize and characterized a flow of particles moving relative to an effectively stationary background. Thus, laser light scattered by moving particles, such as blood cells, may have a different wavelength than that of the original illuminating laser source. In contrast, laser light scattered by the effectively stationary background (for example, the vascular tissue) may have the same wavelength of that of the original illuminating laser source. The change in wavelength of the scattered light from the blood cells may reflect both the direction of the flow of the blood cells relative to the laser source as well as the blood cell velocity. As previously disclosed, FIGS. 156A-C illustrate the change in wavelength of light scattered from blood cells that may be moving away from (FIG. 156A) or towards (FIG. 156C) the laser light source.

[1316] In each of FIGS. 156A-C, the original illuminating light 2502 is depicted having a relative central wavelength of 0. It may be observed from FIG. 156A that light scattered from blood cells moving away from the laser source 2504 has a wavelength shifted by some amount 2506 to a greater wavelength relative to that of the laser source (and is thus red shifted). It may also be observed from FIG. 154C that light scattered from blood cells moving towards from the laser source 2508 has a wavelength shifted by some amount 2510 to a shorter wavelength relative to that of the laser source (and is thus blue shifted). The amount of wavelength shift (for example 2506 or 2510) may be dependent on the velocity of the motion of the blood cells. In some aspects, an amount of a red shift (2506) of some blood cells may be about the same as the amount of blue shift (2510) of some other blood cells. Alternatively, an amount of a red shift (2506) of some blood cells may differ from the amount of blue shift (2510) of some other blood cells. Thus, the velocity of the blood cells flowing away from the laser source as depicted in FIG. 154A may be less than the velocity of the blood cells flowing towards the laser source as depicted in FIG. 156C based on the relative magnitude of the wavelength shifts (2506 and 2510). In contrast, and as depicted in FIG. 156B, light scattered from tissue not moving relative to the laser light source (for example blood vessels 2512 or non-vascular tissue 2514) may not demonstrate any change in wavelength.

[1317] As previously disclosed, FIG. 157 depicts an aspect of instrumentation 2530 that may be used to detect a Doppler shift in laser light scattered from portions of a tissue 2540. Light 2534 originating from a laser 2532 may pass through a beam splitter 2544. Some portion of the laser light 2536 may be transmitted by the beam splitter 2544 and may illuminate tissue 2540. Another portion of the laser light may be reflected 2546 by the beam splitter 2544 to impinge on a detector 2550. The light back-scattered 2542 by the tissue

2540 may be directed by the beam splitter 2544 and also impinge on the detector 2550. The combination of the light 2534 originating from the laser 2532 with the light back-scattered 2542 by the tissue 2540 may result in an interference pattern detected by the detector 2550. The interference pattern received by the detector 2550 may include interference fringes resulting from the combination of the light 2534 originating from the laser 2532 and the Doppler shifted (and thus wavelength shifted) light back-scattered 2452 from the tissue 2540.

[1318] It may be recognized that back-scattered light 2542 from the tissue 2540 may also include back scattered light from boundary layers within the tissue 2540 and/or wavelength-specific light absorption by material within the tissue 2540. As a result, the interference pattern observed at the detector 2550 may incorporate interference fringe features from these additional optical effects and may therefore confound the calculation of the Doppler shift unless properly analyzed.

[1319] It may be recognized that light reflected from the tissue may also include back scattered light from boundary layers within the tissue and/or wavelength-specific light absorption by material within the tissue. As a result, the interference pattern observed at the detector may incorporate fringe features that may confound the calculation of the Doppler shift unless properly analyzed.

[1320] As previously disclosed, FIG. 158 depicts some of these additional optical effects. It is well known that light traveling through a first optical medium having a first refractive index,  $n_1$ , may be reflected at an interface with a second optical medium having a second refractive index,  $n_2$ . The light transmitted through the second optical medium will have a transmission angle relative to the interface that differs from the angle of the incident light based on a difference between the refractive indices  $n_1$  and  $n_2$  (Snell's Law). FIG. 156 illustrates the effect of Snell's Law on light impinging on the surface of a multi-component tissue 2150, as may be presented in a surgical field. The multi-component tissue 2150 may be composed of an outer tissue layer 2152 having a refractive index  $n_1$  and a buried tissue, such as a blood vessel having a vessel wall 2156. The blood vessel wall 2156 may be characterized by a refractive index  $n_2$ . Blood may flow within the lumen of the blood vessel 2160. In some aspects, it may be important during a surgical procedure to determine the position of the blood vessel 2160 below the surface 2154 of the outer tissue layer 2152 and to characterize the blood flow using Doppler shift techniques.

[1321] An incident laser light 2170a may be used to probe for the blood vessel 2160 and may be directed on the top surface 2154 of the outer tissue layer 2152. A portion 2172 of the incident laser light 2170a may be reflected at the top surface 2154. Another portion 2170b of the incident laser light 2170a may penetrate the outer tissue layer 2152. The reflected portion 2172 at the top surface 2154 of the outer tissue layer 2152 has the same path length of the incident light 2170a, and therefore has the same wavelength and phase of the incident light 2170a. However, the portion 2170b of light transmitted into the outer tissue layer 2152 will have a transmission angle that differs from the incidence angle of the light impinging on the tissue surface because the outer tissue layer 2152 has an index of refraction  $n_1$  that differs from the index of refraction of air.

[1322] If the portion of light transmitted through the outer tissue layer 2152 impinges on a second tissue surface 2158,

for example of the blood vessel wall **2156**, some portion **2174a,b** of light will be reflected back towards the source of the incident light **2170a**. The light thus reflected **2174a** at the interface between the outer tissue layer **2152** and the blood vessel wall **2156** will have the same wavelength as the incident light **2170a**, but will be phase shifted due to the change in the light path length. Projecting the light reflected **2174a,b** from the interface between the outer tissue layer **2152** and the blood vessel wall **2156** along with the incident light on the sensor, will produce an interference pattern based on the phase difference between the two light sources.

[1323] Further, a portion of the incident light **2170c** may be transmitted through the blood vessel wall **2156** and penetrate into the blood vessel lumen **2160**. This portion of the incident light **2170c** may interact with the moving blood cells in the blood vessel lumen **2160** and may be reflected back **2176a-c** towards the source of the impinging light having a wavelength Doppler shifted according to the velocity of the blood cells, as disclosed above. The Doppler shifted light reflected **2176a-c** from the moving blood cells may be projected along with the incident light on the sensor, resulting in an interference pattern having a fringe pattern based on the wavelength difference between the two light sources.

[1324] In FIG. **158**, a light path **2178** is presented of light impinging on the red blood cells in the blood vessel lumen **2160** if there are no changes in refractive index between the emitted light and the light reflected by the moving blood cells. In this example, only a Doppler shift in the reflected light wavelength can be detected. However, the light reflected by the blood cells (**2176a-c**) may incorporate phase changes due to the variation in the tissue refractive indices in addition to the wavelength changes due to the Doppler Effect.

[1325] Thus, it may be understood that if the light sensor receives the incident light, the light reflected from one or more tissue interfaces (**2172**, and **2174a,b**) and the Doppler shifted light from the blood cells (**2176a-c**), the interference pattern thus produced on the light sensor may include the effects due to the Doppler shift (change in wavelength) as well as the effects due to the change in refractive index within the tissue (change in phase). As a result, a Doppler analysis of the light reflected by the tissue sample may produce erroneous results if the effects due to changes in the refractive index within the sample are not compensated for.

[1326] As previously disclosed, FIG. **159** illustrates an example of the effects on a Doppler analysis of light that impinge **2250** on a tissue sample to determine the depth and location of an underlying blood vessel. If there is no intervening tissue between the blood vessel and the tissue surface, the interference pattern detected at the sensor may be due primarily to the change in wavelength reflected from the moving blood cells. As a result, a spectrum **2252** derived from the interference pattern may generally reflect only the Doppler shift of the blood cells. However, if there is intervening tissue between the blood vessel and the tissue surface, the interference pattern detected at the sensor may be due to a combination of the change in wavelength reflected from the moving blood cells and the phase shift due to the refractive index of the intervening tissue. A spectrum **2254** derived from such an interference pattern, may result in the calculation of the Doppler shift that is confounded due to the additional phase change in the reflected light. In some aspects, if information regarding the characteristics (thick-

ness and refractive index) of the intervening tissue is known, the resulting spectrum **2256** may be corrected to provide a more accurate calculation of the change in wavelength.

[1327] It may be recognized that the phase shift in the reflected light from a tissue may provide additional information regarding underlying tissue structures, regardless of Doppler effects.

[1328] FIG. **167** illustrates that the location and characteristics of non-vascular structures may be determined based on the phase difference between the incident light **2372** and the light reflected from the deep tissue structures (**2374**, **2376**, **2378**). As noted above, the penetration depth of light impinging on a tissue is dependent on the wavelength of the impinging illumination. Red laser light (having a wavelength in the range of about 635 nm to about 660 nm) may penetrate the tissue to a depth of about 1 mm. Green laser light (having a wavelength in the range of about 520 nm to about 532 nm) may penetrate the tissue to a depth of about 2-3 mm. Blue laser light (having a wavelength in the range of about 405 nm to about 445 nm) may penetrate the tissue to a depth of about 4 mm or greater. In one aspect, an interface **2381a** between two tissues differing in refractive index that is located less than or about 1 mm below a tissue surface **2380** may reflect **2374** red, green, or blue laser light. The phase of the reflected light **2374** may be compared to the incident light **2372** and thus the difference in the refractive index of the tissues at the interface **2381a** may be determined. In another aspect, an interface **2381b** between two tissues differing in refractive index that is located between 2 and 3 mm **2381b** below a tissue surface **2380** may reflect **2376** green or blue laser light, but not red light. The phase of the reflected light **2376** may be compared to the incident light **2372** and thus the difference in the refractive index of the tissues at the interface **2381b** may be determined. In yet another aspect, an interface **2381c** between two tissues differing in refractive index that is located between 3 and 4 mm **2381c** below a tissue surface **2380** may reflect **2378** only blue laser light, but not red or green light. The phase of the reflected light **2378** may be compared to the incident light **2372** and thus the difference in the refractive index of the tissues at the interface **2381c** may be determined.

[1329] A phase interference measure of a tissue illuminated by light having different wavelengths may therefore provide information regarding the relative indices of refraction of the reflecting tissue as well as the depth of the tissue. The indices of refraction of the tissue may be assessed using the multiple laser sources and their intensity, and thereby relative indices of refraction may be calculated for the tissue. It is recognized that different tissues may have different refractive indices. For example, the refractive index may be related to the relative composition of collagen and elastin in a tissue or the amount of hydration of the tissue. Therefore, a technique to measure relative tissue index of refraction may result in the identification of a composition of the tissue.

[1330] In some aspects, smart surgical instruments include algorithms to determine parameters associated with the function of the instruments. One non-limiting example of such parameters may be the pressure of an anvil against a tissue for a smart stapling device. The amount of pressure of an anvil against a tissue may depend on the type and composition of the tissue. For example, less pressure may be required to staple a highly compressive tissue, while a greater amount of pressure may be required to staple a more

non-compressive tissue. Another non-limiting example of a parameter associated with a smart surgical device may include a rate of firing of an i-beam knife to cut the tissue. For example, a stiff tissue may require more force and a slower cutting rate than a less stiff tissue. Another non-limiting example of such parameters may be the amount of current provided to an electrode in a smart cauterizing or RF sealing device. Tissue composition, such as percent tissue hydration, may determine an amount of current necessary to heat seal the tissue. Yet another non-limiting example of such parameters may be the amount of power provided to an ultrasonic transducer of a smart ultrasound cutting device or the driving frequency of the cutting device. A stiff tissue may require more power for cutting, and contact of the ultrasonic cutting tool with a stiff tissue may shift the resonance frequency of the cutter.

[1331] It may be recognized that a tissue visualization system that can identify tissue type and depth may provide such data to one or more smart surgical devices. The identification and location data may then be used by the smart surgical devices to adjust one or more of their operating parameters thereby allowing them to optimize their manipulation of the tissue. It may be understood that an optical method to characterize a type of tissue may permit automation of the operating parameters of the smart surgical devices. Such automation of the operation of smart surgical instruments may be preferable to relying on human estimation to determine the operational parameters of the instruments.

[1332] In one aspect, Optical Coherence Tomography (OCT) is a technique that can visual subsurface tissue structures based on the phase difference between an illuminating light source, and light reflected from structures located within the tissue. FIG. 168 depicts schematically one example of instrumentation 2470 for Optical Coherence Tomography. In FIG. 168, a laser source 2472 may emit light 2482 according to any optical wavelength of interest (red, green, blue, infrared, or ultraviolet). The light 2482 may be directed to a beam splitter 2486. The beam splitter 2486 directs one portion of the light 2488 to a tissue sample 2480. The beam splitter 2486 may also direct a portion of the light 2492 to a stationary reference mirror 2494. The light reflected from the tissue sample 2480 and from the stationary mirror 2494 may be recombined 2498 at the beam splitter 2486 and directed to a detector 2496. The phase difference between the light from the reference mirror 2494 and from the tissue sample 2480 may be detected at the detector 2496 as an interference pattern. Appropriate computing devices may then calculate phase information from the interference pattern. Additional computation may then provide information regarding structures below the surface of the tissue sample. Additional depth information may also be obtained by comparing the interference patterns generated from the sample when illuminated at different wavelengths of laser light.

[1333] As disclosed above, depth information regarding subsurface tissue structures may be ascertained from a combination of laser light wavelength and the phase of light reflected from a deep tissue structure. Additionally, local tissue surface inhomogeneity may be ascertained by comparing the phase as well as amplitude difference of light reflected from different portions of the same sub-surface tissues. Measurements of a difference in the tissue surface properties at a defined location compared to those at a

neighboring location may be indicative of adhesions, disorganization of the tissue layers, infection, or a neoplasm in the tissue being probed.

[1334] FIG. 169 illustrates this effect. The surface characteristics of a tissue determine the angle of reflection of light impinging on the surface. A smooth surface 2551a reflects the light essentially with the same spread 2544 as the light impinging on the surface 2542 (specular reflection). Consequently, the amount of light received by a light detector having a known fixed aperture may effectively receive the entire amount of light reflected 2544 from the smooth surface 2551a. However, increased surface roughness at a tissue surface may result in an increase spread in the reflected light with respect to the incident light (diffuse reflection).

[1335] Some amount of the reflected light 2546 from a tissue surface having some amount of surface irregularities 2551b will fall outside the fixed aperture of the light detector due to the increased spread of the reflected light 2546. As a result, the light detector will detect less light (shown in FIG. 169 as a decrease in the amplitude of the reflected light signal 2546). It may be understood that the amount of reflected light spread will increase as the surface roughness of a tissue increases. Thus, as depicted in FIG. 169, the amplitude of light reflected 2548 from a surface 2551c having significant surface roughness may have a smaller amplitude than the light reflected 2544 from a smooth surface 2551a, or light reflected 2546 from a surface having only a moderate amount of surface roughness 2551b. Therefore, in some aspects, a single laser source may be used to investigate the quality of a tissue surface or subsurface by comparing the optical properties of reflected light from the tissue with the optical properties of reflected light from adjacent surfaces.

[1336] In other aspects, light from multiple laser sources (for example, lasers emitting light having different central wavelengths) may be used sequentially to probe tissue surface characteristics at a variety of depths below the surface 2550. As disclosed above (with reference to FIG. 167), the absorbance profile of a laser light in a tissue is dependent on the central wavelength of the laser light. Laser light having a shorter (more blue) central wavelength can penetrate tissue deeper than laser light having a longer (more red) central wavelength. Therefore, measurements related to light diffuse reflection made at different light wavelengths can indicate both an amount of surface roughness as well as the depth of the surface being measured.

[1337] FIG. 170 illustrates one method of displaying image processing data related to a combination of tissue visualization modalities. Data used in the display may be derived from image phase data related to tissue layer composition, image intensity (amplitude) data related to tissue surface features, and image wavelength data related to tissue mobility (such as blood cell transport) as well as tissue depth. As one example, light emitted by a laser in the blue optical region 2562 may impinge on blood flowing at a depth of about 4 mm below the surface of the tissue. The reflected light 2564 may be red shifted due to the Doppler effect of the blood flow. As a result, information may be obtained regarding the existence of a blood vessel and its depth below the surface.

[1338] In another example, a layer of tissue may lie at a depth of about 2-3 mm below the surface of the surgical site. This tissue may include surface irregularities indicative of

scarring or other pathologies. Emitted red light **2572** may not penetrate to the 2-3 mm depth, so consequently, the reflected red light **2580** may have about the same amplitude of the emitted red light **2572** because it is unable to probe structures more than 1 mm below the top surface of the surgical site. However, green light reflected from the tissue **2578** may reveal the existence of the surface irregularities at that depth in that the amplitude of the reflected green light **2578** may be less than the amplitude of the emitted green light **2570**. Similarly, blue light reflected from the tissue **2574** may reveal the existence of the surface irregularities at that depth in that the amplitude of the reflected blue light **2574** may be less than the amplitude of the emitted blue light **2562**. In one example of an image processing step, the image **2582** may be smoothed using a moving window filter **2584** to reduce inter-pixel noise as well as reduce small local tissue anomalies **2586** that may hide more important features **2588**.

[1339] FIGS. **171A-C** illustrate several aspects of displays that may be provided to a surgeon for a visual identification of surface and sub-surface structures of a tissue in a surgical site. FIG. **171A** may represent a surface map of the surgical site with color coding to indicate structures located at varying depths below the surface of the surgical site. FIG. **171B** depicts an example of one of several horizontal slices through the tissue at varying depths, which may be color coded to indicate depth and further include data associated with differences in tissue surface anomalies (for example, as displayed in a 3D bar graph). FIG. **171C** depicts yet another visual display in which surface irregularities as well as Doppler shift flowmetry data may indicate sub-surface vascular structures as well as tissue surface characteristics.

[1340] FIG. **172** is a flow chart **2950** of a method for providing information related to a characteristic of a tissue to a smart surgical instrument. An image acquisition system may illuminate **2960** a tissue with a first light beam having a first central frequency and receive **2962** a first reflected light from the tissue illuminated by the first light beam. The image acquisition system may then calculate **2964** a first tissue surface characteristic at a first depth based on the first emitted light beam and the first reflected light from the tissue. The image acquisition system may then illuminate **2966** the tissue with a second light beam having a second central frequency and receive **2968** a second reflected light from the tissue illuminated by the second light beam. The image acquisition system may then calculate **2970** a second tissue surface characteristic at a second depth based on the second emitted light beam and the second reflected light from the tissue. Tissue features that may include a tissue type, a tissue composition, and a tissue surface roughness metric may be determined from the first central light frequency, the second central light frequency, the first reflected light from the tissue, and the second reflected light from the tissue. The tissue characteristic may be used to calculate **2972** one or more parameters related to the function of a smart surgical instrument such as jaw pressure, power to effect tissue cauterization, or current amplitude and/or frequency to drive a piezoelectric actuator to cut a tissue. In some additional examples, the parameter may be transmitted **2974** either directly or indirectly to the smart surgical instrument which may modify its operating characteristics in response to the tissue being manipulated.

#### Multifocal Minimally Invasive Camera

[1341] In a minimally invasive procedure, e.g., laparoscopic, a surgeon may visualize the surgical site using imaging instruments including a light source and a camera. The imaging instruments may allow the surgeon to visualize the end effector of a surgical device during the procedure. However, the surgeon may need to visualize tissue away from the end effector to prevent unintended damage during the surgery. Such distant tissue may lie outside the field of view of the camera system when focused on the end effector. The imaging instrument may be moved in order to change the field of view of the camera, but it may be difficult to return the camera system back to its original position after being moved.

[1342] The surgeon may attempt to move the imaging system within the surgical site to visualize different portions of the site during the procedure. Repositioning of the imaging system is time consuming and the surgeon is not guaranteed to visualize the same field of view of the surgical site when the imaging system is returned to its original location.

[1343] It is therefore desirable to have a medical imaging visualization system that can provide multiple fields of view of the surgical site without the need to reposition the visualization system. Medical imaging devices include, without limitation, laparoscopes, endoscopes, thoroscopes, and the like, as described herein. In some aspects, a single display system may display each of the multiple fields of view of the surgical site at about the same time. The display of each of the multiple fields of view may be independently updated depending on a display control system composed of one or more hardware modules, one or more software modules, one or more firmware modules, or any combination or combinations thereof.

[1344] Some aspects of the present disclosure further provide for a control circuit configured to control the illumination of a surgical site using one or more illumination sources such as laser light sources and to receive imaging data from one or more image sensors. In some aspects, the control circuit may be configured to control the operation of one or more light sensor modules to adjust a field of view. In some aspects, the present disclosure provides for a non-transitory computer readable medium storing computer readable instructions that, when executed, cause a device to adjust one or more components of the one or more light sensor modules and to process an image from each of the one or more light sensor modules.

[1345] An aspect of a minimally invasive image acquisition system may comprise a plurality of illumination sources wherein each illumination source is configured to emit light having a specified central wavelength, a first light sensing element having a first field of view and configured to receive illumination reflected from a first portion of the surgical site when the first portion of the surgical site is illuminated by at least one of the plurality of illumination sources, a second light sensing element having a second field of view and configured to receive illumination reflected from a second portion of the surgical site when the second portion of the surgical site is illuminated by at least one of the plurality of illumination sources, wherein the second field of view overlaps at least a portion of the first field of view; and a computing system.

[1346] The computing system may be configured to receive data from the first light sensing element, receive data

from the second light sensing element, compute imaging data based on the data received from the first light sensing element and the data received from the second light sensing element, and transmit the imaging data for receipt by a display system.

[1347] A variety of surgical visualization systems have been disclosed above. Such systems provide for visualizing tissue and sub-tissue structures that may be encountered during one or more surgical procedures. Non-limiting examples of such systems may include: systems to determine the location and depth of subsurface vascular tissue such as veins and arteries; systems to determine an amount of blood flowing through the subsurface vascular tissue; systems to determine the depth of non-vascular tissue structures; systems to characterize the composition of such non-vascular tissue structures; and systems to characterize one or more surface characteristics of such tissue structures.

[1348] It may be recognized that a single surgical visualization system may incorporate components of any one or more of these visualization modalities. FIGS. 152A-D depict some examples of such a surgical visualization system 2108.

[1349] As disclosed above, in one non-limiting aspect, a surgical visualization system 2108 may include an imaging control unit 2002 and a hand unit 2020. The hand unit 2020 may include a body 2021, a camera scope cable 2015 attached to the body 2021, and an elongated camera probe 2024. The elongated camera probe 2024 may also terminate at its distal end with at least one window. In some non-limiting examples, a light sensor 2030 may be incorporated in the hand unit 2020, for example either in the body of the hand unit 2032b, or at a distal end 2032a of the elongated camera probe, as depicted in FIG. 152C. The light sensor 2030 may be fabricated using a CMOS sensor array or a CCD sensor array. As illustrated in FIG. 153C, a typical CMOS or CCD sensor array may generate an RGB (red-green-blue) image from light impinging on a mosaic of sensor elements, each sensor element having one of a red, green, or blue optical filter.

[1350] Alternatively, the illumination of the surgical site may be cycled among visible illumination sources as depicted in FIG. 160D. In some example, the illumination sources may include any one or more of a red laser 2360a, a green laser 2360b, or a blue laser 2360c. In some non-limiting examples, a red laser 2360a light source may source illumination having a peak wavelength that may range between 635 nm and 660 nm, inclusive. Non-limiting examples of a red laser peak wavelength may include about 635 nm, about 640 nm, about 645 nm, about 650 nm, about 655 nm, about 660 nm, or any value or range of values therebetween. In some non-limiting examples, a green laser 2360b light source may source illumination having a peak wavelength that may range between 520 nm and 532 nm, inclusive. Non-limiting examples of a red laser peak wavelength may include about 520 nm, about 522 nm, about 524 nm, about 526 nm, about 528 nm, about 530 nm, about 532 nm, or any value or range of values therebetween. In some non-limiting examples, the blue laser 2360c light source may source illumination having a peak wavelength that may range between 405 nm and 445 nm, inclusive. Non-limiting examples of a blue laser peak wavelength may include about 405 nm, about 410 nm, about 415 nm, about 420 nm, about 425 nm, about 430 nm, about 435 nm, about 440 nm, about 445 nm, or any value or range of values therebetween.

[1351] Additionally, illumination of the surgical site may be cycled to include non-visible illumination sources that may supply infrared or ultraviolet illumination. In some non-limiting examples, an infrared laser light source may source illumination having a peak wavelength that may range between 750 nm and 3000 nm, inclusive. Non-limiting examples of an infrared laser peak wavelength may include about 750 nm, about 1000 nm, about 1250 nm, about 1500 nm, about 1750 nm, about 2000 nm, about 2250 nm, about 2500 nm, about 2750 nm, 3000 nm, or any value or range of values therebetween. In some non-limiting examples, an ultraviolet laser light source may source illumination having a peak wavelength that may range between 200 nm and 360 nm, inclusive. Non-limiting examples of an ultraviolet laser peak wavelength may include about 200 nm, about 220 nm, about 240 nm, about 260 nm, about 280 nm, about 300 nm, about 320 nm, about 340 nm, about 360 nm, or any value or range of values therebetween.

[1352] The outputs of the sensor array under the different illumination wavelengths may be combined to form the RGB image, for example, if the illumination cycle time is sufficiently fast and the laser light is in the visible range. FIGS. 173A and 173B illustrate a multi-pixel light sensor receiving by light reflected by a tissue illuminated, for example, by sequential exposure to red, green, blue, infrared, (FIG. 173A) or red, green, blue, and ultraviolet laser light sources (FIG. 173B).

[1353] FIG. 174A depicts the distal end of a flexible elongated camera probe 2120 having a flexible camera probe shaft 2122 and a single light sensor module 2124 disposed at the distal end 2123 of the flexible camera probe shaft 2122. In some non-limiting examples, the flexible camera probe shaft 2122 may have an outer diameter of about 5 mm. The outer diameter of the flexible camera probe shaft 2122 may depend on geometric factors that may include, without limitation, the amount of allowable bend in the shaft at the distal end 2123. As depicted in FIG. 174A, the distal end 2123 of the flexible camera probe shaft 2122 may bend about 90° with respect to a longitudinal axis of an un-bent portion of the flexible camera probe shaft 2122 located at a proximal end of the elongated camera probe 2120. It may be recognized that the distal end 2123 of the flexible camera probe shaft 2122 may bend any appropriate amount as may be required for its function. Thus, as non-limiting examples, the distal end 2123 of the flexible camera probe shaft 2122 may bend any amount between about 0° and about 90°. Non-limiting examples of the bend angle of the distal end 2123 of the flexible camera probe shaft 2122 may include about 0°, about 10°, about 20°, about 30°, about 40°, about 50°, about 60°, about 70°, about 80°, about 90°, or any value or range of values therebetween. In some examples, the bend angle of the distal end 2123 of the flexible camera probe shaft 2122 may be set by a surgeon or other health care professional prior to or during a surgical procedure. In some other example, the bend angle of the distal end 2123 of the flexible camera probe shaft 2122 may be a fixed angle set at a manufacturing site.

[1354] The single light sensor module 2124 may receive light reflected from the tissue when illuminated by light emitted by one or more illumination sources 2126 disposed at the distal end of the elongated camera probe. In some examples, the light sensor module 2124 may be a 4 mm sensor module such as 4 mm mount 2136b, as depicted in FIG. 152D. It may be recognized that the light sensor

module **2124** may have any appropriate size for its intended function. Thus, the light sensor module **2124** may include a 5.5 mm mount **2136a**, a 2.7 mm mount **2136c**, or a 2 mm mount **2136d** as depicted in FIG. 152D.

[1355] It may be recognized that the one or more illumination sources **2126** may include any number of illumination sources **2126** including, without limitation, one illumination source, two illumination sources, three illumination sources, four illumination sources, or more than four illumination sources. It may be further understood that each illumination source may source illumination having any central wavelength including a central red illumination wavelength, a central green illumination wavelength, a central blue illumination wavelength, a central infrared illumination wavelength, a central ultraviolet illumination wavelength, or any other wavelength. In some examples, the one or more illumination sources **2126** may include a white light source, which may illuminate tissue with light having wavelengths that may span the range of optical white light from about 390 nm to about 700 nm.

[1356] FIG. 174B depicts the distal end **2133** of an alternative elongated camera probe **2130** having multiple light sensor modules, for example the two light sensor modules **2134a,b**, each disposed at the distal end **2133** of the elongated camera probe **2130**. In some non-limiting examples, the alternative elongated camera probe **2130** may have an outer diameter of about 7 mm. In some examples, the light sensor modules **2134a,b** may each comprise a 4 mm sensor module, similar to light sensor module **2124** in FIG. 174A. Alternatively, each of the light sensor modules **2134a,b** may comprise a 5.5 mm light sensor module, a 2.7 mm light sensor module, or a 2 mm light sensor module as depicted in FIG. 152D. In some examples, both light sensor modules **2134a,b** may have the same size. In some examples, the light sensor modules **2134a,b** may have different sizes. As one non-limiting example, an alternative elongated camera probe **2130** may have a first 4 mm light sensor and two additional 2 mm light sensors. In some aspects, a visualization system may combine the optical outputs from the multiple light sensor modules **2134a,b** to form a 3D or quasi-3D image of the surgical site. In some other aspects, the outputs of the multiple light sensor modules **2134a,b** may be combined in such a manner as to enhance the optical resolution of the surgical site, which may not be otherwise practical with only a single light sensor module.

[1357] Each of the multiple light sensor modules **2134a,b** may receive light reflected from the tissue when illuminated by light emitted by one or more illumination sources **2136a,b** disposed at the distal end **2133** of the alternative elongated camera probe **2130**. In some non-limiting examples, the light emitted by all of the illumination sources **2136a,b** may be derived from the same light source (such as a laser). In other non-limiting examples, the illumination sources **2136a** surrounding a first light sensor module **2134a** may emit light at a first wavelength and the illumination sources **2136b** surrounding a second light sensor module **2134b** may emit light at a second wavelength. It may be further understood that each illumination source **2136a,b** may source illumination having any central wavelength including a central red illumination wavelength, a central green illumination wavelength, a central blue illumination wavelength, a central infrared illumination wavelength, a central ultraviolet illumination wavelength, or any other wavelength. In some examples, the one or more illumination

sources **2136a,b** may include a white light source, which may illuminate tissue with light having wavelengths that may span the range of optical white light from about 390 nm to about 700 nm.

[1358] In some additional aspects, the distal end **2133** of the alternative elongated camera probe **2130** may include one or more working channels **2138**. Such working channels **2138** may be in fluid communication with an aspiration port of a device to aspirate material from the surgical site, thereby permitting the removal of material that may potentially obscure the field of view of the light sensor modules **2134a,b**. Alternatively, such working channels **2138** may be in fluid communication with a fluid source port of a device to provide a fluid to the surgical site, to flush debris or material away from the surgical site. Such fluids may be used to clear material from the field of view of the light sensor modules **2134a,b**.

[1359] FIG. 174C depicts a perspective view of an aspect of a monolithic sensor **2160** having a plurality of pixel arrays for producing a three dimensional image in accordance with the teachings and principles of the disclosure. Such an implementation may be desirable for three dimensional image capture, wherein the two pixel arrays **2162** and **2164** may be offset during use. In another implementation, a first pixel array **2162** and a second pixel array **2164** may be dedicated to receiving a predetermined range of wave lengths of electromagnetic radiation, wherein the first pixel array **2162** is dedicated to a different range of wave length electromagnetic radiation than the second pixel array **2164**.

[1360] Additional disclosures regarding a dual sensor array may be found in U.S. Patent Application Publication No. 2014/0267655, titled SUPER RESOLUTION AND COLOR MOTION ARTIFACT CORRECTION IN A PULSED COLOR IMAGING SYSTEM, filed on Mar. 14, 2014, which issued on May 2, 2017 as U.S. Pat. No. 9,641,815, the contents thereof being incorporated by reference herein in its entirety and for all purposes.

[1361] In some aspects, a light sensor module may comprise a multi-pixel light sensor such as a CMOS array in addition to one or more additional optical elements such as a lens, a reticle, and a filter.

[1362] In some alternative aspects, the one or more light sensors may be located within the body **2021** of the hand unit **2020**. Light reflected from the tissue may be acquired at a light receiving surface of one or more optical fibers at the distal end of the elongated camera probe **2024**. The one or more optical fibers may conduct the light from the distal end of the elongated camera probe **2024** to the one or more light sensors, or to additional optical elements housed in the body of the hand unit **2020** or in the imaging control unit **2002**. The additional optical elements may include, without limitation, one or more dichroic mirrors, one or more reference mirrors, one or more moving mirrors, and one or more beam splitters and/or combiners, and one or more optical shutters. In such alternative aspects, the light sensor module may include any one or more of a lens, a reticle and a filter, disposed at the distal end of the elongated camera probe **2024**.

[1363] Images obtained from each of the multiple light sensors for example **2134a,b** may be combined or processed in several different manners, either in combination or separately, and then displayed in a manner to allow a surgeon to visualize different aspects of the surgical site.

[1364] In one non-limiting example, each light sensor may have an independent field of view. In some additional examples, the field of view of a first light sensor may partially or completely overlap the field of view of a second light sensor.

[1365] As disclosed above, an imaging system may include a hand unit 2020 having an elongated camera probe 2024 with one or more light sensor modules 2124, 2134a,b disposed at its distal end 2123, 2133. As an example, the elongated camera probe 2024 may have two light sensor modules 2134a,b, although it may be recognized that there may be three, four, five, or more light sensor modules at the distal end of the elongated camera probe 2024. Although FIGS. 175 and 176A-D depict examples of the distal end of an elongated camera probe having two light sensor modules, it may be recognized that the description of the operation of the light sensor modules is not limited to solely two light sensor modules. As depicted in FIGS. 175, and 46A-D, the light sensor modules may include an image sensor, such as a CCD or CMOS sensor that may be composed of an array of light sensing elements (pixels). The light sensor modules may also include additional optical elements, such as lenses. Each lens may be adapted to provide a field of view for the light sensor of the respective light sensor module.

[1366] FIG. 175 depicts a generalized view of a distal end 2143 of an elongated camera probe having multiple light sensor modules 2144a,b. Each light sensor module 2144a,b may be composed of a CCD or CMOS sensor and one or more optical elements such as filters, lenses, shutters, and similar. In some aspects, the components of the light sensor modules 2144a,b may be fixed within the elongated camera probe. In some other aspects, one or more of the components of the light sensor modules 2144a,b may be adjustable. For example, the CCD or CMOS sensor of a light sensor module 2144a,b may be mounted on a movable mount to permit automated adjustment of the center 2145a,b of a field of view 2147a,b of the CCD or CMOS sensor. In some other aspects, the CCD or CMOS sensor may be fixed, but a lens in each light sensor modules 2144a,b may be adjustable to change the focus. In some aspects, the light sensor modules 2144a,b may include adjustable irises to permit changes in the visual aperture of the sensor modules 2144a,b.

[1367] As depicted in FIG. 175, each of the sensor modules 2144a,b may have a field of view 2147a,b having an acceptance angle. As depicted in FIG. 175, the acceptance angle for each sensor modules 2144a,b may have an acceptance angle of greater than 90°. In some examples, the acceptance angle may be about 100°. In some examples, the acceptance angle may be about 120°. In some examples, if the sensor modules 2144a,b have an acceptance angle of greater than 90° (for example, 100°), the fields of view 2147a and 2147b may form an overlap region 2150a,b. In some aspects, an optical field of view having an acceptance angle of 100° or greater may be called a “fish-eyed” field of view. A visualization system control system associated with such an elongated camera probe may include computer readable instructions that may permit the display of the overlap region 2150a,b in such a manner so that the extreme curvature of the overlapping fish-eyed fields of view is corrected, and a sharpened and flattened image may be displayed. In FIG. 175, the overlap region 2150a may represent a region wherein the overlapping fields of view 2147a,b of the sensor modules 2144a,b have their respective centers 2145a,b directed in a forward direction. However, if

any one or more components of the sensor modules 2144a,b is adjustable, it may be recognized that the overlap region 2150b may be directed to any attainable angle within the fields of view 2147a,b of the sensor modules 2144a,b.

[1368] FIGS. 176A-D depict a variety of examples of an elongated light probe having two light sensor modules 2144a,b with a variety of fields of view. The elongated light probe may be directed to visualize a surface 2152 of a surgical site.

[1369] In FIG. 176A, the first light sensor module 2144a has a first sensor field of view 2147a of a tissue surface 2154a, and the second light sensor module 2144b has a second sensor field of view 2147b of a tissue surface 2154b. As depicted in FIG. 176A, the first field of view 2147a and the second field of view 2147b have approximately the same angle of view. Additionally, the first sensor field of view 2147a is adjacent to but does not overlap the second sensor field of view 2147b. The image received by the first light sensor module 2144a may be displayed separately from the image received by the second light sensor module 2144b, or the images may be combined to form a single image. In some non-limiting examples, the angle of view of a lens associated with the first light sensor module 2144a and the angle of view of a lens associated with the second light sensor module 2144b may be somewhat narrow, and image distortion may not be great at the periphery of their respective images. Therefore, the images may be easily combined edge to edge.

[1370] As depicted in FIG. 176B, the first field of view 2147a and the second field of view 2147b have approximately the same angular field of view, and the first sensor field of view 2147a overlaps completely the second sensor field of view 2147b. This may result in a first sensor field of view 2147a of a tissue surface 2154a being identical to the view of a tissue surface 2154b as obtained by the second light sensor module 2144b from the second sensor field 2147b of view. This configuration may be useful for applications in which the image from the first light sensor module 2144a may be processed differently than the image from the second light sensor module 2144b. The information in the first image may complement the information in the second image and refer to the same portion of tissue.

[1371] As depicted in FIG. 176C, the first field of view 2147a and the second field of view 2147b have approximately the same angular field of view, and the first sensor field of view 2147a partially overlaps the second sensor field of view 2147b. In some non-limiting examples, a lens associated with the first light sensor module 2144a and a lens associated with the second light sensor module 2144b may be wide angle lenses. These lenses may permit the visualization of a wider field of view than that depicted in FIG. 176A. Wide angle lenses are known to have significant optical distortion at their periphery. Appropriate image processing of the images obtained by the first light sensor module 2144a and the second light sensor module 2144b may permit the formation of a combined image in which the central portion of the combined image is corrected for any distortion induced by either the first lens or the second lens. It may be understood that a portion of the first sensor field of view 2147a of a tissue surface 2154a may thus have some distortion due to the wide angle nature of a lens associated with the first light sensor module 2144a and a portion of the second sensor field of view 2147b of a tissue surface 2154b may thus have some distortion due to the wide angle nature



of a lens associated with the second light sensor module **2144b**. However, a portion of the tissue viewed in the overlap region **2150'** of the two light sensor modules **2144a,b** may be corrected for any distortion induced by either of the light sensor modules **2144a,b**. The configuration depicted in FIG. **176C** may be useful for applications in which it is desired to have a wide field of view of the tissue around a portion of a surgical instrument during a surgical procedure. In some examples, lenses associated with each light sensor module **2144a,b** may be independently controllable, thereby controlling the location of the overlap region **2150'** of view within the combined image.

[**1372**] As depicted in FIG. **176D**, the first light sensor module **2144a** may have a first angular field of view **2147a** that is wider than the second angular field of view **2147b** of the second light sensor module **2144b**. In some non-limiting examples, the second sensor field of view **2147b** may be totally disposed within the first sensor field of view **2147a**. In alternative examples, the second sensor field of view may lie outside of or tangent to the wide angle field of view **2147a** of the first sensor **2144a**. A display system that may use the configuration depicted in FIG. **176D** may display a wide angle portion of tissue **2154a** imaged by the first sensor module **2144a** along with a magnified second portion of tissue **2154b** imaged by the second sensor module **2144b** and located in an overlap region **2150"** of the first field of view **2147a** and the second field of view **2147b**. This configuration may be useful to present a surgeon with a close-up image of tissue proximate to a surgical instrument (for example, imbedded in the second portion of tissue **2154b**) and a wide-field image of the tissue surrounding the immediate vicinity of the medical instrument (for example, the proximal first portion of tissue **2154a**). In some non-limiting examples, the image presented by the narrower second field of view **2147b** of the second light sensor module **2144b** may be a surface image of the surgical site. In some additional examples, the image presented in the first wide field view **2147a** of the first light sensor module **2144a** may include a display based on a hyperspectral analysis of the tissue visualized in the wide field view.

[**1373**] FIGS. **177A-C** illustrate an example of the use of an imaging system incorporating the features disclosed in FIG. **176D**. FIG. **177A** illustrates schematically a proximal view **2170** at the distal end of the elongated camera probe depicting the light sensor arrays **2172a,b** of the two light sensor modules **2174a,b**. A first light sensor module **2174a** may include a wide angle lens, and the second light sensor module **2174b** may include a narrow angle lens. In some aspects, the second light sensor module **2174b** may have a narrow aperture lens. In other aspects, the second light sensor module **2174b** may have a magnifying lens. The tissue may be illuminated by the illumination sources disposed at the distal end of the elongated camera probe. The light sensor arrays **2172'** (either light sensor array **2172a** or **2172b**, or both **2172a** and **2172b**) may receive the light reflected from the tissue upon illumination. The tissue may be illuminated by light from a red laser source, a green laser source, a blue laser source, an infrared laser source, and/or an ultraviolet laser source. In some aspects, the light sensor arrays **2172'** may sequentially receive the red laser light **2175a**, green laser light **2175b**, blue laser light **2175c**, infrared laser light **2175d**, and the ultra-violet laser light **2175e**. The tissue may be illuminated by any combination of such laser sources simultaneously, as depicted in FIGS.

**153E** and **153F**. Alternatively, the illuminating light may be cycled among any combination of such laser sources, as depicted for example in FIGS. **153D**, and FIGS. **173A** and **173B**.

[**1374**] FIG. **177B** schematically depicts a portion of lung tissue **2180** which may contain a tumor **2182**. The tumor **2182** may be in communication with blood vessels including one or more veins **2184** and/or arteries **2186**. In some surgical procedures, the blood vessels (veins **2184** and arteries **2186**) associated with the tumor **2182** may require resection and/or cauterization prior to the removal of the tumor.

[**1375**] FIG. **177C** illustrates the use of a dual imaging system as disclosed above with respect to FIG. **177A**. The first light sensor module **2174a** may acquire a wide angle image of the tissue surrounding a blood vessel **2187** to be severed with a surgical knife **2190**. The wide angle image may permit the surgeon to verify the blood vessel to be severed **2187**. In addition, the second light sensor module **2174b** may acquire a narrow angle image of the specific blood vessel **2187** to be manipulated. The narrow angle image may show the surgeon the progress of the manipulation of the blood vessel **2187**. In this manner, the surgeon is presented with the image of the vascular tissue to be manipulated as well as its environs to assure that the correct blood vessel is being manipulated.

[**1376**] FIGS. **178A** and **178B** depict another example of the use of a dual imaging system. FIG. **178A** depicts a primary surgical display providing an image of a section of a surgical site. The primary surgical display may depict a wide view image **2800** of a section of intestine **2802** along with its vasculature **2804**. The wide view image **2800** may include a portion of the surgical field **2809** that may be separately displayed as a magnified view **2810** in a secondary surgical display (FIG. **178B**). As disclosed above with respect to surgery to remove a tumor from a lung (FIGS. **177A-C**), it may be necessary to dissect blood vessels supplying a tumor **2806** before removing the cancerous tissue. The vasculature **2804** supplying the intestines **2802** is complex and highly ramified. It may necessary to determine which blood vessels supply the tumor **2806** and to identify blood vessels supplying blood to healthy intestinal tissue. The wide view image **2800** permits a surgeon to determine which blood vessel may supply the tumor **2806**. The surgeon may then test a blood vessel using a clamping device **2812** to determine if the blood vessel supplies the tumor **2806** or not.

[**1377**] FIG. **178B** depicts a secondary surgical display that may only display a narrow magnified view image **2810** of one portion of the surgical field **2809**. The narrow magnified view image **2810** may present a close-up view of the vascular tree **2814** so that the surgeon can focus on dissecting only the blood vessel of interest **2815**. For resecting the blood vessel of interest **2815**, a surgeon may use a smart RF cauterization device **2816**. It may be understood that any image obtained by the visualization system may include not only images of the tissue in the surgical site but also images of the surgical instruments inserted therein. In some aspects, such a surgical display (either the primary display in FIG. **178A** or the secondary display in FIG. **178B**) may also include indicia **2817** related to functions or settings of any surgical device used during the surgical procedure. For example, the indicia **2817** may include a power setting of the smart RF cauterization device **2816**. In some aspects, such smart medical

devices may transmit data related to their operating parameters to the visualization system to incorporate in display data to be transmitted to one or more display devices.

[1378] FIGS. 179A-C illustrate examples of a sequence of surgical steps for the removal of an intestinal/colon tumor and which may benefit from the use of multi-image analysis at the surgical site. FIG. 179A depicts a portion of the surgical site, including the intestines 2932 and the ramified vasculature 2934 supplying blood and nutrients to the intestines 2932. The intestines 2932 may have a tumor 2936 surrounded by a tumor margin 2937. A first light sensor module of a visualization system may have a wide field of view 2930, and it may provide imaging data of the wide field of view 2930 to a display system. A second light sensor module of the visualization system may have a narrow or standard field of view 2940, and it may provide imaging data of the narrow field of view 2940 to the display system. In some aspects, the wide field image and the narrow field image may be displayed by the same display device. In another aspect, the wide field image and the narrow field image may be displayed by separate display devices.

[1379] During the surgical procedure, it may be important to remove not just the tumor 2936 but the margin 2937 surrounding it to assure complete removal of the tumor. A wide angle field of view 2930 may be used to image both the vasculature 2934 as well as the section of the intestines 2932 surrounding the tumor 2936 and the margin 2637. As noted above, the vasculature feeding the tumor 2936 and the margin 2637 should be removed, but the vasculature feeding the surrounding intestinal tissue must be preserved to provide oxygen and nutrients to the surrounding tissue. Transection of the vasculature feeding the surrounding colon tissue will remove oxygen and nutrients from the tissue, leading to necrosis. In some examples, laser Doppler imaging of the tissue visualized in the wide angle field 2630 may be analyzed to provide a speckle contrast analysis 2933, indicating the blood flow within the intestinal tissue.

[1380] FIG. 179B illustrates a step during the surgical procedure. The surgeon may be uncertain which part of the vascular tree supplies blood to the tumor 2936. The surgeon may test a blood vessel 2944 to determine if it feeds the tumor 2936 or the healthy tissue. The surgeon may clamp a blood vessel 2944 with a clamping device 2812 and determine the section of the intestinal tissue 2943 that is no longer perfused by means of the speckle contrast analysis. The narrow field of view 2940 displayed on an imaging device may assist the surgeon in the close-up and detailed work required to visualize the single blood vessel 2944 to be tested. When the suspected blood vessel 2944 is clamped, a portion of the intestinal tissue 2943 is determined to lack perfusion based on the Doppler imaging speckle contrast analysis. As depicted in FIG. 159B, the suspected blood vessel 2944 does not supply blood to the tumor 2935 or the tumor margin 2937, and therefore is recognized as a blood vessel to be spared during the surgical procedure.

[1381] FIG. 179C depicts a following stage of the surgical procedure. In stage, a supply blood vessel 2984 has been identified to supply blood to the margin 2937 of the tumor. When this supply blood vessel 2984 has been severed, blood is no longer supplied to a section of the intestine 2987 that may include at least a portion of the margin 2937 of the tumor 2936. In some aspects, the lack of perfusion to the section 2987 of the intestines may be determined by means of a speckle contrast analysis based on a Doppler analysis of

blood flow into the intestines. The non-perfused section 2987 of the intestines may then be isolated by a seal 2985 applied to the intestine. In this manner, only those blood vessels perfusing the tissue indicated for surgical removal may be identified and sealed, thereby sparing healthy tissue from unintended surgical consequences.

[1382] In some additional aspects, a surgical visualization system may permit imaging analysis of the surgical site.

[1383] In some aspects, the surgical site may be inspected for the effectiveness of surgical manipulation of a tissue.

[1384] Non-limiting examples of such inspection may include the inspection of surgical staples or welds used to seal tissue at a surgical site. Cone beam coherent tomography using one or more illumination sources may be used for such methods.

[1385] In some additional aspects, an image of a surgical site may have landmarks denoted in the image. In some examples, the landmarks may be determined through image analysis techniques. In some alternative examples, the landmarks may be denoted through a manual intervention of the image by the surgeon.

[1386] In some additional aspects, non-smart ready visualizations methods may be imported for used in Hub image fusion techniques.

[1387] In additional aspects, instruments that are not integrated in the Hub system may be identified and tracked during their use within the surgical site. In this aspect, computational and/or storage components of the Hub or in any of its components (including, for example, in the cloud system) may include a database of images related to EES and competitive surgical instruments that are identifiable from one or more images acquired through any image acquisition system or through visual analytics of such alternative instruments. The imaging analysis of such devices may further permit identification of when an instrument is replaced with a different instrument to do the same or a similar job. The identification of the replacement of an instrument during a surgical procedure may provide information related to when an instrument is not doing the job or a failure of the device.

#### Cloud System Hardware and Functional Modules

[1388] Aspects of the present disclosure include a cloud-based medical analytics system that communicatively couples to multiple Hub systems, as described above, and multiple robotic surgical devices, described more below. The cloud-based medical analytics system is configured to receive data pertaining to a patient and/or medical procedure and provide various integrated processes that span multiple Hub systems and multiple robotic surgical devices. The cloud-based medical analytics system generally aggregates data and forms insights based on the aggregated data that may not otherwise be concluded without gathering the various disparate data sources that span the multiple Hub systems and robotic devices. Described below are various examples of different types of functions and structures present in the cloud-based medical analytics system that provide more detail toward these ends.

[1389] FIG. 180 is a block diagram of the computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure. In one aspect, the computer-implemented interactive surgical system is configured to monitor and analyze data related to the operation of various surgical systems that include surgical hubs,

surgical instruments, robotic devices and operating theaters or healthcare facilities. The computer-implemented interactive surgical system comprises a cloud-based analytics system. Although the cloud-based analytics system is described as a surgical system, it is not necessarily limited as such and could be a cloud-based medical system generally. As illustrated in FIG. 180, the cloud-based analytics system comprises a plurality of surgical instruments 7012 (may be the same or similar to instruments 112), a plurality of surgical hubs 7006 (may be the same or similar to hubs 106), and a surgical data network 7001 (may be the same or similar to network 201) to couple the surgical hubs 7006 to the cloud 7004 (may be the same or similar to cloud 204). Each of the plurality of surgical hubs 7006 is communicatively coupled to one or more surgical instruments 7012. The hubs 7006 are also communicatively coupled to the cloud 7004 of the computer-implemented interactive surgical system via the network 7001. The cloud 7004 is a remote centralized source of hardware and software for storing, manipulating, and communicating data generated based on the operation of various surgical systems. As shown in FIG. 180, access to the cloud 7004 is achieved via the network 7001, which may be the Internet or some other suitable computer network. Surgical hubs 7006 that are coupled to the cloud 7004 can be considered the client side of the cloud computing system (i.e., cloud-based analytics system). Surgical instruments 7012 are paired with the surgical hubs 7006 for control and implementation of various surgical procedures or operations as described herein.

[1390] In addition, surgical instruments 7012 may comprise transceivers for data transmission to and from their corresponding surgical hubs 7006 (which may also comprise transceivers). Combinations of surgical instruments 7012 and corresponding hubs 7006 may indicate particular locations, such as operating theaters in healthcare facilities (e.g., hospitals), for providing medical operations. For example, the memory of a surgical hub 7006 may store location data. As shown in FIG. 180, the cloud 7004 comprises central servers 7013 (may be same or similar to remote server 7013), hub application servers 7002, data analytics modules 7034, and an input/output (“I/O”) interface 7006. The central servers 7013 of the cloud 7004 collectively administer the cloud computing system, which includes monitoring requests by client surgical hubs 7006 and managing the processing capacity of the cloud 7004 for executing the requests. Each of the central servers 7013 comprises one or more processors 7008 coupled to suitable memory devices 7010 which can include volatile memory such as random-access memory (RAM) and non-volatile memory such as magnetic storage devices. The memory devices 7010 may comprise machine executable instructions that when executed cause the processors 7008 to execute the data analytics modules 7034 for the cloud-based data analysis, operations, recommendations and other operations described below. Moreover, the processors 7008 can execute the data analytics modules 7034 independently or in conjunction with hub applications independently executed by the hubs 7006. The central servers 7013 also comprise aggregated medical data databases 2212, which can reside in the memory 2210.

[1391] Based on connections to various surgical hubs 7006 via the network 7001, the cloud 7004 can aggregate data from specific data generated by various surgical instruments 7012 and their corresponding hubs 7006. Such aggregat-

ed data may be stored within the aggregated medical databases 7012 of the cloud 7004. In particular, the cloud 7004 may advantageously perform data analysis and operations on the aggregated data to yield insights and/or perform functions that individual hubs 7006 could not achieve on their own. To this end, as shown in FIG. 180, the cloud 7004 and the surgical hubs 7006 are communicatively coupled to transmit and receive information. The I/O interface 7006 is connected to the plurality of surgical hubs 7006 via the network 7001. In this way, the I/O interface 7006 can be configured to transfer information between the surgical hubs 7006 and the aggregated medical data databases 7011. Accordingly, the I/O interface 7006 may facilitate read/write operations of the cloud-based analytics system. Such read/write operations may be executed in response to requests from hubs 7006. These requests could be transmitted to the hubs 7006 through the hub applications. The I/O interface 7006 may include one or more high speed data ports, which may include universal serial bus (USB) ports, IEEE 1394 ports, as well as Wi-Fi and Bluetooth I/O interfaces for connecting the cloud 7004 to hubs 7006. The hub application servers 7002 of the cloud 7004 are configured to host and supply shared capabilities to software applications (e.g., hub applications) executed by surgical hubs 7006. For example, the hub application servers 7002 may manage requests made by the hub applications through the hubs 7006, control access to the aggregated medical data databases 7011, and perform load balancing. The data analytics modules 7034 are described in further detail with reference to FIG. 181.

[1392] The particular cloud computing system configuration described in the present disclosure is specifically designed to address various issues arising in the context of medical operations and procedures performed using medical devices, such as the surgical instruments 7012, 112. In particular, the surgical instruments 7012 may be digital surgical devices configured to interact with the cloud 7004 for implementing techniques to improve the performance of surgical operations. Various surgical instruments 7012 and/or surgical hubs 7006 may comprise touch controlled user interfaces such that clinicians may control aspects of interaction between the surgical instruments 7012 and the cloud 7004. Other suitable user interfaces for control such as auditory controlled user interfaces can also be used.

[1393] FIG. 181 is a block diagram which illustrates the functional architecture of the computer-implemented interactive surgical system, in accordance with at least one aspect of the present disclosure. The cloud-based analytics system includes a plurality of data analytics modules 7034 that may be executed by the processors 7008 of the cloud 7004 for providing data analytic solutions to problems specifically arising in the medical field. As shown in FIG. 181, the functions of the cloud-based data analytics modules 7034 may be assisted via hub applications 7014 hosted by the hub application servers 7002 that may be accessed on surgical hubs 7006. The cloud processors 7008 and hub applications 7014 may operate in conjunction to execute the data analytics modules 7034. Application program interfaces (APIs) 7016 define the set of protocols and routines corresponding to the hub applications 7014. Additionally, the APIs 7016 manage the storing and retrieval of data into and from the aggregated medical databases 7012 for the operations of the applications 7014. The caches 7018 also store data (e.g., temporarily) and are coupled to the APIs 7016 for more

efficient retrieval of data used by the applications 7014. The data analytics modules 7034 in FIG. 181 include modules for resource optimization 7020, data collection and aggregation 7022, authorization and security 7024, control program updating 7026, patient outcome analysis 7028, recommendations 7030, and data sorting and prioritization 7032. Other suitable data analytics modules could also be implemented by the cloud 7004, according to some aspects. In one aspect, the data analytics modules are used for specific recommendations based on analyzing trends, outcomes, and other data.

[1394] For example, the data collection and aggregation module 7022 could be used to generate self-describing data (e.g., metadata) including identification of notable features or configuration (e.g., trends), management of redundant data sets, and storage of the data in paired data sets which can be grouped by surgery but not necessarily keyed to actual surgical dates and surgeons. In particular, pair data sets generated from operations of surgical instruments 7012 can comprise applying a binary classification, e.g., a bleeding or a non-bleeding event. More generally, the binary classification may be characterized as either a desirable event (e.g., a successful surgical procedure) or an undesirable event (e.g., a misfired or misused surgical instrument 7012). The aggregated self-describing data may correspond to individual data received from various groups or sub-groups of surgical hubs 7006. Accordingly, the data collection and aggregation module 7022 can generate aggregated metadata or other organized data based on raw data received from the surgical hubs 7006. To this end, the processors 7008 can be operationally coupled to the hub applications 7014 and aggregated medical data databases 7011 for executing the data analytics modules 7034. The data collection and aggregation module 7022 may store the aggregated organized data into the aggregated medical data databases 2212.

[1395] The resource optimization module 7020 can be configured to analyze this aggregated data to determine an optimal usage of resources for a particular or group of healthcare facilities. For example, the resource optimization module 7020 may determine an optimal order point of surgical stapling instruments 7012 for a group of healthcare facilities based on corresponding predicted demand of such instruments 7012. The resource optimization module 7020 might also assess the resource usage or other operational configurations of various healthcare facilities to determine whether resource usage could be improved. Similarly, the recommendations module 7030 can be configured to analyze aggregated organized data from the data collection and aggregation module 7022 to provide recommendations. For example, the recommendations module 7030 could recommend to healthcare facilities (e.g., medical service providers such as hospitals) that a particular surgical instrument 7012 should be upgraded to an improved version based on a higher than expected error rate, for example. Additionally, the recommendations module 7030 and/or resource optimization module 7020 could recommend better supply chain parameters such as product reorder points and provide suggestions of different surgical instrument 7012, uses thereof, or procedure steps to improve surgical outcomes. The healthcare facilities can receive such recommendations via corresponding surgical hubs 7006. More specific recommendations regarding parameters or configurations of various surgical instruments 7012 can also be provided. Hubs

7006 and/or surgical instruments 7012 each could also have display screens that display data or recommendations provided by the cloud 7004.

[1396] The patient outcome analysis module 7028 can analyze surgical outcomes associated with currently used operational parameters of surgical instruments 7012. The patient outcome analysis module 7028 may also analyze and assess other potential operational parameters. In this connection, the recommendations module 7030 could recommend using these other potential operational parameters based on yielding better surgical outcomes, such as better sealing or less bleeding. For example, the recommendations module 7030 could transmit recommendations to a surgical 7006 regarding when to use a particular cartridge for a corresponding stapling surgical instrument 7012. Thus, the cloud-based analytics system, while controlling for common variables, may be configured to analyze the large collection of raw data and to provide centralized recommendations over multiple healthcare facilities (advantageously determined based on aggregated data). For example, the cloud-based analytics system could analyze, evaluate, and/or aggregate data based on type of medical practice, type of patient, number of patients, geographic similarity between medical providers, which medical providers/facilities use similar types of instruments, etc., in a way that no single healthcare facility alone would be able to analyze independently. The control program updating module 7026 could be configured to implement various surgical instrument 7012 recommendations when corresponding control programs are updated. For example, the patient outcome analysis module 7028 could identify correlations linking specific control parameters with successful (or unsuccessful) results. Such correlations may be addressed when updated control programs are transmitted to surgical instruments 7012 via the control program updating module 7026. Updates to instruments 7012 that are transmitted via a corresponding hub 7006 may incorporate aggregated performance data that was gathered and analyzed by the data collection and aggregation module 7022 of the cloud 7004. Additionally, the patient outcome analysis module 7028 and recommendations module 7030 could identify improved methods of using instruments 7012 based on aggregated performance data.

[1397] The cloud-based analytics system may include security features implemented by the cloud 7004. These security features may be managed by the authorization and security module 7024. Each surgical hub 7006 can have associated unique credentials such as username, password, and other suitable security credentials. These credentials could be stored in the memory 7010 and be associated with a permitted cloud access level. For example, based on providing accurate credentials, a surgical hub 7006 may be granted access to communicate with the cloud to a predetermined extent (e.g., may only engage in transmitting or receiving certain defined types of information). To this end, the aggregated medical data databases 7011 of the cloud 7004 may comprise a database of authorized credentials for verifying the accuracy of provided credentials. Different credentials may be associated with varying levels of permission for interaction with the cloud 7004, such as a predetermined access level for receiving the data analytics generated by the cloud 7004. Furthermore, for security purposes, the cloud could maintain a database of hubs 7006, instruments 7012, and other devices that may comprise a

“black list” of prohibited devices. In particular, a surgical hubs **7006** listed on the black list may not be permitted to interact with the cloud, while surgical instruments **7012** listed on the black list may not have functional access to a corresponding hub **7006** and/or may be prevented from fully functioning when paired to its corresponding hub **7006**. Additionally or alternatively, the cloud **7004** may flag instruments **7012** based on incompatibility or other specified criteria. In this manner, counterfeit medical devices and improper reuse of such devices throughout the cloud-based analytics system can be identified and addressed.

**[1398]** The surgical instruments **7012** may use wireless transceivers to transmit wireless signals that may represent, for example, authorization credentials for access to corresponding hubs **7006** and the cloud **7004**. Wired transceivers may also be used to transmit signals. Such authorization credentials can be stored in the respective memory devices of the surgical instruments **7012**. The authorization and security module **7024** can determine whether the authorization credentials are accurate or counterfeit. The authorization and security module **7024** may also dynamically generate authorization credentials for enhanced security. The credentials could also be encrypted, such as by using hash based encryption. Upon transmitting proper authorization, the surgical instruments **7012** may transmit a signal to the corresponding hubs **7006** and ultimately the cloud **7004** to indicate that the instruments **7012** are ready to obtain and transmit medical data. In response, the cloud **7004** may transition into a state enabled for receiving medical data for storage into the aggregated medical data databases **7011**. This data transmission readiness could be indicated by a light indicator on the instruments **7012**, for example. The cloud **7004** can also transmit signals to surgical instruments **7012** for updating their associated control programs. The cloud **7004** can transmit signals that are directed to a particular class of surgical instruments **7012** (e.g., electro-surgical instruments) so that software updates to control programs are only transmitted to the appropriate surgical instruments **7012**. Moreover, the cloud **7004** could be used to implement system wide solutions to address local or global problems based on selective data transmission and authorization credentials. For example, if a group of surgical instruments **7012** are identified as having a common manufacturing defect, the cloud **7004** may change the authorization credentials corresponding to this group to implement an operational lockout of the group.

**[1399]** The cloud-based analytics system may allow for monitoring multiple healthcare facilities (e.g., medical facilities like hospitals) to determine improved practices and recommend changes (via the recommendations module **2030**, for example) accordingly. Thus, the processors **7008** of the cloud **7004** can analyze data associated with an individual healthcare facility to identify the facility and aggregate the data with other data associated with other healthcare facilities in a group. Groups could be defined based on similar operating practices or geographical location, for example. In this way, the cloud **7004** may provide healthcare facility group wide analysis and recommendations. The cloud-based analytics system could also be used for enhanced situational awareness. For example, the processors **7008** may predictively model the effects of recommendations on the cost and effectiveness for a particular facility (relative to overall operations and/or various medical procedures). The cost and effectiveness associated with that

particular facility can also be compared to a corresponding local region of other facilities or any other comparable facilities.

**[1400]** The data sorting and prioritization module **7032** may prioritize and sort data based on criticality (e.g., the severity of a medical event associated with the data, unexpectedness, suspiciousness). This sorting and prioritization may be used in conjunction with the functions of the other data analytics modules **7034** described above to improve the cloud-based analytics and operations described herein. For example, the data sorting and prioritization module **7032** can assign a priority to the data analysis performed by the data collection and aggregation module **7022** and patient outcome analysis modules **7028**. Different prioritization levels can result in particular responses from the cloud **7004** (corresponding to a level of urgency) such as escalation from an expedited response, special processing, exclusion from the aggregated medical data databases **7011**, or other suitable responses. Moreover, if necessary, the cloud **7004** can transmit a request (e.g., a push message) through the hub application servers for additional data from corresponding surgical instruments **7012**. The push message can result in a notification displayed on the corresponding hubs **7006** for requesting supporting or additional data. This push message may be required in situations in which the cloud detects a significant irregularity or outlier and the cloud cannot determine the cause of the irregularity. The central servers **7013** may be programmed to trigger this push message in certain significant circumstances, such as when data is determined to be different from an expected value beyond a predetermined threshold or when it appears security has been compromised, for example.

**[1401]** Additional example details for the various functions described are provided in the ensuing descriptions below. Each of the various descriptions may utilize the cloud architecture as described in FIGS. **180** and **181** as one example of hardware and software implementation.

#### Usage, Resource, and Efficiency Modeling for Medical Facility

**[1402]** Aspects of the present disclosure are presented for a cloud-based analytics system, communicatively coupled to a plurality of hubs and smart medical instruments, and configured to provide customized recommendations to localized medical care facilities regarding usage of medical supplies and other resources to improve efficiency and optimize resource allocation. A medical care facility, such as a hospital or medical clinic, may develop a set of practices for procuring, using, and disposing of various medical supplies that are often derived from routines and traditions maintained over time. The behaviors of a medical facility typically are risk-averse, and generally would be hesitant to adopt new and better practices unless and until convincingly shown of a better practice. Similarly, even if a better usage or efficiency model has been developed in a nearby facility, it is difficult for a local facility to adopt the improved practice because 1) each facility may be more natively resistant to change from the outside and 2) there are many unknowns for how or why the improved practice works in the nearby facility in relation to what the local facility does instead. Furthermore, even if a medical facility desired to improve its practices, it may be unable to do so optimally because it lacks enough knowledge from other similarly

situated facilities, either in its region, according to a similar size, and/or according to similar practices or patients, and the like.

**[1403]** To help facilitate the dissemination of improved practices across multiple medical facilities, it would be desirable if a common source could have knowledge of the contexts from multiple medical facilities and be able to determine what changes should be made for any particular medical facility, based on the knowledge of the practices of any or all of the multiple facilities.

**[1404]** In some aspects, a cloud-based system communicatively coupled to knowledge centers in a medical facility, such as one or more medical hubs, may be configured to aggregate medical resource usage data from multiple medical facilities. The cloud-based system may then correlate the medical resource usage data with outcomes from those facilities, and may be able to derive various patterns within the data. For example, in some aspects, the cloud-based system may find which hospitals generate the least amount of waste per unit cost, based on an aggregation of all waste and procurement data obtained from medical facilities in a wide geographic region (e.g., all surgery centers in Japan). The cloud-based system may be configured to identify which medical facility produced the least amount of waste per unit cost, and then may analyze what practices differentiate that medical facility. If a trend is found, the cloud-based system may disseminate this information to all of the similarly situated medical facilities to improve their practices. This analysis may help improve inventory management, throughput efficiency, or overall efficiency of a medical facility. The improved inventory management may help surgical devices and other medical resources be utilized at their peak performance levels for longer periods of time, compared to if resources were badly managed, and therefore medical devices may be continuously used while they are older and more worn down.

**[1405]** In general, the cloud-based system may be configured to aggregate data from multiple medical facilities, something that no single facility alone would be able to accomplish on its own. Furthermore, the cloud-based system may be configured to analyze the large collection of data, controlling for common variables, such as type of practice, type of patient, number of patients, geographic similarity, which facilities use similar types of instruments, etc., that no single facility alone would be able to analyze on its own.

**[1406]** In this way, the cloud-based system of the present disclosure may be able to find more accurate causalities that lead to best practices at a particular facility, which can then be disseminated to all of the other facilities. Furthermore, the cloud-based system may be able to provide the data from all of the disparate sources that no single facility may be able to do on its own.

**[1407]** Referring to FIG. 182, shown is an example illustration of a tabulation of various resources correlated to particular types of surgical categories. There are two bars for each category, with the dashed line bars **7102**, **7106**, and **7110** representing unused and/or scrap resources, and the solid line bars **7104**, **7108**, and **7112** showing a totality of resourced in use for that category. In this example, bars **7104**, **7108**, and **7112** show a total amount of endocutter cartridges, sponges, saline, fibrin sealants, sutures, and stapler buttresses, for thoracic, colorectal, and bariatric procedures, respectively, compared to the lower amounts **7102**,

**7106**, and **7110** representing an amount of unused resources for the thoracic, colorectal, and bariatric procedures, respectively.

**[1408]** The cloud system may be configured to identify wasted product that was gathered and not used or gathered and used in a manner that was not beneficial to the patient or the surgery. To do this, the cloud system may record in memory all records of inventory intake and disposal. During each intake, the inventory may be scanned and entered, and the bar codes of each inventory item may identify what type of product it is, as an example. In some aspects, smart disposal bins may be utilized to automatically tabulate when a product is being disposed of. These may be connected to the cloud system ultimately, either through one or more surgical hubs or through a separate inventory management system throughout the entire facility. Each facility may be tracked by its location, for example through a set GPS coordinate, inputted address or the like. This data may be organized in memory using one or more databases with various meta data associated with it, such as date and time of use, location of origin, type of procedure used for if applicable, cost per item, expiration date if applicable, and so on.

**[1409]** In addition, the cloud system may be configured to identify misfired or misused product and tracking of where the product was used, and may archive these results. For example, each surgical instrument communicatively coupled to a surgical hub may transmit a record of when the instrument was fired, such as to fire a staple or apply ultrasonic energy. Each record may be transmitted through the instrument and recorded at the cloud system ultimately. The action by the instrument may be tied with an outcome, either at that instant or with an overall outcome stating whether the procedure was successful or not. The action may be associated with a precise timestamp that places the action at an exact point during a surgery, where all of the actions of the surgery are also automatically recorded to the cloud, including start and end times of the surgery. This enables all of the human medical care workers to focus on their respective duties during surgery, rather than worry about an exact instance an action of a medical instrument occurred. The recordings of the medical instruments can be used to identify what products may be wasted during surgery, and the cloud system may be configured to also identify usage trends in this way.

**[1410]** In some aspects, the cloud system may be configured to perform trending analysis of the product tied to the overall length or amount of the product to identify short fires, or discarded product. For example, the cloud system may place the use of a product within a known period of when a surgical procedure is occurring, with a time stamp. The cloud system may then record an amount of resources utilized during that procedure, and may compare the materials used in that procedure with similarly situated procedures performed elsewhere. Out of this, several conclusions may be reached by the cloud system. For example, the cloud system may provide recommendations of a mix that provides smaller portions or an alternative usage that results in less wasted product. As another example, the cloud system may provide a suggestion or specified protocol change of specialized kits that would assemble the product in a manner more aligned to the detected institution usage. As yet another example, the cloud system may provide a suggestion or a change in protocol for alternative product mixes that would

be more aligned to the detected usage and therefore should result in less wasted product. As yet another example, the cloud system may provide a recommendation on how to adjust a medical procedure during surgery based on timings of actions occurring before or after an event that typically results in wasteful resources, such as misfirings or multiple firings, based on identifying a correlation or pattern that actions during surgery occurring within a certain time interval relative to a prior action tend to result in wasteful actions. These analyses may be derived in part using algorithms that attempt to optimize the available resources with the rates of their disposals, taking into account various factors such as misfirings, native practices of the surgeons or the facility at large, and so forth.

[1411] Still referring to FIG. 182, based on the tabulation of the used and unused product, the cloud system can also generate several other conclusions. For example, the cloud system may be configured to generate a correlation of unused product to cost overhead. The cloud system may also generate a calculation of expired product and how that impacts rates of change with inventory. It may also generate an indication of where in the supply chain the product is being unused and how it is being accounted for. It may also generate ways to reduce costs or inventory space by finding substitutes of some resources over others for the same procedure. This may be based on comparing similar practices at different medical facilities that use different resources to perform the same procedures.

[1412] In some aspects, the cloud system may be configured to analyze the inventory usage of any and all medical products and conduct procurement management for when to acquire new product. The cloud system may optimize the utilization of inventory space to determine how best to utilize what space is available, in light of rates of usage for certain products compared to others. It may often be the case that inventory is not closely monitored in terms of how long a product remains in storage. If certain products are utilized at slower rates, but there is a large amount of it, it may be determined that the storage space is allocated poorly. Therefore, the cloud system may better apportion the storage space to reflect actual resource usage.

[1413] To improve in this area, in some aspects, the cloud system may for example, identify missing or insufficient product within an operating room (OR) for a specified procedure. The cloud system may then provide an alert or notification or transmit data to display that deficiency at the surgical hub in the OR. As another example, when a product is used in the OR, it may communicate its usage information to the cloud, such as activate a sensor or activation identification. The product may be registered with a scan or a power on switch. Analysis of this information for a given hospital coupled with its ordering information, may eventually inform the supply status and can enable ordering recommendations. This may occur automatically, once the cloud system registers that products are being used in the OR, or through other means.

[1414] In some aspects, device utilization within a procedure is monitored by the cloud system and compared for a given segment (e.g., individual surgeon, individual hospital, network of hospitals, region, etc.) against device utilization for similar procedures in other segments. Recommendations are presented to optimize utilization based on unit resource used or expenditure spent to supply such resource. In

general, the cloud system may focus on a comparison of product utilization between different institutions that it is connected with.

[1415] FIG. 183 provides an example illustration of how the data is analyzed by the cloud system to provide a comparison between multiple facilities to compare use of resources. In general, the cloud system 7200 may obtain usage data from all facilities, such as any of the types of data described with respect to FIG. 182, and may associate each datum with various other meta data, such as time, procedure, outcome of the procedure, cost, date of acquisition, and so forth. FIG. 183 shows an example set of data 7202 being uploaded to the cloud 7200, each circle in the set 7202 representing an outcome and one or more resources and contextual metadata that may be relevant to leading to the outcome. In addition, high performing outcomes 7204 and their associated resources and contextual metadata are also uploaded to the cloud 7200, though at the time of upload, it may not be known which data has very good outcomes or simply average (or below average) outcomes. The cloud system may identify which use of resources is associated with better results compared to an average or expected outcome. This may be based on determining which resources last longer, are not wasted as often, ultimately cost less per unit time or unit resource, as some examples. The cloud system may analyze the data to determine best outcomes based on any and all of these variables, or even one or more combinations of them. The trends identified may then be used to find a correlation or may prompt request of additional data associated with these data points. If a pattern is found, these recommendations may be alerted to a user to examine as possible ways to improve resource usage and efficiency.

[1416] The example graph 7206 provides a visual depiction of an example trend or pattern that the cloud may derive from examining the resource and outcome data, according to some aspects. In this example, the cloud system may have analyzed resource and outcome data of number of stapler firings and their relation to performance in surgery. The cloud system may have gathered the data from multiple medical facilities, and multiple surgeons within each facility, based on automatically recorded firing data during each surgery that is generated directly from the operation of the surgical staplers themselves. The performance outcomes may be based on post-op examinations and evaluations, and/or immediate outcomes during surgery, such as whether there is a bleeding event or a successful wound closure. Based on all of the data, trends may be determined, and here, it may be discovered that there is a small window of the number of firings that results in the best performance outcomes, at interval "a" as shown. The magnitude of this performance compared to the most common number of firings is shown as interval "b." Because the number of firings that results in the best outcomes may not be what is commonly practiced, it may not be readily easily to have discovered these outcomes without the aggregation and analytical abilities of the cloud system.

[1417] As another example: cartridge type, color, and adjunct usage that are monitored for sleeve gastrectomy procedures for individual surgeons within the same hospital may be obtained. The data may reveal an average procedure cost for one surgeon is higher for this surgeon when compared to others within the same hospital, yet short term patient outcomes remain the same. The hospital is then

informed and is encouraged to look into differences in device utilization, techniques, etc. in search of optimizing costs potentially through the elimination of adjuncts.

**[1418]** In some aspects, the cloud system may also identify specialty cases. For example, specific cost information provided within the hospital, including OR time, device utilization, and staff, may be identified. These aspects may be unique to a particular OR, or facility. The cloud system may be configured to suggest efficiencies in OR time usage (scheduling), device inventory, etc. across specialties (orthopedics, thoracic, colorectal, bariatric, etc.) for these specialty cases.

**[1419]** In some aspects, the cloud system may also be configured to compare cost-benefit of robotic surgery vs traditional methods, such as laparoscopic procedures for given procedure type. The cloud system may compare device costs, OR time, patient discharge times, efficacy of the procedure done by the robot vs performed by surgeons exclusively, and the like.

#### Linking of Local Usage Trends with the Resource Acquisition Behaviors of the Larger Data Set (Individualized Change)

**[1420]** According to some aspects of the cloud system, whereas the above disclosure focuses on a determination of efficiency (i.e., value) and optimizing based on that, here, this section centers around on identifying which local practices may be best disseminated to other similarly situated medical facilities.

**[1421]** A medical care facility, such as a hospital or medical clinic, may develop a set of practices for how to utilize medical devices for aiding medical procedures that are often derived from routines and traditions maintained over time. The behaviors of a medical facility typically are risk-averse, and generally would be hesitant to adopt new and better practices unless and until convincingly shown of a better practice. Similarly, even if a better practice for utilizing a device or for adjusting a procedure has been developed in a nearby facility, it is difficult for a local facility to adopt the improved practice because 1) each facility may be more natively resistant to change from the outside and 2) there are many unknowns for how or why the improved practice works in the nearby facility in relation to what the local facility does instead. Furthermore, even if a medical facility desired to improve its practices, it may be unable to do so optimally because it lacks enough knowledge from other similarly situated facilities, either in its region, according to a similar size, and/or according to similar practices or patients, and the like.

**[1422]** To help facilitate the dissemination of improved practices across multiple medical facilities, it would be desirable if a common source could have knowledge of the contexts from multiple medical facilities and be able to determine what changes should be made for any particular medical facility, based on the knowledge of the practices of any or all of the multiple facilities.

**[1423]** In some aspects, a cloud-based system communicatively coupled to knowledge centers in a medical facility, such as one or more medical hubs, may be configured to aggregate resource utilization data and patient outcomes from multiple medical facilities. The cloud-based system may then correlate the resource utilization data with the outcomes from those facilities, and may be able to derive various patterns within the data. For example, in some

aspects, the cloud-based system may find which hospitals produce better outcomes for a particular type of procedure, based on an aggregation of all the patient outcome data for that particular procedure collected in a wide geographic region (e.g., all surgery centers in Germany). The cloud-based system may be configured to identify which medical facility produced a better procedural outcome compared to the average across the geographic region, and then may analyze what differences in that procedure occur in that medical facility. If a trend is found and one or more differences are identified, the cloud-based system may disseminate this information to all of the similarly situated medical facilities to improve their practices.

**[1424]** In general, the cloud-based system may be configured to aggregate data from multiple medical facilities, something that no single facility alone would be able to accomplish on its own. Furthermore, the cloud-based system may be configured to analyze the large collection of data, controlling for common variables, such as type of practice, type of patient, number of patients, geographic similarity, which facilities use similar types of instruments, etc., that no single facility alone would be able to analyze on its own.

**[1425]** In this way, the cloud-based system of the present disclosure may be able to find more accurate causalities that give rise to best practices at a particular facility, which can then be disseminated to all of the other facilities. Furthermore, the cloud-based system may be able to provide the data from all of the disparate sources that no single facility may be able to do on its own.

**[1426]** The cloud system may be configured to generate conclusions about the efficacy of any local facility in a number of ways. For example, the cloud system may determine if a local treatment facility is using a product mixture or usage that differs from the larger community and their outcomes are superior. The cloud system may then correlate the differences and highlight them for use in other facilities, other surgical hub, or in clinical sales as some examples. In general, this information may be disseminated widely in a way that no single facility may have had access or knowledge of, including the facility that practiced this improve procedure.

**[1427]** As another example, the cloud system may determine if the local facility has equal to or inferior outcomes to the larger community. The cloud system may then correlate suggestions and provide that information back to the local facility as recommendations. The system may display data showing their performance in relation to others, and may also display suggestions on what that facility is doing compared to what everybody else is doing. Again, the local facility may not even know they have an inefficiency in that respect, nor may everybody else realize they are utilizing their resources more efficiently, and thus nobody would ever know to examine these issues without the cloud system having a bigger picture of all of the data.

**[1428]** These suggestions can come in various forms. For example, the cloud system may provide recommendations at the purchasing level that suggest improvements in cost for similar outcomes. As another example, the cloud system may provide recommendations at the OR level when the procedure is being planned and outfitted as the less desirable products are being pulled suggest other techniques and product mixes that would be in line with the broader community which is achieving higher outcomes. As yet another example, the cloud system may display outcomes



comparison needs to account for surgeon experience, possibly through a count of similar cases performed by that surgeon from cloud data. In some aspects, the learning curve of an individual may be reported against an aggregated larger dataset, as expectation of improved outcomes, or of surgeon performance relative to peers in obtaining a steady state outcome level.

[1429] FIG. 184 illustrates one example of how the cloud system 7300 may determine efficacy trends from an aggregated set of data 7302 across whole regions, according to some aspects. Here, for each circle of the set of data 7302, device utilization, cost, and procedure outcomes for a procedure is monitored and compared for a given segment (e.g., individual surgeon, individual hospital, network of hospitals, region, etc.) against device utilization, cost, and procedure outcomes for similar procedures in other segments. These data may possess metadata that associates it to a particular facility. In general, an outcome of a procedure may be linked to multiple types of data associated with it, such as what resources were used, what procedure was performed, who performed the procedure, where the procedure was performed, and so on. The data linked to the outcome may then be presented as a data pair. The data may be subdivided in various ways, such as between good and inferior outcomes, filtered by particular facilities, particular demographics, and so forth. A regional filter 7304 is visually depicted as an example. The data set 7302 contains both good outcomes and inferior outcomes, with the inferior outcomes being darkened for contrast.

[1430] FIG. 184 also shows examples of charts that have these distinctions made and may be derived from the aggregated data set 7302, using one or more data pairs. Chart 7306 shows a global analysis in one example, while a regionally segmented analysis is provided in the other chart 7308. Statistical analysis may be performed to determine whether the outcomes are statistically significant. In chart 7306, the cloud system may determine that no statistical difference was found between good outcomes and inferior outcomes based on rates of occurrence. In contrast, in chart 7308, the cloud system may determine that there is a statistically higher occurrence of inferior outcomes for a given region, when filtering for a particular region. Recommendations are presented to share outcomes vs. cost vs. device utilization and all combinations therein to help inform optimization of outcomes against procedure costs with device utilization potentially being a key contributor of differences, according to some aspects.

[1431] As another example, a cartridge type and color are monitored for lobectomy procedures for individual surgeons within the same hospital. The data reveals average cost for one surgeon is higher on average for this surgeon, yet average length of stay is less. The hospital is informed by the cloud system and is encouraged to look into differences in device utilization, techniques, etc. in search of improving patient outcomes.

[1432] In some aspects, the cloud system may also be configured to provide predictive modeling of changes to procedures, product mixes, and timing for a given localized population or for the general population as a whole. The predictive modeling may be used to assess impact on resource utilization, resource efficiency, and resource performance, as some examples.

[1433] FIG. 185 provides an example illustration of some types of analysis the cloud system may be configured to

perform to provide the predicting modeling, according to some aspects. The cloud system may combine its knowledge of the required steps and instruments for performing a procedure, and may compare the different avenues via various metrics, such as resources utilized, time, procedural cost, and the like. In this example of chart 7400, a thoracic lobectomy procedure is analyzed using two different types of methods to perform the same procedure. Option A describes a disposable ultrasonic instrument as the method for performing the procedure, while Option B shows a combination of different methods that in the aggregate perform the same procedure. The graphical illustration may help a surgeon or administrator see how the resources are utilized and their cost. Option B is broken down into multiple sections, including sterilization cost, reusable dissectors and additional time in the OR for performing the procedure. The cloud system may be configured to convert these somewhat abstract notions into a quantitative cost value based on combining its knowledge of time spent in the OR, staff salaries and resource costs per unit time in the OR, and resources utilized for sterilization and reusable dissectors and their associated costs. The cloud system may be configured to associate the various amounts of resources and costs with its knowledge of the required steps to perform the thoracic lobectomy procedure using the prescribed method in Option B.

[1434] As another example, chart 7404 in FIG. 185 shows a comparison between using an ultrasonic long dissector and a monopolar reusable dissector to perform various portions of a procedure. Chart 7404 shows a comparison in terms of time needed to perform each portion of the procedure for each instrument. The surgeon may then be able to select which instrument may be desired for a particular procedure. The breakout times may be automatically recorded empirically during live procedures, with the times for each portion of the overall procedure broken out due to the cloud system's knowledge of the expected sequence to perform the procedure. Demarcations between each portion may be set by a surgeon providing an input to manually denote when each change occurs. In other cases, the cloud system may utilize situational awareness to determine when a portion of the procedure has ended based on the way the devices are used and not used. The cloud system may aggregate a number of these procedures, performed across multiple surgeons and multiple facilities, and then compute an average time for each section, as an example.

[1435] As another example, chart 7402 in FIG. 185 shows an example graphical interface for comparing relative cost when utilizing the ultrasonic long dissector or a monopolar reusable dissector, according to some aspect. The value of each instrument per unit time is displayed for a particular procedure. The data used to generate these values may be similar to those obtained for charts 7400 and 7404, as some examples. The graphical display may allow for a succinct description of the key points of efficiency that would be most useful to make a determination. This analysis may help a surgeon see how valuable each instrument is for a given procedure.

[1436] In general, to perform the predictive modeling, the cloud system may combine its knowledge of the exact steps to perform a procedure, what instruments may be used to perform each step, and its aggregated data for how each instrument performs each particular step. A surgeon may not have the combination of such knowledge in order to provide

such an assessment alone. The predictive modeling therefore may be the result of continued monitoring and acquisition of data across multiple facilities, the likes of which would not be possible without the cloud system.

[1437] In some aspects, the cloud system may also derive the distilled information from multiple sources (e.g., HUB data collection sources, literature, etc.) to identify the optimal procedure technique. Various other examples for how predictive modeling may be utilized include:

- (1) sigmoidectomy: multi-quadrant surgery; which is the best order of operations, etc.;
- (2) RYGB: what is the ideal limb length, etc. based on the circumstances for this patient;
- (3) Lobectomy: how many and which lymph nodes should be removed; and
- (4) VSG: Bougie size and distance from pylorus.

[1438] In some aspects, when a suggestion is made to a surgeon, the surgeon is given the option to decline future suggestions like this, or to continue. In addition, through interface with the hub, the surgeon may inquire to the cloud system additional information to inform his or her decision. For example, the surgeon may want to isolate the times to a more localized set of data, such as the particular facility or a certain demographic that better caters to the patient undergoing the surgery. The data may change, for example, if the patient is a child or the patient is a woman.

#### Device Setup Modifications Based on Surgeon, Regional, Hospital, or Patient Parameters (Preoperatively)

[1439] Similar to the above section, the cloud-based system may also be configured to monitor smart instrument configurations and, more generally, configurations that utilize multiple smart instruments, such as an operating room preparing for surgery. For similar reasons as described above, such as to improve medical efficacy and efficiency, it may be useful to compare a procedural setup at any particular medical facility to aggregate data pertaining to the procedural setups at multiple other medical facilities.

[1440] The cloud-based system of the present disclosure may be configured to aggregate data pertaining to smart medical instrument configurations and operating room (OR) setups that utilize multiple smart medical instruments. The smart medical instruments may include: manual devices that are communicatively coupled to a medical data tower and are configured to generate sensor data; and robotic instruments that perform procedures in a more automated fashion.

[1441] The cloud-based system may be configured to detect irregularities in an OR setup, either pertaining to what devices are present in the room and/or what materials are used to create a product mix for a medical procedure. The irregularities may be based on comparing the materials and equipment present in the OR with other setups from other medical facilities for a similar situation. The cloud system may then generate a change in firmware, software, or other settings and transmit those changes to the surgical devices like a device update.

[1442] In this way, the cloud-based system of the present disclosure may be able to identify errors and find more accurate causalities that give rise to best practices at a particular facility, which can then be disseminated to all of the other facilities. Furthermore, the cloud-based system may be able to provide the data from all of the disparate sources that no single facility may be able to do on its own.

This can lead to safe and more efficient operating room procedures and medical practices in general.

[1443] In some aspects, the cloud system may be configured to provide recommendations of instrument configurations, and even generate the appropriate device settings changes, to customize performance to that of a pre-specified user.

[1444] For example, the cloud system may focus on a surgical device user or surgeon based on a comparison of current usage of a device with the historic trends of a larger data set. As some examples, the cloud system may provide recommendations of what type of cartridge to use based on what the user has previously used for the particular procedure or just what the particular surgeon desires in general. The cloud system may access data based on the particular surgeon, the type of procedure, and the type of instruments used in order to make this determination.

[1445] As another example, the cloud system may provide a recommendation based on an identified anatomy indicated in a display of the cartridge. As another example, the cloud system may provide a recommendation by referring to a baseline surgical device clamping and firing speed, based on local previous usage data that it has stored in its memory.

[1446] As yet another example, the cloud system may conduct a comparison of current device tissue interaction against a historical average for the same surgeon, or for the same step in the same procedure for a segment of surgeons in the database. The cloud system again may have access to all steps used to perform a procedure, and may access a catalog of all data when performing a particular step in a procedure across all surgeons who have ever performed that procedure in its network. The recommendation may also come from an analysis of how the current surgical device has been observed to interact with tissue historically. This type of analysis may be useful because it is often not the case that large amounts of live patient data can be collected for how a surgical device interacts precisely with the tissue. Furthermore, a surgeon typically knows only his or her experience, and does not have outside knowledge of what other surgeons experience for the same procedure. The cloud, on the other hand, is capable of collecting all of this data and providing new insights that any individual surgeon would not know alone.

[1447] As another example: In stapling, more than one of the following are known: cartridge color, stapler type, procedure, procedure step, patient information, clamp force over time, prior firing information, end effector deformations, etc. This information is compared against a historical average for a similar dataset. The current situation is compared against this average, informing the user about the nature of the current firing.

[1448] FIG. 186 provides a graphical illustration of a type of example analysis the cloud system may perform to provide these recommendations, according to some aspects. In this example, chart 7500 shows data for parenchyma staple firing analysis. In the bar graphs 7502 are various types of staples used, where each color of staple reflects a different amount of force applied to the surgical site. The y axis (on the left) associated with the bar graphs 7502 reflects a percent level of usage of that type of staple color, and each color shows bar graphs for three different categories: regional average usage (in Japan in this case), global average usage with best outcomes, and the local facility average usage. Based on this data, the cloud system may be config-

ured to develop a recommendation for what staples to change to for a given situation. A series of suggested actions is shown in chart 7506 as a result. The chart 7500 also shows a set of line graphs 7504 that reflect a percentage of prolonged air leaks (the y axis on the right) for each color used, and for each type of category (regional, global average, facility average). If staples are too thick and do not match the level of tissue thickness, there could be holes in the staples that lead to undesirable air leaks. Here, the cloud system may provide a recommendation based on all of the data shown as well as data not shown, according to some aspects. The cloud system may simply provide a recommendation in the form of a letter as the label, and the surgeon may verify whether the data supports such a finding and decide to accept the cloud system's recommendation.

[1449] As another example, the cloud system may be configured to provide a recommendation of ultrasonic blade lengths or capacities based on likely to encounter vascular structures in a procedure. Similar to what is described above in reference to FIG. 186, the cloud system may collect the relevant data for blade lengths, and their outcomes that have been obtained from multiple surgical hubs, and illustrate the various outcomes for using different blade lengths on a particular procedure. A recommendation may be provided in a graphical display where the surgeon can verify the recommendation using the graphical presentation created by the cloud system.

[1450] In some aspects, the cloud system is also configured to provide recommendations to the staff about which devices to pull for an upcoming procedure. These recommendations may be based on a combination of surgeon preference (pick list) against historical device utilization rates for the same procedures performed by some segment of the larger database, as well as average recommendations or utilizations across different facilities that produce the best results. The data may be obtained by pairing good outcomes with the metadata, such as what devices were used to achieve those good outcomes. Recommendations can be influenced by other factors, including patient information, demographic data, etc.

[1451] Relatedly, in some aspects, the cloud system may also provide identification of pulled instruments that might not be the preferred device for a given procedure. The blacklisting of sorts can more clearly eliminate any obviously flawed uses of devices to help surgeons make the best decisions. This data may be obtained from manufacturer input, analysis of poor outcomes, specific input provided to the cloud system, and so on.

[1452] In addition, based on interrogating tissue for properties (elasticity, impedance, perfusion rate), a specific device with a given parameter set (clamp preload) could be suggested to be used from current stock in inventory by the cloud system. Some of the metadata associated with the outcomes of past procedures may include a description of the type of tissue being operated on, and an associated description of the physical characteristics of that tissue. The cloud system may then draw trends or patterns based on different types of procedures, but having in common all procedures that deal with similar types of tissue. This kind of analysis may be used as a secondary recommendation, when a new or unknown procedure must take place and new suggestions are welcome. If the recommendation is accepted, the cloud system may be configured to generate the change in parameters and transmit them to the intercon-

nected medical device, through the surgical hub, to make the medical device readily available for use in the adjusted procedure.

[1453] In some aspects, the device setup recommendations can include suggestions of adjuncts for devices based on the pre-surgery imaging or locally collected data during the beginning of a procedure. That is, this suggestion of adjuncts may be for use on or with devices based on the local correlation of use to efficacy of the device. As an example, based on a given procedure, surgeon, and patient information, bleeding in a case must be tightly controlled, and therefore the cloud system may conclude that a buttress is recommended on all staple firings.

[1454] In some aspects, the cloud system may also be configured to provide awareness of any newly-launched products that are available and suitable for operation as well as instructions for use (IFU). The data may be gathered from one or more surgical hubs, or from direct factory input for the newly-launched products. The cloud system can download the information and make the information displayable to multiple medical hubs across multiple facilities.

[1455] In some aspects, regarding any of the above examples for recommendations being provided by the cloud system, the cloud system may also conversely provide alerts or other signals when a device or suggested setup is not followed or is disregarded. The cloud system may be configured to access procedural data from a surgical hub during a surgical procedure, for example. The surgical hub may collect data for what type of devices are in use during a procedure. The cloud system may monitor the progress of the procedure by verifying if an accepted method or device is used in the correct or prescribed order for the procedure. If there is a deviation, in that a particular device is not expected or a step is missed, the cloud system may send an alert to the surgical hub that a particular device is not being used properly, as an example. This would occur in real time, as the timing of the procedure is important for the patient's safety.

#### Medical Facility Segmented Individualization of Instrument Function

[1456] In some aspects, the cloud-based system may also be configured to provide recommendations or automatically adjust surgical instrument settings to account for specific differences at a medical facility. While there are a number of similarities that can be normalized across multiple facilities, there may also be particular differences that should be accounted for. For example, patient demographic differences, patient physiological differences more native to a local population, procedural differences—for example preferences by each individual surgeon—and region specific instrument availability or other differences may inspire certain adjustments to be made at any particular medical facility.

[1457] The cloud-based system of the present disclosure may be configured to aggregate not only data pertaining to smart medical instrument configurations and operating room (OR) setups that utilize multiple smart medical instruments, but also data that highlight specific differences that may be unique to that region or that particular medical facility. The cloud-based system may then factor in adjustments to device settings or recommendations to changes in procedures based on these differences. For example, the cloud-based system may first provide a baseline recommendation for how a

smart instrument should be used, based on best practices discovered in the aggregate data. Then, the cloud-based system may augment the recommendation to account for one or more unique differences specific to a medical facility. Examples of these differences are described above. The cloud-based system may be made aware of what demographics and patient data gave rise to the optimal baseline procedure, and then compare the local facility demographics and patient data against that. The cloud-based system may develop or extrapolate a correlation from that baseline setting in order to develop an adjustment or offset that accounts for the differences in demographics and patient data.

**[1458]** In this way, the cloud-based system of the present disclosure may be able to make optimal adjustments specific to each medical facility or even specific to each operating room, or surgeon. The adjustments may offer improved performance that take into account the observed best practices as well as any unique differences.

**[1459]** In some aspects, the cloud system may be configured to provide changes to instrument variation of usage to improve outcomes. For example, the cloud system may determine a localized undesirable effect that is due to a specific manner of utilizing a surgical device. FIG. 187 provides an illustration of how the cloud system may conduct analysis to identify a statistical correlation to a local issue that is tied to how a device is used in the localized setting. The cloud 7600 may aggregate usage data of all types of devices and record their outcomes. The data set may be filtered down to only those outcomes that utilized the particular device in question. The cloud system may then perform statistical analysis to determine if there is a trend in how the procedures are performed at a particular facility when utilizing that device. A pattern may emerge that suggests there is a consistent flaw in how the device is used at that facility, represented as the data points 7602 that demonstrate the statistical correlation. Additional data may then be examined, to see if a second pattern may arise in comparison to how others are using the device in the aggregate. A suggestion may be provided once a pattern is identified and addressed to the local outlier 7604. In other cases, the cloud system may provide a facility-specific update to the device to offset the local practice of how that device is used.

**[1460]** In some aspects, the cloud system may be configured to communicate the deviation to the specific user and the recommendation of a differing technique or usage to improve outcomes from the specific device. The cloud system may transmit the data for display at the surgical hub to illustrate what changes ought to be made.

**[1461]** As an example: A stapler configured with a means to sense the force required to clamp the device transmits data indicating that the clamp force is still rapidly changing (viscoelastic creep) when the surgeon initiates firing of the staple, and it is observed that the staple line bleeds more often than expected. The cloud system and/or device is able to communicate a need to wait longer (e.g., 15 seconds) before firing the device to improve outcomes. This may be based on performing the statistical analysis described in FIG. 187 using data points from similar procedures aggregated from multiple surgeons and multiple facilities. In the moment of the surgery, it would be infeasible or impractical for anybody on the surgery team to come to these conclu-

sions without the help of the cloud system aggregating such knowledge and arriving at such conclusions.

**[1462]** In some aspects, the cloud system may also be configured for intentional deployment of control algorithms to devices with an in-use criteria meeting specific criteria. For regional differences, the cloud system may adjust the control algorithms of various surgical devices. A different amount of force may be applied to a device for patients in a different demographic, for example. As another example, surgeons may have different uses for a type of surgical device, and control algorithms can be adjusted to account for this. The cloud system may be configured to send out a wide area update to a device, and may target the regional and specific instrument IDs which allow for targeted updates to their control programs.

**[1463]** In some aspects, the cloud system may provide for coding of the serial numbers of sales units and/or individual devices, which enables updated control programs to be pushed to a specific device or specific groups of devices based on meeting a specific criteria or threshold.

**[1464]** In addition, according to some aspects, the cloud system may be configured to perform analysis of peri-operative data against outcomes data seeking correlations that identify exceptional results (positive and negative). The analysis may be performed at multiple levels (e.g., individual, hospital, and geographic (e.g., city, county, state, country, etc.) filters). Furthermore, regional corroboration of improved outcomes may be target for only a limited geographic area, as it is known that the changes occur only within a limited area. The ability to tune devices to regional preferences, techniques, and surgical preferences may allow for nuanced improvements for regionally specific areas.

**[1465]** In addition to directly changing instrument settings, the cloud system may also be configured to provide recommendations on different instruments or equivalent device suggestions due to regional availability. That is, an equivalent suggestion to a device to perform a particular function may be recommended by the cloud system, in the event a device is lacking and a particular region has an excess or general availability of the different device that may be used to serve an equivalent purpose.

**[1466]** For example, the cloud system may determine that PPH hemorrhoid stapling devices or curved cutter 30 devices are only available in Italy due to a unique procedure configuration or teaching hospital procedure design. As another example, the cloud system may determine that there is an Asia-specific TX and open vascular stapler use due to cost sensitivity, lack of laparoscopic adoption, and teaching hospital preferred techniques and patient thoracic cavity size. As another example, the cloud system may provide awareness messages to OR staff of sub-standard knock-off products available in a certain region. This data may be derived from an ingestion of information from multiple sources, such as inputs provided by experts and doctors, and employing machine learning and natural language processing to interpret trends and news related to a local area. FIG. 188 provides a graphical illustration of an example of how some devices may satisfy an equivalent use compared to an intended device. Here, a circular stapling device 7702 is compared to a compression ring 7704 for use in a PPH stapler 7700 for hemorrhoidopexy procedures. The type of analysis performed to reach the recommendations by the cloud system may be similar to those described in FIG. 187. The cloud system may provide a display of this suggestion,

as well as an analysis of its efficiency and resource utilization, in example display 7706 that may be shown at a display in a surgical hub. In this case, the instrument cost is compared, as well as time and efficacy for each type of instrument. The cloud system may derive these recommendations by obtaining usage examples from different facilities, observing how other facilities and doctors treat the same procedure.

[1467] In some aspects, the cloud system may also be configured to provide a surgical hub decision tree and local suggestions of post-operative care, based on data processed during the procedure and Cloud Analytics trending of results or performance of the devices aggregated from larger population sets.

[1468] In some aspects, the cloud system may provide update-able decision trees for post-operative care suggestions, based on device measured situational usage. The post-operative care decisions may initially be derived from traditionally known responses that doctors would normally recommend. Once additional data becomes available, say from aggregating types of post-operative care from other facilities, or from analyzing new types of care from literature or from research on new surgical devices, the decision can be updated by the cloud system. The decision tree may be displayable at a surgical hub and in a graphical form.

[1469] In using this decision tree, feedback can be provided for each node to state how effective the current solutions are. The data may be inputted based on whatever feedback patients may provide. A doctor or data admin need not perform any analysis at the time, but the cloud system can aggregate all of the data and observe what trends may arise. Feedback can then be provided to update the decision tree.

[1470] In some aspects, the cloud system may incorporate operative data & device performance to propose post-operative monitoring & activities. For example, various patient measures may change what decisions in post-operative care should be taken. These measurements can include but are not limited to: (a) blood pressure; (b) low hematocrit; (c) PTT (partial thromboplastin time); (d) INR (international normalized ratio); (e) Oxygen saturation; (f) Ventilation changes; and (g) X-Ray data.

[1471] As another example, anesthesia protocol can dictate what post-operative decisions should be taken. This may account for: (a) any fluids administered; (b) Anesthesia time; and (3) Medications, as some non-limiting examples.

[1472] As another example, the types of medications may also play a role. The application of Warfarin is one notable example. A patient post-operatively has abnormal PTT and INR, for example. Because the patient is on Warfarin, potential treatments could include vitamin K, factor 7, or the delivery of plasma (fpp). Plavix can be another example. A patient post-operatively has abnormal PTT and INR. Because patient is on Plavix, potential treatments for Warfarin would be ineffective. Deliver platelets instead may be the suggestion in the decision tree.

[1473] As a fourth example, post-operative instructions may be provided that are dependent on the type of procedure. Some non-limiting examples include colorectal time to solid food (motility); and (b) time to physical activity & PT. These varying decisions can be reflected in the decision tree, and all of the types of branching decisions may be stored in the cloud system and updated when additional data is gained from any connected facility.

[1474] FIG. 189 provides various examples of how some data may be used as variables in deciding how the post-operative decision tree may branch out. As shown, some factors 7802 may include the parameters used in surgical devices, such as the force to fire (FTF) used in an operation, or the force to close (FTC) used in a surgical device. Graph 7800 shows a visual depiction of how the FTC and FTF curves may interrelate with one another. Other factors include compression rate, wait time, and staple adaptability. Based on some of these variables, a type of post-operative care should be adjusted. In this case, a multi-factored analysis is applied, which may be too complex to calculate or modify without the aid of the processing power of a system like the cloud system. This example suggests that a decision tree 7804 provided by the cloud system can be more than a simple two dimensional decision tree. To account for multiple variables to make a single decision, the decision tree generated by the cloud may be visually available for perhaps just a portion, and the ultimate conclusion may have to be displayed without a full display of all of the other branches that were not considered. The chart 7806 may be an example of providing additional information of how to respond within the decision tree.

#### Adaptive Control Program Updates for Surgical Devices

[1475] Modular devices include the modules (as described in connection with FIGS. 3 and 9, for example) that are receivable within a surgical hub and the surgical devices or instruments that can be connected to the various modules. The modular devices include, for example, intelligent surgical instruments, medical imaging devices, suction/irrigation devices, smoke evacuators, energy generators, ventilators, and insufflators. Various operations of the modular devices described herein can be controlled by one or more control algorithms. The control algorithms can be executed on the modular device itself, on the surgical hub to which the particular modular device is paired, or on both the modular device and the surgical hub (e.g., via a distributed computing architecture). In some exemplifications, the modular devices' control algorithms control the devices based on data sensed by the modular device itself (i.e., by sensors in, on, or connected to the modular device). This data can be related to the patient being operated on (e.g., tissue properties or insufflation pressure) or the modular device itself (e.g., the rate at which a knife is being advanced, motor current, or energy levels). For example, a control algorithm for a surgical stapling and cutting instrument can control the rate at which the instrument's motor drives its knife through tissue according to resistance encountered by the knife as it advances.

[1476] Although an "intelligent" device including control algorithms that respond to sensed data can be an improvement over a "dumb" device that operates without accounting for sensed data, if the device's control program does not adapt or update over time in response to collected data, then the devices may continue to repeat errors or otherwise perform suboptimally. One solution includes transmitting operational data collected by the modular devices in combination with the outcomes of each procedure (or step thereof) to an analytics system. In one exemplification, the procedural outcomes can be inferred by a situational awareness system of a surgical hub to which the modular devices are paired, as described in U.S. patent application Ser. No.

15/940,654, titled SURGICAL HUB SITUATIONAL AWARENESS, which is herein incorporated by reference in its entirety. The analytics system can analyze the data aggregated from a set of modular devices or a particular type of modular device to determine under what conditions the control programs of the analyzed modular devices are controlling the modular devices suboptimally (i.e., if there are repeated faults or errors in the control program or if an alternative algorithm performs in a superior manner) or under what conditions medical personnel are utilizing the modular devices suboptimally. The analytics system can then generate an update to fix or improve the modular devices' control programs. Different types of modular devices can be controlled by different control programs; therefore, the control program updates can be specific to the type of modular device that the analytics system determines is performing suboptimally. The analytics system can then push the update to the appropriate modular devices connected to the analytics system through the surgical hubs.

[1477] FIG. 190 illustrates a block diagram of a computer-implemented adaptive surgical system 9060 that is configured to adaptively generate control program updates for modular devices 9050, in accordance with at least one aspect of the present disclosure. In one exemplification, the surgical system includes a surgical hub 9000, multiple modular devices 9050 communicably coupled to the surgical hub 9000, and an analytics system 9100 communicably coupled to the surgical hub 9000. Although a single surgical hub 9000 is depicted, it should be noted that the surgical system 9060 can include any number of surgical hubs 9000, which can be connected to form a network of surgical hubs 9000 that are communicably coupled to the analytics system 9100. In one exemplification, the surgical hub 9000 includes a processor 9010 coupled to a memory 9020 for executing instructions stored thereon and a data relay interface 9030 through which data is transmitted to the analytics system 9100. In one exemplification, the surgical hub 9000 further includes a user interface 9090 having an input device 9092 (e.g., a capacitive touchscreen or a keyboard) for receiving inputs from a user and an output device 9094 (e.g., a display screen) for providing outputs to a user. Outputs can include data from a query input by the user, suggestions for products or mixes of products to use in a given procedure, and/or instructions for actions to be carried out before, during, or after surgical procedures. The surgical hub 9000 further includes an interface 9040 for communicably coupling the modular devices 9050 to the surgical hub 9000. In one aspect, the interface 9040 includes a transceiver that is communicably connectable to the modular device 9050 via a wireless communication protocol. The modular devices 9050 can include, for example, surgical stapling and cutting instruments, electrosurgical instruments, ultrasonic instruments, insufflators, respirators, and display screens. In one exemplification, the surgical hub 9000 can further be communicably coupled to one or more patient monitoring devices 9052, such as EKG monitors or BP monitors. In another exemplification, the surgical hub 9000 can further be communicably coupled to one or more databases 9054 or external computer systems, such as an EMR database of the medical facility at which the surgical hub 9000 is located.

[1478] When the modular devices 9050 are connected to the surgical hub 9000, the surgical hub 9000 can sense or receive perioperative data from the modular devices 9050 and then associate the received perioperative data with

surgical procedural outcome data. The perioperative data indicates how the modular devices 9050 were controlled during the course of a surgical procedure. The procedural outcome data includes data associated with a result from the surgical procedure (or a step thereof), which can include whether the surgical procedure (or a step thereof) had a positive or negative outcome. For example, the outcome data could include whether a patient suffered from postoperative complications from a particular procedure or whether there was leakage (e.g., bleeding or air leakage) at a particular staple or incision line. The surgical hub 9000 can obtain the surgical procedural outcome data by receiving the data from an external source (e.g., from an EMR database 9054), by directly detecting the outcome (e.g., via one of the connected modular devices 9050), or inferring the occurrence of the outcomes through a situational awareness system. For example, data regarding postoperative complications could be retrieved from an EMR database 9054 and data regarding staple or incision line leakages could be directly detected or inferred by a situational awareness system. The surgical procedural outcome data can be inferred by a situational awareness system from data received from a variety of data sources, including the modular devices 9050 themselves, the patient monitoring device 9052, and the databases 9054 to which the surgical hub 9000 is connected.

[1479] The surgical hub 9000 can transmit the associated modular device 9050 data and outcome data to the analytics system 9100 for processing thereon. By transmitting both the perioperative data indicating how the modular devices 9050 are controlled and the procedural outcome data, the analytics system 9100 can correlate the different manners of controlling the modular devices 9050 with surgical outcomes for the particular procedure type. In one exemplification, the analytics system 9100 includes a network of analytics servers 9070 that are configured to receive data from the surgical hubs 9000. Each of the analytics servers 9070 can include a memory and a processor coupled to the memory that is executing instructions stored thereon to analyze the received data. In some exemplifications, the analytics servers 9070 are connected in a distributed computing architecture and/or utilize a cloud computing architecture. Based on this paired data, the analytics system 9100 can then learn optimal or preferred operating parameters for the various types of modular devices 9050, generate adjustments to the control programs of the modular devices 9050 in the field, and then transmit (or "push") updates to the modular devices' 9050 control programs.

[1480] Additional detail regarding the computer-implemented interactive surgical system 9060, including the surgical hub 9000 and various modular devices 9050 connectable thereto, are described in connection with FIGS. 9-10.

[1481] FIG. 191 illustrates a logic flow diagram of a process 9200 for updating the control program of a modular device 9050, in accordance with at least one aspect of the present disclosure. In the following description of the process 9200, reference should also be made to FIG. 190. The process 9200 can be executed by, for example, one or more processors of the analytics servers 9070 of the analytics system 9100. In one exemplification, the analytics system 9100 can be a cloud computing system. For economy, the following description of the process 9200 will be described as being executed by the analytics system 9100; however, it should be understood that the analytics system 9100

includes processor(s) and/or control circuit(s) that are executing the describe steps of the process 9200.

[1482] The analytics system 9100 receives 9202 modular device 9050 perioperative data and surgical procedural outcome data from one or more of the surgical hubs 9000 that are communicably connected to the analytics system 9100. The perioperative data includes preoperative data, intraoperative data, and/or postoperative data detected by a modular device 9050 in association with a given surgical procedure. For modular devices 9050 or particular functions of modular devices 9050 that are manually controlled, the perioperative data indicates the manner in which a surgical staff member operated the modular devices 9050. For modular devices 9050 or particular functions of modular devices 9050 that are controlled by the modular devices' control programs, the perioperative data indicates the manner in which the control programs operated the modular devices 9050. The manner in which the modular devices 9050 function under particular sets of conditions (either due to manual control or control by the modular devices' 9050 control programs) can be referred to as the "operational behavior" exhibited by the modular device 9050. The modular device 9050 perioperative data includes data regarding the state of the modular device 9050 (e.g., the force to fire or force to close for a surgical stapling and cutting instrument or the power output for an electrosurgical or ultrasonic instrument), tissue data measured by the modular device 9050 (e.g., impedance, thickness, or stiffness), and other data that can be detected by a modular device 9050. The perioperative data indicates the manner in which the modular devices 9050 were programmed to operate or were manually controlled during the course of a surgical procedure because it indicates how the modular devices 9050 functioned in response to various detected conditions.

[1483] The surgical procedural outcome data includes data pertaining to an overall outcome of a surgical procedure (e.g., whether there was a complication during the surgical procedure) or data pertaining to an outcome of a specific step within a surgical procedure (e.g., whether a particular staple line bled or leaked). The procedural outcome data can, for example, be directly detected by the modular devices 9050 and/or surgical hub 9000 (e.g., a medical imaging device can visualize or detect bleeding), determined or inferred by a situational awareness system of the surgical hub 9000 as described in U.S. patent application Ser. No. 15/940,654, or retrieved from a database 9054 (e.g., an EMR database) by the surgical hub 9000 or the analytics system 9100. The procedural outcome data can include whether each outcome represented by the data was a positive or negative result. Whether each outcome was positive or negative can be determined by the modular devices 9050 themselves and included in the perioperative data transmitted to the surgical hubs 9000 or determined or inferred by the surgical hubs 9000 from the received perioperative data. For example, the procedural outcome data for a staple line that bled could include that the bleeding represented a negative outcome. Similarly, the procedural outcome data for a staple line that did not bleed could include that the lack of bleeding represented a positive outcome. In another exemplification, the analytics system 9100 can be configured to determine whether a procedural outcome is a positive or negative outcome based upon the received procedural outcome data. In some exemplifications, correlating the modular device 9050 data to positive or negative procedural outcomes

allows the analytics system 9100 to determine whether a control program update should be generated 9208.

[1484] Upon the analytics system 9100 receiving 9202 the data, the analytics system 9100 analyzes the modular device 9050 and procedural outcome data to determine 9204 whether the modular devices 9050 are being utilized suboptimally in connection with the particular procedure or the particular step of the procedure. A modular device 9050 can be controlled suboptimally if the particular manner in which the modular device 9050 is being controlled is repeatedly causing an error or if an alternative manner of controlling the modular device 9050 is superior under the same conditions. The analytics system 9100 can thus determine whether a modular device 9050 is being controlled suboptimally (either manually or by its control program) by comparing the rate of positive and/or negative outcomes produced by the modular device 9050 relative to set thresholds or the performance of other modular devices 9050 of the same type.

[1485] For example, the analytics system 9100 can determine whether a type of modular device 9050 is being operated suboptimally if the rate of negative procedural outcomes produced by the modular device 9050 under a particular set of conditions in association with a particular operational behavior exceeds an average or threshold level. As a specific example, the analytics system 9100 can analyze 9204 whether a control program for a surgical stapling instrument that dictates a particular force to fire (or ranges of forces to fire) is suboptimal for a particular tissue thickness and tissue type. If the analytics system 9100 determines that the instrument generates an abnormally high rate of leaky staple lines when fired at the particular force (e.g., causing the staples to be malformed, not fully penetrate the tissue, or tear the tissue) relative to an average or threshold staple line leakage rate, then the analytics system 9100 can determine that the control program for the surgical stapling instrument is performing suboptimally given the tissue conditions.

[1486] As another example, the analytics system 9100 can determine whether a type of modular device 9050 is being operated suboptimally if the rate of positive outcomes produced by an alternative manner of control under a particular set of conditions in association with a particular operational behavior exceeds the rate of positive outcomes generated by the analyzed manner of control under the same conditions. In other words, if one subpopulation of the type of modular device 9050 exhibits a first operational behavior under a certain set of conditions and a second subpopulation of the same type of modular device 9050 exhibits a second operational behavior under the same set of conditions, then the analytics system 9100 can determine whether to update the control programs of the modular devices 9050 according to whether the first or second operational behavior is more highly correlated to a positive procedural outcome. As a specific example, the analytics system 9100 can analyze 9204 whether a control program for an RF electrosurgical or ultrasonic instrument that dictates a particular energy level is suboptimal for a particular tissue type and environmental conditions. If the analytics system 9100 determines that a first energy level given a set of tissue conditions and environmental conditions (e.g., the instrument being located in a liquid-filled environment, as in an arthroscopic procedure) produces a lower rate of hemostasis than a second energy level, then the analytics system 9100 can determine that the control program for the electrosurgical or ultrasonic

instrument dictating the first energy level is performing suboptimally for the given tissue and environmental conditions.

[1487] After analyzing 9204 the data, the analytics system 9100 determines 9206 whether to update the control program. If the analytics system 9100 determines that the modular device 9050 is not being controlled suboptimally, then the process 9200 continues along the NO branch and the analytics system 9100 continues analyzing 9204 received 9202 data, as described above. If the analytics system 9100 determines that the modular device 9050 is being controlled suboptimally, then the process 9200 continues along the YES branch and the analytics system 9100 generates 9208 a control program update. The generated 9208 control program update includes, for example, a new version of the control program for the particular type of modular device 9050 to overwrite the prior version or a patch that partially overwrites or supplements the prior version.

[1488] The type of control program update that is generated 9208 by the analytics system 9100 depends upon the particular suboptimal behavior exhibited by the modular device 9050 that is identified by the analytics system 9100. For example, if the analytics system 9100 determines that a particular force to fire a surgical stapling instrument results in an increased rate of leaking staple lines, then the analytics system 9100 can generate 9208 a control program update that adjusts the force to fire from a first value to a second value that corresponds to a higher rate of non-leaking staple lines or a lower rate of leaking staple lines. As another example, if the analytics system 9100 determines that a particular energy level for an electrosurgical or ultrasonic instrument produces a low rate of hemostasis when the instrument is used in a liquid-filled environment (e.g., due to the energy dissipating effects of the liquid), then the analytics system 9100 can generate 9208 a control program update that adjusts the energy level of the instrument when it is utilized in surgical procedures where the instrument will be immersed in liquid.

[1489] The type of control program update that is generated 9208 by the analytics system 9100 also depends upon whether the suboptimal behavior exhibited by the modular device 9050 is caused by manual control or control by the control program of the modular device 9050. If the suboptimal behavior is caused by manual control, the control program update can be configured to provide warnings, recommendations, or feedback to the users based upon the manner in which they are operating the modular devices 9050. Alternatively, the control program update can change the manually controlled operation of the modular device 9050 to an operation that is controlled by the control program of the modular device 9050. The control program update may or may not permit the user to override the control program's control of the particular function. In one exemplification, if the analytics system 9100 determines 9204 that surgeons are manually setting an RF electrosurgical instrument to a suboptimal energy level for a particular tissue type or procedure type, then the analytics system 9100 can generate 9208 a control program update that provides an alert (e.g., on the surgical hub 9000 or the RF electrosurgical instrument itself) recommending that the energy level be changed. In another exemplification, the generated 9208 control program update can automatically set the energy level to a default or recommended level given the particular

detected circumstances, which could then be changed as desired by the medical facility staff. In yet another exemplification, the generated 9208 control program update can automatically set the energy level to a set level determined by the analytics system 9100 and not permit the medical facility staff to change the energy level. If the suboptimal behavior is caused by the control program of the modular device 9050, then the control program update can alter how the control program functions under the particular set of circumstances that the control program is performing suboptimally under.

[1490] Once the control program update has been generated 9208 by the analytics system 9100, the analytics system 9100 then transmits 9210 or pushes the control program update to all of the modular devices 9050 of the relevant type that are connected to the analytics system 9100. The modular devices 9050 can be connected to the analytics system 9100 through the surgical hubs 900, for example. In one exemplification, the surgical hubs 9000 are configured to download the control program updates for the various types of modular devices 9050 from the analytics system 9100 each time an update is generated 9208 thereby. When the modular devices 9050 subsequently connect to or pair with a surgical hub 9000, the modular devices 9050 then automatically download any control program updates therefrom. In one exemplification, the analytics system 9100 can thereafter continue receiving 9202 and analyzing 9204 data from the modular devices 9050, as described above.

[1491] In one exemplification, instead of the modular devices 9050 transmitting recorded data to a surgical hub 9000 to which the modular devices 9050 are connected, the modular devices 9050 are configured to record the perioperative data and the procedural outcome data on a memory of the modular device 9050. The data can be stored for indefinitely or until the data is downloaded from the modular devices 9050. This allows the data to be retrieved at a later time. For example, the modular devices 9050 could be returned to the manufacturer after they are utilized in a surgical procedure. The manufacturer could then download the data from the modular devices 9050 and then analyze the data as described above to determine whether a control program update should be generated for the modular devices 9050. In one exemplification, the data could be uploaded to an analytics system 9100 for analysis, as described above. The analytics system 9100 could then generate update control programs according to the recorded data and then either incorporate that update in future manufactured product or push the update to modular devices 9050 currently in the field.

[1492] In order to assist in the understanding of the process 9200 illustrated in FIG. 191 and the other concepts discussed above, FIG. 192 illustrates a diagram of an illustrative analytics system 9100 updating a surgical instrument control program, in accordance with at least one aspect of the present disclosure. In one exemplification, a surgical hub 9000 or network of surgical hubs 9000 is communicably coupled to an analytics system 9100, as illustrated above in FIG. 190. The analytics system 9100 is configured to filter and analyze modular device 9050 data associated with surgical procedural outcome data to determine whether adjustments need to be made to the control programs of the modular devices 9050. The analytics system 9100 can then push updates to the modular devices 9050 through the surgical hubs 9000, as necessary. In the depicted exempli-



fication, the analytics system **9100** comprises a cloud computing architecture. The modular device **9050** perioperative data received by the surgical **9000** hubs from their paired modular devices **9050** can include, for example, force to fire (i.e., the force required to advance a cutting member of a surgical stapling instrument through a tissue), force to close (i.e., the force required to clamp the jaws of a surgical stapling instrument on a tissue), the power algorithm (i.e., change in power over time of electrosurgical or ultrasonic instruments in response to the internal states of the instrument and/or tissue conditions), tissue properties (e.g., impedance, thickness, stiffness, etc.), tissue gap (i.e., the thickness of the tissue), and closure rate (i.e., the rate at which the jaws of the instrument clamped shut). It should be noted that the modular device **9050** data that is transmitted to the analytics system **9100** is not limited to a single type of data and can include multiple different data types paired with procedural outcome data. The procedural outcome data for a surgical procedure (or step thereof) can include, for example, whether there was bleeding at the surgical site, whether there was air or fluid leakage at the surgical site, and whether the staples of a particular staple line were formed properly. The procedural outcome data can further include or be associated with a positive or negative outcome, as determined by the surgical hub **9000** or the analytics system **9100**, for example. The modular device **9050** data and the procedural outcome data corresponding to the modular device **9050** perioperative data can be paired together or otherwise associated with each other when they are uploaded to the analytics system **9100** so that the analytics system **9100** is able to recognize trends in procedural outcomes based on the underlying data of the modular devices **9050** that produced each particular outcome. In other words, the analytics system **9100** can aggregate the modular device **9050** data and the procedural outcome data to search for trends or patterns in the underlying device modular data **9050** that can indicate adjustments that can be made to the modular devices' **9050** control programs.

[1493] In the depicted exemplification, the analytics system **9100** executing the process **9200** described in connection with FIG. **190** is receiving **9202** modular device **9050** data and procedural outcome data. When transmitted to the analytics system **9100**, the procedural outcome data can be associated or paired with the modular device **9050** data corresponding to the operation of the modular device **9050** that caused the particular procedural outcome. The modular device **9050** perioperative data and corresponding procedural outcome data can be referred to as a data pair. The data is depicted as including a first group **9212** of data associated with successful procedural outcomes and a second group **9214** of data associated with negative procedural outcomes. For this particular exemplification, a subset of the data **9212**, **9214** received **9202** by the analytics system **9100** is highlighted to further elucidate the concepts discussed herein.

[1494] For a first data pair **9212a**, the modular device **9050** data includes the force to close (FTC) overtime, the force to fire (FTF) over time, the tissue type (parenchyma), the tissue conditions (the tissue is from a patient suffering from emphysema and had been subject to radiation), what number firing this was for the instrument (third), an anonymized time stamp (to protect patient confidentiality while still allowing the analytics system to calculate elapsed time between firings and other such metrics), and an anonymized patient identifier (**002**). The procedural outcome data

includes data indicating that there was no bleeding, which corresponds to a successful outcome (i.e., a successful firing of the surgical stapling instrument). For a second data pair **9212b**, the modular device **9050** data includes the wait time prior the instrument being fired (which corresponds to the first firing of the instrument), the FTC over time, the FTF over time (which indicates that there was a force spike near the end of the firing stroke), the tissue type (1.1 mm vessel), the tissue conditions (the tissue had been subject to radiation), what number firing this was for the instrument (first), an anonymized time stamp, and an anonymized patient identifier (**002**). The procedural outcome data includes data indicating that there was a leak, which corresponds to a negative outcome (i.e., a failed firing of the surgical stapling instrument). For a third data pair **9212c**, the modular device **9050** data includes the wait time prior the instrument being fired (which corresponds to the first firing of the instrument), the FTC over time, the FTF over time, the tissue type (1.8 mm vessel), the tissue conditions (no notable conditions), what number firing this was for the instrument (first), an anonymized time stamp, and an anonymized patient identifier (**012**). The procedural outcome data includes data indicating that there was a leak, which corresponds to a negative outcome (i.e., a failed firing of the surgical stapling instrument). It should be noted again that this data is intended solely for illustrative purposes to assist in the understanding of the concepts discussed herein and should not be interpreted to limit the data that is received and/or analyzed by the analytics system **9100** to generate control program updates.

[1495] When the analytics system **9100** receives **9202** perioperative data from the communicably connected surgical hubs **9000**, the analytics system **9100** proceeds to aggregate and/or store the data according to the procedure type (or a step thereof) associated with the data, the type of the modular device **9050** that generated the data, and other such categories. By collating the data accordingly, the analytics system **9100** can analyze the data set to identify correlations between particular ways of controlling each particular type of modular device **9050** and positive or negative procedural outcomes. Based upon whether a particular manner of controlling a modular device **9050** correlates to positive or negative procedural outcomes, the analytics system **9100** can determine **9204** whether the control program for the type of modular device **9050** should be updated.

[1496] For this particular exemplification, the analytics system **9100** performs a first analysis **9216a** of the data set by analyzing the peak FTF **9213** (i.e., the maximum FTF for each particular firing of a surgical stapling instrument) relative to the number of firings **9211** for each peak FTF value. In this exemplary case, the analytics system **9100** can determine that there is no particular correlation between the peak FTF **9213** and the occurrence of positive or negative outcomes for the particular data set. In other words, there are not distinct distributions for the peak FTF **9213** for positive and negative outcomes. As there is no particular correlation between peak FTF **9213** and positive or negative outcomes, the analytics system **9100** would thus determine that a control program update to address this variable is not necessary. Further, the analytics system **9100** performs a second analysis **9216b** of the data set by analyzing the wait time **9215** prior to the instrument being fired relative to the number of firings **9211**. For this particular analysis **9216b**,

the analytics system **9100** can determine that there is a distinct negative outcome distribution **9217** and a positive outcome distribution **9219**.

[1497] In this exemplary case, the negative outcome distribution **9217** has a mean of 4 seconds and the positive outcome distribution has a mean of 11 seconds. Thus, the analytics system **9100** can determine that there is a correlation between the wait time **9215** and the type of outcome for this surgical procedure step. Namely, the negative outcome distribution **9217** indicates that there is a relatively large rate of negative outcomes for wait times of 4 seconds or less. Based on this analysis **9216b** demonstrating that there is a large divergence between the negative outcome distribution **9217** and the positive outcome distribution **9219**, the analytics system **9100** can then determine **9204** that a control program update should be generated **9208**.

[1498] Once the analytics system **9100** analyzes the data set and determines **9204** that an adjustment to the control program of the particular module device **9050** that is the subject of the data set would improve the performance of the modular device **9050**, the analytics system **9100** then generates **9208** a control program update accordingly. In this exemplary case, the analytics system **9100** can determine based on the analysis **9216b** of the data set that a control program update **9218** recommending a wait time of more than 5 seconds would prevent 90% of the distribution of the negative outcomes with a 95% confidence interval. Alternatively, the analytics system **9100** can determine based on the analysis **9216b** of the data set that a control program update **9218** recommending a wait time of more than 5 seconds would result in the rate of positive outcomes being greater than the rate of negative outcomes. The analytics system **9100** could thus determine that the particular type of surgical instrument should wait more than 5 seconds before being fired under the particular tissue conditions so that negative outcomes are less common than positive outcomes. Based on either or both of these constraints for generating **9208** a control program update that the analytics system **9100** determines are satisfied by the analysis **9216b**, the analytics system **9100** can generate **9208** a control program update **9218** for the surgical instrument that causes the surgical instrument, under the given circumstances, to either impose a 5 second or longer wait time before the particular surgical instrument can be fired or causes the surgical instrument to display a warning or recommendation to the user that indicates to the user that the user should wait at least 5 seconds before firing the instrument. Various other constraints can be utilized by the analytics system **9100** in determining whether to generate **9208** a control program update, such as whether a control program update would reduce the rate of negative outcomes by a certain percentage or whether a control program update maximizes the rate of positive outcomes.

[1499] After the control program update **9218** is generated **9208**, the analytics system **9100** then transmits **9210** the control program update **9218** for the appropriate type of modular devices **9050** to the surgical hubs **9000**. In one exemplification, when a modular device **9050** that corresponds to the control program update **9218** is next connected to a surgical hub **9000** that has downloaded the control program update **9218**, the modular device **9050** then automatically downloads the update **9218**. In another exemplification, the surgical hub **9000** controls the modular device **9050** according to the control program update **9218**, rather

than the control program update **9218** being transmitted directly to the modular device **9050** itself.

[1500] In one aspect, the surgical system **9060** is configured to push down verification of software parameters and updates if modular devices **9050** are detected to be out of date in the surgical hub **9000** data stream. FIG. **193** illustrates a diagram of an analytics system **9100** pushing an update to a modular device **9050** through a surgical hub **9000**, in accordance with at least one aspect of the present disclosure. In one exemplification, the analytics system **9000** is configured to transmit a generated control program update for a particular type of modular device **9050** to a surgical hub **9000**. In one aspect, each time a modular device **9050** connects to a surgical hub **9000**, the modular device **9050** determines whether there is an updated version of its control program on or otherwise accessible via the surgical hub **9000**. If the surgical hub **9000** does have an updated control program (or the updated control program is otherwise available from the analytics system **9100**) for the particular type of modular device **9050**, then the modular device **9050** downloads the control program update therefrom.

[1501] In one exemplification, any data set being transmitted to the analytics systems **9100** includes a unique ID for the surgical hub **9000** and the current version of its control program or operating system. In one exemplification, any data set being sent to the analytics systems **9100** includes a unique ID for the modular device **9050** and the current version of its control program or operating system. The unique ID of the surgical hub **9000** and/or modular device **9050** being associated with the uploaded data allows the analytics system **9100** to determine whether the data corresponds to the most recent version of the control program. The analytics system **9100** could, for example, elect to discount (or ignore) data generated by a modular device **9050** or surgical hub **9000** being controlled by an out of date control program and/or cause the updated version of the control program to be pushed to the modular device **9050** or surgical hub **9000**.

[1502] In one exemplification, the operating versions of all modular devices **9050** the surgical hub **9000** has updated control software for could also be included in a surgical hub **9000** status data block that is transmitted to the analytics system **9100** on a periodic basis. If the analytics system **9100** identifies that the operating versions of the control programs of the surgical hub **9100** and/or any of the connectable modular devices **9050** are out of date, the analytics system **9100** could push the most recent revision of the relevant control program to the surgical hub **9000**.

[1503] In one exemplification, the surgical hub **9000** and/or modular devices **9050** can be configured to automatically download any software updates. In another exemplification, the surgical hub **9000** and/or modular devices **9050** can be configured to provide a prompt for the user to ask at the next setup step (e.g., between surgical procedures) if the user wants to update the out of date control program(s). In another exemplification, the surgical hub **9000** could be programmable by the user to never allow updates or only allow updates of the modular devices **9050** and not the surgical hub **9000** itself.

#### Adaptive Control Program Updates for Surgical Hubs

[1504] As with the modular devices **9050** described above, the surgical hubs **9000** can likewise include control pro-

grams that control the various operations of the surgical hub 9000 during the course of a surgical procedure. If the surgical hubs' 9000 control programs do not adapt over time in response to collected data, then the surgical hubs 9000 may continue to repeat errors, not provide warnings or recommendations to the surgical staff based on learned information, and not adjust to the surgical staff's preferences. One solution includes transmitting operational data from the surgical hubs 9000 that indicates how the surgical hubs 9000 are being utilized or controlled during the course of a surgical procedure to an analytics system 9100. The analytics system 9100 can then analyze the data aggregated from the network of surgical hubs 9000 connected to the analytics system 9100 to determine if a particular manner of operating the surgical hubs 9000 corresponds to improved patient outcomes or is otherwise preferred across the population of the surgical hubs 9000. In one exemplification, if a particular manner in which the surgical hubs 9000 are operated satisfies a defined condition or set of conditions, then the analytics system 9100 can determine that this particular manner should be implemented across the network of surgical hubs 9000. The analytics system 9100 can generate an update to the surgical hubs' 9000 control program to fix or improve the control program and then push the update to the surgical hubs 9000 so that the improvement is shared across every surgical hub 9000 that is connected to the analytics system 9100. For example, if a threshold number of the surgical hubs 9000 are controlled in a particular manner and/or if a particular manner of controlling the surgical hubs 9000 correlates to an improvement in the surgical procedure outcomes that exceeds a threshold level, then the analytics system 9100 can generate a control program update that controls the surgical hubs 9000 in a manner corresponding to the preferred or improved manner of control. The control program update can then be pushed to the surgical hubs 9000.

[1505] In one exemplification, an analytics system 9100 is configured to generate and push control program updates to surgical hubs 9000 in the field based on perioperative data relating to the manner in which the surgical hubs 9000 are controlled or utilized. In other words, the surgical hubs 9000 can be updated with improved decision-making abilities according to data generated from the hub network. In one aspect, external and perioperative data is collected by an analytics system. The data is then analyzed to generate a control update to improve the performance of the surgical hubs 9000. The analytics system 9100 can analyze the data aggregated from the surgical hubs 9000 to determine the preferred manner for the surgical hubs 9000 to operate, under what conditions the surgical hubs' 9000 control programs are controlling the surgical hubs 9000 suboptimally (i.e., if there are repeated faults or errors in the control program or if an alternative algorithm performs in a superior manner), or under what conditions medical personnel are utilizing the surgical hubs 9000 suboptimally. The analytics system 9100 can then push the update to the surgical hubs 9000 connected thereto.

[1506] FIG. 194 illustrates a diagram of a computer-implemented adaptive surgical system 9060 that is configured to adaptively generate control program updates for surgical hubs 9000, in accordance with at least one aspect of the present disclosure. The surgical system 9060 includes several surgical hubs 9000 that are communicably coupled to the analytics system 9100. Subpopulations of surgical

hubs 9000 (each of which can include individual surgical hubs 9000 or groups of surgical hubs 9000) within the overall population connected to the analytics system 9100 can exhibit different operational behaviors during the course of a surgical procedure. The differences in operational behavior between groups of surgical hubs 9000 within the population can result from the surgical hubs 9000 running different versions of their control program, by the surgical hubs' 9000 control programs being customized or programmed differently by local surgical staff, or by the local surgical staff manually controlling the surgical hubs 9000 differently. In the depicted example, the population of surgical hubs 9000 includes a first subpopulation 9312 that is exhibiting a first operational behavior and a second subpopulation 9314 that is exhibiting a second operational behavior for a particular task. Although the surgical hubs 9000 are divided into a pair of subpopulations 9312, 9314 in this particular example, there is no practical limit to the number of different behaviors exhibited within the population of surgical hubs 9000. The tasks that the surgical hubs 9000 can be executing include, for example, controlling a surgical instrument or analyzing a dataset in a particular manner.

[1507] The surgical hubs 9000 can be configured to transmit perioperative data pertaining to the operational behavior of the surgical hubs 9000 to the analytics system 9100. The perioperative data can include preoperative data, intraoperative data, and postoperative data. The preoperative data can include, for example, patient-specific information, such as demographics, health history, preexisting conditions, preoperative workup, medication history (i.e., medications currently and previously taken), genetic data (e.g., SNPs or gene expression data), EMR data, advanced imaging data (e.g., MRI, CT, or PET), metabolomics, and microbiome. Various additional types of patient-specific information that can be utilized by the analytics system 9100 are described by U.S. Pat. No. 9,250,172, U.S. patent application Ser. No. 13/631,095, U.S. patent application Ser. No. 13/828,809, and U.S. Pat. No. 8,476,227, each of which is incorporated by reference herein to the extent that they describe patient-specific information. The preoperative data can also include, for example, operating theater-specific information, such as geographic information, hospital location, operating theater location, operative staff performing the surgical procedure, the responsible surgeon, the number and type of modular devices 9050 and/or other surgical equipment that could potentially be used in the particular surgical procedure, the number and type of modular devices 9050 and/or other surgical equipment that are anticipated to be used in the particular surgical procedure, patient identification information, and the type of procedure being performed.

[1508] The intraoperative data can include, for example, modular device 9050 utilization (e.g., the number of firings by a surgical stapling instrument, the number of firings by an RF electrosurgical instrument or an ultrasonic instrument, or the number and types of stapler cartridges utilized), operating parameter data of the modular devices 9050 (e.g., the FTF curve for a surgical stapling instrument, a FTC curve for a surgical stapling instrument, the energy output of a generator, the internal pressure or pressure differential of a smoke evacuator), unexpected modular device 9050 utilization (i.e., the detection of the utilization of a modular device that is nonstandard for the procedure type), adjunctive therapies administered to the patient, and utilization of

equipment other than the modular devices **9050** (e.g., sealants to address leaks). The intraoperative data can also include, for example, detectable misuse of a modular device **9050** and detectable off-label use of a modular device **9050**.

[1509] The postoperative data can include, for example, a flag if the patient does not leave the operating theater and/or is sent for nonstandard postoperative care (e.g., a patient undergoing a routine bariatric procedure is sent to the ICU after the procedure), a postoperative patient evaluation relating to the surgical procedure (e.g., data relating to a spirometric performance after a thoracic surgery or data relating to a staple line leakage after bowel or bariatric procedures), data related to postoperative complications (e.g., transfusions or air leaks), or the patient's length of stay in the medical facility after the procedure. Because hospitals are increasingly being graded on readmission rates, complication rates, average length of stay, and other such surgical quality metrics, the postoperative data sources can be monitored by the analytics system **9100** either alone or in combination with surgical procedural outcome data (discussed below) to assess and institute updates to the controls programs of the surgical hubs **9000** and/or modular devices **9050**.

[1510] In some exemplifications, the intraoperative and/or postoperative data can further include data pertaining to the outcome of each surgical procedure or a step of the surgical procedure. The surgical procedural outcome data can include whether a particular procedure or a particular step of a procedure had a positive or negative outcome. In some exemplifications, the surgical procedural outcome data can include procedure step and/or time stamped images of modular device **9050** performance, a flag indicating whether a modular device **9050** functioned properly, notes from the medical facility staff, or a flag for poor, suboptimal, or unacceptable modular device **9050** performance. The surgical procedural outcome data can, for example, be directly detected by the modular devices **9050** and/or surgical hub **9000** (e.g., a medical imaging device can visualize or detect bleeding), determined or inferred by a situational awareness system of the surgical hub **9000** as described in U.S. patent application Ser. No. 15/940,654, or retrieved from a database **9054** (e.g., an EMR database) by the surgical hub **9000** or the analytics system **9100**. In some exemplifications, perioperative data including a flag indicating that a modular device **9050** failed or otherwise performed poorly during the course of a surgical procedure can be prioritized for communication to and/or analysis by the analytics system **9100**.

[1511] In one exemplification, the perioperative data can be assembled on a procedure-by-procedure basis and uploaded by the surgical hubs **9000** to the analytics system **9100** for analysis thereby. The perioperative data indicates the manner in which the surgical hubs **9000** were programmed to operate or were manually controlled in association with a surgical procedure (i.e., the operational behavior of the surgical hubs **9000**) because it indicates what actions the surgical hub **9000** took in response to various detected conditions, how the surgical hubs **9000** controlled the modular devices **9050**, and what inferences the situationally aware surgical hubs **9000** derived from the received data. The analytics system **9100** can be configured to analyze the various types and combinations of preoperative, intraoperative, and post-operative data to determine whether a control program update should be generated and then push

the update to the overall population or one or more sub-populations of surgical hubs **9000**, as necessary.

[1512] FIG. 195 illustrates a logic flow diagram of a process **9300** for updating the control program of a surgical hub **9000**, in accordance with at least one aspect of the present disclosure. During the following description of the process **9300**, reference should also be made to FIGS. 190 and 194. The process **9200** can be executed by, for example, one or more processors of the analytics servers **9070** of the analytics system **9100**. In one exemplification, the analytics system **9100** can be a cloud computing system. For economy, the following description of the process **9300** will be described as being executed by the analytics system **9100**; however, it should be understood that the analytics system **9100** includes processor(s) and/or control circuit(s) that are executing the describe steps of the process **9300**.

[1513] The analytics system **9100** executing the process **9300** receives **9302** perioperative data from the surgical hubs **9000** that are communicably connected to the analytics system **9100**. The perioperative data indicates the manner in which the surgical hubs **9000** are programmed to operate by their control programs or are controlled by the surgical staff during a surgical procedure. In some aspects, the perioperative data can include or being transmitted to the analytics system **9100** in association with surgical procedural outcome data. The surgical procedural outcome data can include data pertaining to an overall outcome of a surgical procedure (e.g., whether there was a complication during the surgical procedure) or data pertaining to a specific step within a surgical procedure (e.g., whether a particular staple line bled or leaked).

[1514] After an analytics system **9100** executing the process **9300** has received **9302** the perioperative data, the analytics system **9100** then analyzes **9304** the data to determine whether an update condition has been satisfied. In one exemplification, the update condition includes whether a threshold number or percentage of surgical hubs **9000** within the population exhibit a particular operational behavior. For example, the analytics system **9100** can determine that a control program update should be generated to automatically active an energy generator at a particular step in a type of surgical procedure when a majority of the surgical hubs **9000** are utilized to active the energy generator at that procedural step. In another exemplification, the update condition includes whether the rate of positive procedural outcomes (or lack of negative procedural outcomes) correlated to a particular operational behavior exceeds a threshold value (e.g., an average rate of positive procedural outcomes for a procedure step). For example, the analytics system **9100** can determine that a control program update should be generated to recommend that the energy generator be set at a particular energy level when the associated rate of hemostasis (i.e., lack of bleeding) at that energy level for the particular tissue type exceeds a threshold rate. In another exemplification, the update condition includes whether the rate of positive procedural outcomes (or lack of negative procedural outcomes) for a particular operational behavior is higher than the rate of positive procedural outcomes (or a lack of negative procedural outcomes) for related operational behaviors. In other words, if one subpopulation of surgical hubs **9000** exhibits a first operational behavior under a certain set of conditions and a second subpopulation of surgical hubs **9000** exhibits a second operational behavior under the same set of conditions, then the analytics system

**9100** can determine whether to update the control programs of the surgical hubs **9000** according to whether the first or second operational behavior is more highly correlated to a positive procedural outcome. In another exemplification, the analytics system **9100** analyzes **9304** the data to determine whether multiple update conditions have been satisfied.

[1515] If an update condition has not been satisfied, the process **9300** continues along the NO branch and the analytics system **9100** continues receiving **9302** and analyzing **9304** perioperative data from the surgical hubs **9000** to monitor for the occurrence of an update condition. If an update condition has been satisfied, the process **9300** continues along the YES branch and the analytics system **9100** proceeds to generate **9308** a control program update. The nature of the generated **9308** control program update corresponds to the particular operational behavior of the surgical hub **9000** that is identified by the analytics system **9100** as triggering the update condition. In other words, the control program update adds, removes, or otherwise alters functions performed by the surgical hub **9000** so that the surgical hub **9000** operates differently under the conditions that gave rise to the identified operational behavior. Furthermore, the type of control program update also depends upon whether the identified operational behavior results from manual control or control by the control program of the surgical hub **9000**. If the identified operational behavior results from manual control, the control program update can be configured to provide warnings, recommendations, or feedback to the users based upon the manner in which they are operating the surgical hub **9000**. For example, if the analytics system **9100** determines that taking a particular action or utilizing a particular instrument for a step in a surgical procedure improves outcomes, then the analytics system **9100** can generate **9308** a control program update that provides a prompt or warning to the surgical staff when the surgical hub **9000** determines that the designated step of the surgical procedure is occurring or will subsequently occur. Alternatively, the control program update can change one or more functions of the surgical hub **9000** from being manually controllable to being controlled by the control program of the surgical hub **9000**. For example, if the analytics system **9100** determines that a display of the visualization system **108** (FIG. 2) is set to a particular view by the surgical staff in a predominant number of surgical procedures at a particular step, the analytics system **9100** can generate a control program update that causes the surgical hub **9000** to automatically change the display to that view under those conditions. If the identified operational behavior results from the control program of the surgical hub **9000**, then the control program update can alter how the control program functions under the set of circumstances that cause the identified operational behavior. For example, if the analytics system **9100** determines that a particular energy level for an RF electrosurgical or ultrasonic instrument correlates to poor or negative outcomes under a certain set of conditions, then the analytics system **9100** can generate **9308** a control program update that causes the surgical hub **9000** to adjust the energy level of the connected instrument to a different value when the set of conditions is detected (e.g., when the surgical hub **9000** determines that an arthroscopic procedure is being performed).

[1516] The analytics system **9100** then transmits **9310** the control program update to the overall population of surgical hubs **9000** or the subpopulation(s) of surgical hubs **9000** that

are performing the operational behavior that is identified by the analytics system **9100** as triggering the update condition. In one exemplification, the surgical hubs **9000** are configured to download the control program updates from the analytics system **9100** each time an update is generated **9308** thereby. In one exemplification, the analytics system **9100** can thereafter continue the process **9300** of analyzing **9304** the data received **9302** from the surgical hubs **9000**, as described above.

[1517] FIG. 196 illustrates a representative implementation of the process **9300** depicted in FIG. 195. FIG. 196 illustrates a logic flow diagram of a process **9400** for updating the data analysis algorithm of a control program of a surgical hub **9000**, in accordance with at least one aspect of the present disclosure. As with the process **9300** depicted in FIG. 195, the process **9400** illustrated in FIG. 196 can, in one exemplification, be executed by the analytics system **9100**. In the following description of the process **9400**, reference should also be made to FIG. 194. In one exemplification of the adaptive surgical system **9060** depicted in FIG. 194, the first surgical hub subpopulation **9312** is utilizing a first data analysis algorithm and the second surgical hub subpopulation **9314** is utilizing a second data analysis algorithm. For example, the first surgical hub subpopulation **9312** can be utilizing a normal continuous probability distribution to analyze a particular dataset, whereas the second surgical hub subpopulation **9314** can be utilizing a bimodal distribution for analyzing the particular dataset. In this exemplification, the analytics system **9100** receives **9402**, **9404** the perioperative data from the first and second surgical hub subpopulations **9312**, **9314** corresponding to the respective data analysis algorithms. The analytics system **9100** then analyzes **9406** the perioperative datasets to determine whether one of the perioperative datasets satisfies one or more update conditions. The update conditions can include, for example, a particular analysis method being utilized by a threshold percentage (e.g., 75%) of the surgical hubs **9000** in the overall population and a particular analysis method being correlated to positive surgical procedural outcomes in a threshold percentage (e.g., 50%) of cases.

[1518] In this exemplification, the analytics system **9100** determines **9408** whether one of the data analysis algorithms utilized by the first and second surgical hub subpopulations **9312**, **9314** satisfies both of the update conditions. If the update conditions are not satisfied, then the process **9400** proceeds along the NO branch and the analytics system **9100** continues receiving **9402**, **9404** and analyzing **9406** perioperative data from the first and second surgical hub subpopulations **9312**, **9314**. If the update conditions are satisfied, the process **9400** proceeds along the YES branch and the analytics system **9100** generates **9412** a control program update according to which of the data analysis algorithms the analysis **9406** determined satisfied the update conditions. In this exemplification, the control program update would include causing the surgical hub **9000** to utilize the data analysis algorithm that satisfied the update conditions when performing the corresponding analysis type. The analytics system **9100** then transmits **9414** the generated **9412** control program update to the population of surgical hubs **9000**. In one exemplification, the control program update is transmitted **9414** to the entire population of surgical hubs **9000**. In another exemplification, the control program update is transmitted **9414** to the subpopulation of surgical hubs **9000** that did not utilize the data analysis algorithm that satisfied the

update conditions. In other words, if the analytics system **9100** analyzes **9406** the perioperative data and determines **9408** that the second (bimodal) data analysis method satisfies the update conditions, then the generated **9412** control program update is transmitted **9414** to the first subpopulation of surgical hubs **9000** in this exemplification. Furthermore, the control program update can either force the updated surgical hubs **9000** to utilize the second (bimodal) data analysis algorithm when analyzing the particular dataset or cause the updated surgical hubs **9000** to provide a warning or recommend to the user that the second (bimodal) data analysis algorithm be used under the given conditions (allowing the user to choose whether to follow the recommendation).

**[1519]** This technique improves the performance of the surgical hubs **9000** by updating their control programs generated from data aggregated across the entire network of surgical hubs **9000**. In effect, each surgical hub **9000** can be adjusted according to shared or learned knowledge across the surgical hub **9000** network. This technique also allows the analytics system **9100** to determine when unexpected devices (e.g., modular devices **9050**) are utilized during the course of a surgical procedure by providing the analytics system **9100** with knowledge of the devices being utilized in each type of surgical procedure across the entire surgical hub **9000** network.

#### Security and Authentication Trends and Reactive Measures

**[1520]** In a cloud-based medical system communicatively coupled to multiple communication and data gathering centers located in different geographical areas, security risks are ever present. The cloud-based medical system may aggregate data from the multiple communication and data gathering centers, where the data collected by any data gathering center may originate from one or more medical devices communicatively coupled to the data gathering center.

**[1521]** It may be possible to connect an unauthorized medical device to the data gathering center, such as a pirated device, a knock-off or counterfeit device, or a stolen device. These devices may contain viruses, may possess faulty calibration, lack the latest updated settings, or otherwise fail to pass safety inspections that can be harmful to a patient if used during surgery. Furthermore, the multiple data gathering centers may contain multiple points of entry, such as multiple USB or other input ports, or opportunities to enter user passwords, that if improperly accessed could represent security breaches that can reach the cloud-based medical system, other data gathering centers, and connected medical devices. The risk of devices being tampered with, or data being stolen or manipulated, can lead to severe consequences, particularly because the entire system is purposed for improving medical care.

**[1522]** A security system that reaches all facets of the cloud-based medical system may not be effective unless there is a centralized component that is configured to be made aware of all communication and data gathering centers, and all devices connected therein. If the security systems were merely localized to each data gathering center or at each point of entry, information from one point of entry may not be properly disseminated to other security points. Thus, if a breach occurs at one point, or if improper devices are used at one point, that information may not be properly disseminated to the other centers or devices. Therefore, a

centralized security system, or at least a system configured to communicate with all medical hubs that control access points, would be preferable to be made aware of all of the different issues that may occur and to communicate those issues to other ports as needed.

**[1523]** In some aspects, the cloud-based medical system includes a security and authentication system that is configured to monitor all communication and data gathering centers, such as a medical hub or tower located in an operating room, as well as any smart medical instruments communicatively coupled to those centers. The cloud-based security and authentication system, as part of the cloud-based medical system, may be configured to detect unauthorized or irregular access to any hub system or other protected data sets contained within the cloud. Because of the centralized nature of the cloud-based security system—in the sense that the cloud system is configured to communicate with every hub in the system—if there is any identified irregularity found at one hub, the security system is operable to improve security at all other hubs by communicating this information to the other hubs. For example, if surgical instruments with unauthorized serial numbers are used at a hub in one hospital, the cloud-based security system may learn of this at the local hub located in that hospital, and then communicate that information to all other hubs in the same hospital, as well as all hospitals in the surrounding region.

**[1524]** In some aspects, the cloud-based medical system may be configured to monitor surgical devices and approve or deny access for each surgical device for use with a surgical hub. Each surgical device may be registered with a hub, by performing an authentication protocol exchange with the hub. The cloud-based medical system may possess knowledge of all surgical devices and a status indicating whether the surgical device is acceptable, such as whether the device has been pirated, lacks a proper serial number, was faulty, possesses a virus, as so on. The cloud-based medical system may then be configured to prevent interaction with the surgical device, even if the surgical device is connected to the hub.

**[1525]** In this way, the cloud-based security system can provide the most comprehensive security for any particular hub or medical facility due to its ability to see problems located elsewhere.

**[1526]** FIG. 197 provides an illustration of example functionality by a cloud medical analytics system **10000** for providing improved security and authentication to multiple medical facilities that are interconnected, according to some aspects. Starting at block A reference **10002**, suspicious activity may be registered from one facility or region as a starting point. The suspicious activity may come in various forms. For example, a surgical device may be recorded at a hub as having a duplicate serial number, or a number that is not known to be within an acceptable range, or that the serial number may already be registered at a different location. In some aspects, surgical devices may possess additional authentication mechanisms, such as a type of electronic or digital handshake exchange between the surgical device and the surgical hub when they are connected. Each device may be programmed with a digital signature and/or knowledge of how to perform an authentication process. The firmware of the surgical device may need to be properly programmed to know how to perform during this exchange. The authentication handshake may periodically change, and may be

specified by the cloud on a periodic basis. Any of these may fail during interconnection of the device with a medical hub, triggering an alert with the medical hub and the cloud system **10000**.

[1527] In some aspects, the cloud system **10000** may review the information supplied by the medical device that triggered the suspicious activity, and if the information is unequivocally fraudulent or faulty, an alert and a rejection of the device can occur, such that the medical device will be prevented from operating with the medical hub and/or other medical hubs in the same facility. While the cloud system **10000** may be configured to prevent singularities, the cloud system **10000** may also be capable of utilizing its vast array of knowledge to develop additional security measures that a single hub as an entry port would be unable to perform on its own. An example is described further below.

[1528] At block B reference **10004**, the activity at the local medical hub may be transmitted to the cloud for authentication by at least comparing the surgical device to all known devices within the cloud network. In this scenario, the surgical device may register as being suspicious or having some suspicious activity or property. The cloud may be configured to then undergo a feedback loop of exchange with the local hub or facility from which the suspicious device originated. The cloud may determine to request additional data from that facility. In addition, the medical facility, via one or more surgical hubs, may request authentication or interrogation data about one or more surgical devices from the cloud. In this example, a medical hub in a facility in Texas requests a communication exchange with the cloud system **10000** for more data to determine if the suspicious activity at one of its local hubs is truly problematic.

[1529] At block C reference **10006**, the cloud authentication and security system may then be configured to perform additional data analysis to determine the veracity of any threat and larger context of the nature of this suspicious activity. In this example, the cloud-based security system has performed analysis and brings to light at least two pieces of evidence of a security threat, which is expressed visually in the chart of block C. First, upon comparing the number of data requests and medical interrogations across multiple medical facilities, it is determined that the current requesting facility in Texas has an inordinate number of data requests or medical interrogations compared to all other facilities. The cloud may be configured to flag this as one security issue that needs to be addressed. Second, in comparison to the number of data requests, the number of suspicious data points or findings is also inordinately high at the Texas facility. One or both of these realizations may prompt the cloud security system to enact different security changes at the Texas facility in particular.

[1530] Thus, at block D reference **10008**, in response to the identified anomalous behavior of the facilities in Texas as a whole, the cloud security system may request additional data related to Texas to better understand the nature of the practices and potential threats. For example, additional data regarding purchasing practices, vendors, the type of surgical instruments being used, the type of surgical procedures performed in comparison to other facilities, and so forth, may be obtained from one or more surgical hubs at the Texas facility, or may be accessed in data already stored in the cloud system **10000**. The cloud security system may be configured to look for additional anomalies and patterns that

may help determine how to change security procedures specific to the Texas facility, or the facilities in the Texas region generally.

[1531] At block E reference **10010**, once the additional information has been gathered and analyzed, the cloud security system may initiate a changed security protocol for the Texas facility in particular that triggered this analysis from block A, as well as any new security procedures for any surgical devices that indicate a unique or above average threat. For example, it may be determined that a particular type of surgical devices, such as devices originating from a particular manufacturing facility or having a particular set of unique identification numbers, may be faulty, pirated, or have some other kind of security risk. The cloud system **10000** may have analyzed the suspicious data points originating from the Texas region, determined if there were any commonalities or patterns, and issued a change in security protocol based on these identified patterns. These devices may then be locked out from use at all surgical hubs, even if they are not connected to any surgical hub at the present time. Other example changes regarding security include modifying the types of data gathered to learn more about the types of threats or how widespread the threats are. For example, the suspicious activity in Texas may exhibit a certain pattern or authentication signature of attempting to login in with the system, and so this pattern may be placed on an alert to other facilities in Texas and/or to other facilities to pay special attention to. In some cases, the pattern of suspicious activity may be correlated with another indicator, such as a brand or manufacturer, or a series of serial numbers. The cloud system may send out alerts to those facilities known to associate with these correlated indicators, such as all facilities that utilize medical devices with the same manufacturer.

[1532] In addition, an augmented authentication procedure may be enacted at the localized Texas region. The cloud-security system may opt to perform additional authentication protocols for all devices originating out of the Texas facility, for example. These additional protocols may not be present or required at other facilities, since there is considered a lower level of security risk based on the lack of suspicious activity.

[1533] In some aspects, as alluded to previously, the cloud-based security system may also be configured to protect against unwanted intrusions, either to any hub or to the cloud system itself. This means that the suspect medical device may be unable to access any data from any medical hub, and may also be prevented from operating if it is connected to a medical hub. In a medical system utilizing the cloud system and multiple medical hubs, the common protocol may require that only medical devices connected to a medical hub are authorized to operate on a patient, and therefore the medical hub will have the capability of preventing a device from activating. The limitation of any faulty or fraudulent surgical device may be designed to protect a patient during a surgical procedure, and it can also be used to protect any surgical hub and the cloud itself. The same lockout procedure may be designed to stop both scenarios from occurring.

[1534] In some aspects, the surgical hub may be configured to transmit data to the cloud security system that better characterizes the nature of the security flaws or intrusions. For example, the cloud security system may be configured to store in memory the number of intrusion attempts, the

source of the intrusion attempt (e.g., from which surgical hub or even what port or connection via the surgical hub), and what method for attempted intrusion there is, if any (e.g., virus attack, authentication spoofing, etc.).

[1535] In some aspects, the cloud security system may also determine what types of behaviors by a surgical device or other functions by a surgical hub are irregular, compared to a global average or just by each institution. The cloud security system may better identify what practices seem irregular in this way. The data logs of any surgical hub, or across an entire facility, may be recorded and securely stored in the cloud system. The cloud security system may then analyze the attempted access requests and actions to determine trends, similarities and differences across regions or institutions. The cloud security system may then report any irregularities to the institution and flag any identified irregularities for internal investigation into updates to protect against future breaches. Of note, a local hub or local facility with multiple hubs may not realize if any of their authentication behaviors are irregular, unless they are compared to a broader average or comparison of other facilities. The cloud system may be configured to identify these patterns, because it has access to authentication data and procedures from these multiple facilities.

[1536] In some aspects, the cloud security system may be configured to analyze any current hub control program versions and when it was updated. The cloud security system may verify all updates are correct, and determine where their origins are. This may be an additional check to ensure that the software and firmware systems of the surgical devices are proper and have not been tampered with.

[1537] In some aspects, the cloud security system may also determine larger threats by analyzing multiple facilities at once. The system may determine, after aggregating data from multiple locations, any trends or patterns of suspicious activity across a wider region. The security system may then change security parameters across multiple facilities immediately or in near real time. This may be useful to quickly react to simultaneous attacks, and may make it even easier to solve simultaneous attacks by gathering data from the multiple attacks at once to better increase the chances and speed of finding a pattern to the attacks. Having the cloud system helps confirm whether attacks or suspicious activity occurs in isolation or is part of a grander scheme.

#### Data Handling and Prioritization

[1538] Aspects of the present disclosure are presented for a cloud computing system (computer-implemented interactive surgical system as described above) for providing data handling, sorting, and prioritization, which may be applied to critical data generated during various medical operations. The cloud computing system constitutes a cloud-based analytics system, communicatively coupled to a plurality of surgical hubs 7006 and smart medical instruments such as surgical instruments 7012. Typically, a healthcare facility, such as a hospital or medical clinic, does not necessarily immediately recognize the criticality of data as it is generated. For example, if a medical instrument used during a perioperative period experiences a failure, the response of medical care facility personnel such as nurses and doctors may be directed towards diagnosis of any medical complications, emergency medical assistance, and patient safety generally. In this situation, the criticality of the data might not be analyzed in a time sensitive manner, or at all.

Accordingly, the healthcare facility does not necessarily timely respond to or even recognize critical data as such data is generated. Additionally, a particular healthcare facility can lack knowledge of the management of critical data from other similarly situated facilities, either in its region, according to a similar size, and/or according to similar practices or patients, and the like. The cloud-based analytics system may be specifically designed to address this issue of critical data and particularly the timing of data handling that is performed based on the criticality of data within the context of healthcare facility operations. The cloud-based analytics system may quickly and efficiently identify critical data based on specific criteria. In some situations, aggregate data is determined to be critical after the individual non-critical data comprising the aggregated data are aggregated. As used herein, handling critical data (which could be aggregated) may refer to data sorting, prioritizing, and other data handling based on specific criteria or thresholds.

[1539] To help facilitate timely and improved data sorting, handling, and prioritization, it would be desirable if a common source connected to multiple healthcare facilities could sort, handle, and prioritize critical data from these medical facilities in a holistic manner. In this way, insights could be generated by the common source based on using this aggregated data from the multiple healthcare facilities. In various aspects, the cloud-based analytics system comprises the cloud 7004 that is communicatively coupled to knowledge centers in a medical facility, such as one or more surgical hubs 7006, and is configured to sort, handle, and prioritize medical data from multiple healthcare facilities. In particular, the cloud-based system can identify critical data and respond to such critical data based on the extent of the associated criticality. For example, the cloud-based system could prioritize a response as requiring urgent action based on the critical data indicating a serious perioperative surgical instrument 7012 failure, such as one that requires intensive care unit (ICU) postoperative treatment. The data handling, sorting, and prioritization described herein may be performed by the processors 7008 of the central servers 7013 of the cloud 7004 by, for example, executing one or more data analytics modules 7034.

[1540] Critical data can be determined to be critical based on factors such as severity, unexpectedness, suspiciousness, or security. Other criticality criteria can also be specifically selected such as by a healthcare facility. Criticality can also be indicated by flagging a surgical instrument 7012, which in turn can be based on predetermined screening criteria, which could be the same or different as the factors described above. For example, a surgical instrument 7012 can be flagged based on its usage being correlated with severe post surgical operation complications. Flagging could also be used to trigger the prioritized data handling of the cloud-based analytics system. In connection with a determination of criticality or flagging a surgical instrument 7012, the cloud 7004 can transmit a push message or request to one or more surgical hubs 7006 for additional data associated with the use of the surgical instrument 7012. The additional data could be used for aggregating data associated with the surgical instrument 7012. For example, after receiving the additional data, the cloud 7004 may determine there is a flaw in the surgical instrument 7012 (e.g., malfunctioning generator in an energy surgical instrument) that is common to other corresponding surgical instruments 7012 in a particular healthcare facility. Accordingly, the cloud 7004 could



determine that all such flawed surgical instruments **7012** should be recalled. These flawed surgical instruments **7012** might share a common identification number or quality or a common aspect of a unique identifier, such as a serial number family identifier.

[1541] In general, the cloud-based analytics system may be capable of aggregating, sorting, handling, and prioritizing data in a timely and systematic manner that a single healthcare facility would not be able to accomplish on its own. The cloud-based analytics system further can enable timely response to the aggregated, sorted, and prioritized data by obviating the need for multiple facilities to coordinate analysis of the particular medical data generated during medical operations at each particular facility. In this way, the cloud-based system can aggregate data to determine critical data or flagging for enabling appropriate responses across the entire network of surgical hubs **7006** and instruments **7012**. Specifically, appropriate responses include sorting, handling, and prioritization by the cloud **7004** according to a priority status of the critical data, which can enable timely and consistent responses to aggregated critical data (or critical aggregated data) across the entire network. Criticality of the data may be defined universally and consistently across all surgical hub **7006** and instruments **7012**. Furthermore, the cloud-based analytics system may be able to verify the authenticity of data from the plurality of medical facilities before such data is assigned a priority status or stored in the aggregated medical data databases. As with the categorization of critical data, data verification can also be implemented in a universal and consistent manner across the system which a single facility may not be able to achieve individually.

[1542] FIG. 198 is a flow diagram of the computer-implemented interactive surgical system programmed to use screening criteria to determine critical data and to push requests to a surgical hub to obtain additional data, according to one aspect of the present disclosure. In one aspect, once a surgical hub **7006** receives device data **11002** from a surgical instrument **7012** data may be flagged and/or determined to be critical based on predetermined screening criteria. As shown in FIG. 198, the hub **7006** applies **11004** the screening criteria to flag devices and to identify critical data. The screening criteria include severity, unexpectedness, suspiciousness, and security. Severity can refer to the severity of any adverse medical consequences resulting from an operation performed using the surgical instrument **7012**. Severity could be assessed using a severity threshold for surgical instrument **7012** failures. For example, the severity threshold could be a temporal or loss rate threshold of bleeding such as over 1.0 milliliters per minute (mL/min). Other suitable severity thresholds could be used. Unexpectedness can refer to a medical parameter of a deviation that exceeds a threshold such as an amount of standard deviation from the mean medical parameter value such as a determined tissue compression parameter significantly exceeding the expected mean value at a time during an operation.

[1543] Suspiciousness can refer to data that appears to have been improperly manipulated or tampered with. For example, the total therapeutic energy applied to tissue value indicated by the data may be impossible given a total amount energy applied via the generator of the surgical instrument **7012**. In this situation, the impossibility of the data suggests improper manipulation or tampering. Similarly, security can refer to improperly secured data, such as data including a

force to close parameter that was inadvertently deleted. The screening criteria also may be specified by a particular surgical hub **7006** or by the cloud **7004**. The screening criteria can also incorporate specific thresholds, which can be used for prioritization, for example. In one example, multiple severity thresholds can be implemented such that the extent of perioperative surgical instrument **7012** failures can be sorted into multiple categories according to the multiple severity thresholds. In particular, the multiple severity thresholds could be based on the number of misaligned staples from a stapling surgical instrument **7012** to reflect an extent of the severity of misalignment. By using the cloud-based analytic system, the cloud may systemically identify critical data and flag surgical instruments **7012** for providing a timely and appropriate response which an individual healthcare facility could not achieve on its own. This timely response by the cloud **7004** can be especially advantageous for severe post surgical operation complications.

[1544] Determining critical data and flagging the surgical instrument **7012** by the hub **7006** may include determining a location to store data. Data may be routed or stored based on whether the data is critical and whether the corresponding surgical instrument **7012** is flagged. For example, binary criteria can be used to sort data into two storage locations, namely, a memory of a surgical hub **7006** or the memory **7010** of the cloud **7004**. Surgical instruments **7012** generate this medical data and transmit such data, which is denoted as device data **11002** in FIG. 198, to their corresponding surgical hub devices **7006**. FIG. 198 illustrates an example of this binary sorting process. Specifically, in one aspect, the data routing can be determined based on severity screening criteria as shown at the severity decision steps **11006**, **11008**. At step **11006**, the hub **7006** determines **11006** whether the surgical instrument **7012** that provided the device data **11002** has experienced a failure or malfunction during operation at the perioperative stage and whether this failure is considered severe. The severity thresholds discussed above or other suitable means could be used to determine whether the failure is severe. For example, severe failure may be determined based on whether undesirable patient bleeding occurred during use or firing of the surgical instrument. If the determination at step **11006** is yes, the corresponding data (i.e., critical data) of the surgical instrument **7012** is transmitted **11012** by the hub **7006** to the cloud **7004**. Conversely, if the determination at step **11006** is no, the flow diagram may proceed to step **11008**.

[1545] If the determination at step **11006** is no, then the flow diagram proceeds to step **11008** in FIG. 198, where the surgical hub **7006** determines whether the patient transitioned to non-standard post-operation care (i.e. the ICU) after the operation was performed with the specific surgical instrument **7012**. However, even if the determination at step **11006** is no, the inquiry at step **11008** may still be performed. If the determination at step **11008** is yes, then the critical device data **11002** is transmitted to the cloud **7004**. For example, the determination at step **11008** is yes if a patient transitioned into the ICU from the operating room subsequent to a routine bariatric surgical procedure. Upon transfer of a patient into the ICU, the surgical hub **7006** may receive a timely signal from the surgical instrument **7012** used to perform the bariatric procedure indicating that the patient has experienced complications necessitating entry into the ICU. Since this signal indicates the step **11008** determination is yes, corresponding device data **11002** is

sent **11012** to the cloud **7004**. Additionally, the specific surgical instrument **7012** may be flagged by the cloud **7004** for a prompt specific response by the cloud **7004**, such as designating the surgical instrument **7012** with a prioritization of requiring urgent action. If the determination at step **11008** is no, a signal can be transmitted from the surgical instrument **7012** to the surgical hub **7006** indicating that the procedure was successful. In this scenario, the device data **11002** can be stored **11010** locally in a memory device of the surgical hub **7006**.

[1546] Additionally or alternatively, the specific surgical instrument **7012** may also be flagged by the hub **7006** or the cloud **7004** to trigger data handling by the cloud **7004**, which can comprise an internal response of the cloud **7004**. When the surgical instrument **7012** is flagged or the device data **11002** is determined to be critical, the triggered response may be the cloud **7004** transmitting a signal comprising a request for additional data regarding the surgical instrument **7012**. Additional data may pertain to the critical device data **11002**. The cloud **7004** can also request additional data even if the specific surgical instrument **7012** is not flagged, such as if the device data **11002** is determined to be critical without the surgical instrument **7012** being flagged. Flagging could also indicate an alarm or alert associated with the surgical instrument **7012**. In general, the hub **7006** is configured to execute determination logic for determining whether the device data **11002** should be sent to the cloud **7004**. The determination logic can be considered screening criteria for determining criticality or flagging surgical instruments **7012**. Besides the severity thresholds used at steps decision steps **11006**, **11008**, the data routing can be based on frequency thresholds (e.g., the use of a surgical instrument **7012** exceeds a usage quantity threshold such as a number of times an energy generator is used), data size thresholds, or other suitable thresholds such as the other screening criteria discussed above. Flagging may also result in storing a unique identifier of the specific surgical instrument in a database of the cloud-based system.

[1547] A triggered request **11014** for additional data by the cloud **7004** to the hub **7006** may be made based on a set of inquiries as shown in FIG. **198**. This triggered request **11014** may be a push request sent by the central servers **7013** of the cloud **7004**. In particular, the processors **7008** can execute the data collection and aggregation data analytic module **7022** to implement this trigger condition functionality. This push request may comprise an update request sent by the cloud **7004** to the hub **7006** to indefinitely collect new data associated with the device data **11002**. That is, the hub **7006** may collect additional data until the cloud **7004** transmits another message rescinding the update request. The push request could also be a conditional update request. Specifically, the push request could comprise initiating a prompt for the hub **7006** to send additional information only if certain conditions or events occur. For example, one condition might be if the sealing temperature used by the surgical instrument **7012** to treat tissue exceeds a predetermined threshold. The push request could also have a time bounding component. In other words, the push request could cause the surgical hub **7006** to obtain additional data for a specific predetermined time period, such as three months. The time period could be based on an estimated remaining useful life of the surgical instrument **7012**, for example. As discussed above, the request **11014** for additional data may occur after

the specific surgical instrument **7012** is flagged, which may be due to an affirmative determination at steps **11006**, **11008** described above.

[1548] As shown in FIG. **198**, the triggered request **11014** for additional data may include four inquiries that can be considered trigger conditions for additional information. At the first inquiry, the hub **7006** determines **11016** whether the device data **11002** represents an outlier with no known cause. For example, application of therapeutic energy to tissue during a surgical procedure by the surgical instrument **7012** may cause patient bleeding even though surgical parameters appear to be within a normal range (e.g., temperature and pressure values are within expected range). In this situation, the critical device data **11002** indicates an irregularity without a known reason. The outlier determination **11016** can be made based on comparison of the device data **11002** to an expected value or based on a suitable statistical process control methodology. For example, an actual value of the device data **11002** may be determined to be an outlier based on a comparison of the actual value to a mean expected (i.e., average) value. Calculating that the comparison is beyond a certain threshold can also indicate an outlier. For example, a statistical process control chart could be used to monitor and indicate that the difference between the actual and expected value is a number of standard deviations beyond a threshold (e.g., 3 standard deviations). If the device data **11002** is determined to be an outlier without a known reason, the request **11014** is triggered by the cloud **7004** to the hub **7006**. In response, the hub **7006** timely transmits **11024** additional information to the cloud **7004**, which may provide different, supporting, or additional information to diagnose the reason for the outlier. Other insights into the outlier may also be derived in this way. For example, the cloud **7004** may receive additional surgical procedure parameter information such as the typical clamping force used by other surgical instruments **7012** at the same point in the surgical procedure when the patient bleeding occurred. The expected value may be determined based on aggregated data stored in the aggregated medical data database **7012**, such as by averaging the outcomes or performance of groups of similarly situated surgical instruments **7012**. If at step **11016**, the data is not determined to be an outlier, the flow diagram proceeds to step **11018**.

[1549] The second inquiry is another example of a trigger condition. At step **11018**, the hub **7006** determines **11018** whether device data **11002** involves data that can be classified as suspicious, which can be implemented by the authorization and security module **7024**. For example, suspicious data may include situations in which an unauthorized manipulation is detected. These include situations where the data appears significantly different than expected so as to suggest unauthorized tampering, data or serial numbers appear to be modified, security of surgical instruments **7012** or corresponding hub **7006** appears to be comprised. Significantly different data can refer to, for example, an unexpected overall surgical outcome such as a successful surgical procedure occurring despite a surgical instrument **7012** time of usage being significantly lower than expected or a particular unexpected surgical parameter such as a power level applied to the tissue significantly exceeding what would be expected for the tissue (e.g., calculated based on a tissue impedance property). Significant data discrepancies could indicate data or serial number modification. In one example, a stapling surgical instrument **7012** may

generate a separate unique staple pattern in a surgical operation which may be used to track or verify whether the serial number of that stapling surgical instrument **7012** is subsequently modified. Furthermore, data or serial number modification such as tampering may be detected via other associated information of a surgical instrument **7012** that can be independently verified with the aggregated medical data databases **7011** or some other suitable data modification detection technique.

[1550] Moreover, compromised security, such as unauthorized or irregular access to any surgical hub **7006** or other protected data sets stored within the cloud **7004** can be detected by a cloud-based security and authentication system incorporating the authorization and security module **7024**. The security and authentication system can be a suitable cloud based intrusion detection system (IDS) for detecting compromised security or integrity. The cloud IDS system can analyze the traffic (i.e. network packets) of the cloud computing network **7001** or collect information (e.g., system logs or audit trails) at various surgical hub **7006** for detecting security breaches. Compromised security detection techniques include comparison of collected information against a predefined set of rules corresponding to a known attack which is stored in the cloud **7004** and anomaly based detection. The cloud **7004** can monitor data from a series of surgical operations to determine whether outliers or data variations significantly reduce without an apparent reason, such as a reduction without a corresponding change in parameters of used surgical instruments **7012** or a change in surgical technique. Additionally, suspiciousness can be measured by a predetermined suspiciousness or unexpectedness threshold, unauthorized modification of device data **11002**, unsecure communication of data, or placement of the surgical instrument **7012** on a watch list (as described in further detail below). The suspiciousness or unexpectedness threshold can refer to a deviation (e.g., measured in standard deviations) that exceeds surgical instrument **7012** design specifications. Unauthorized data communication or modification can be determined by the authorization and security module **7024** when the data encryption of the cloud **7004** is violated or bypassed. In sum, if the hub **7006** determines **11018** the data is suspicious for any of the reasons described above, the request **11014** for additional data may be triggered. In response, the hub **7006** timely transmits **11024** additional information to the cloud **7004**, which may provide different, supporting, or additional information to better characterize the suspiciousness. If at step **11018**, the answer to the second inquiry is no, the flow diagram proceeds to step **11020**.

[1551] The third and fourth inquiries depict additional trigger conditions. At step **11020**, the hub **7006** may determine that device data **11002** indicates a unique identifier of the surgical instrument **7012** that matches an identifier maintained on a watchlist (e.g., "black list" of prohibited devices). As described above, the "black list" is a watch list that can be maintained as a set of database records comprising identifiers corresponding to prohibited surgical hubs **7006**, surgical instruments **7012**, and other medical devices. The black list can be implemented by the authorization and security module **7024**. Moreover, surgical instruments **7012** on the black list may be prevented from fully functioning or restricted from access with surgical hubs **7006**. For example, an energy surgical instrument **7012** may be prevented from functioning (i.e. an operational lockout) via the cloud **7004**

or surgical hub **7006** transmitting a signal to the hub **7006** or surgical instrument **7012** to prevent the generator from applying power to the energy surgical instrument **7012**. This operational lockout can generally be implemented in response to an irregularity indicated by the critical device data **11002**. Surgical instruments can be included on the black list for a variety of reasons such as the authorization and security module **7012** determining the presence of counterfeit surgical instruments **7012** using internal authentication codes, unauthorized reselling of surgical instruments **7012** or related products from one region to another, deviation in performance of surgical instruments **7012** that is nonetheless within design specifications, and reuse of surgical instruments **7012** or related products that are designed for single patient use. For example, internal authentication codes may be unique identifiers maintained by the cloud **7004** in the memory devices **7010**. Other unauthorized usage could also result in placement on the black list.

[1552] The use of counterfeit authentication codes may be a security breach that is detectable by the cloud IDS system. Reselling of surgical instruments **7012** into other regions could be detected via region specific indicators of resold surgical instrument **7012** or surgical hubs **7006**, for example. The region specific indicator could be encrypted using a suitable encryption technique. In this way, the cloud **7004** may detect when the region specific indicators of a resold surgical instrument **7012** do not match the corresponding region of intended use. Reuse of a single use surgical instrument **7012** can be monitored by detecting tampering with a lockout mechanism (e.g., a stapler cartridge lockout mechanism of a stapling surgical instrument), programming a microprocessor of the single use surgical instrument **7012** to transmit a warning signal to the corresponding surgical hub **7006** when more than one use occurs, or another suitable detection technique. Performance deviation could be monitored using statistical process control methods as described above. The design specifications of particular surgical instruments **7012** may be considered the control limits of a statistical process control methodology. In one example, when detected by the cloud **7004**, a significant trend toward one of the lower or upper control limits constitutes a sufficient deviation that results in the cloud **7004** adding the corresponding surgical instrument to the black list. As discussed above, a deviation that exceeds design specifications may result determining **11018** the device data **11002** is suspicious. Surgical instruments **7012** may be added to or removed from the black list by the cloud **7004** based on analysis of the requested additional data. In sum, if the hub **7006** determines **11020** the surgical instrument **7012** corresponding to the device data **11002** is on the watchlist, the request **11014** for additional data may be triggered. In response, the hub **7006** timely transmits **11024** additional information to the cloud **7004**, which may provide different, supporting, or additional information. If at step **11020**, the answer to the second inquiry is no, the flow diagram proceeds to step **11022**.

[1553] The trigger condition at step **11022** comprises the hub **7006** determining whether the device data **11002** indicates the surgical instrument **7012** has malfunctioned. In one aspect, a surgical instrument **7012** malfunction results in an automated product inquiry through the corresponding surgical hub **7006**. The hub **7006** sending **11024** additional data to the cloud **7004** may comprise all pertinent data of the

surgical instrument **7012** being immediately transmitted to the cloud through the surgical hub **7006**, which may result in central server **7013** processors **7008** of the cloud **7004** executing an automated product inquiry algorithm. However, such an algorithm may not be immediately executed or at all if the malfunction is not significant. The cloud **7004** may be configured to record this set of pertinent data for all surgical instruments **7012** for contingent use when such automated product inquiries are instituted. The automated product inquiry algorithm comprises the cloud **7004** searching for previous incidents that are related to the malfunction. The cloud **7004** may populate a group of records in the aggregated medical data databases **7011** with any incidents or activity related to the malfunction. Subsequently, a corrective and preventive action (CAPA) portion of the algorithm may be instituted for reducing or eliminating such malfunctions or non-conformities. CAPA and the automated product inquiry algorithm are one example of a possible internal response **11102** of the cloud **7004** of the cloud-based analytics system.

[1554] CAPA involves investigating, recording and analyzing the cause of a malfunction or non-conformity. To implement CAPA, the cloud **7004** may analyze the populated related records in the aggregated medical data databases **7011**, which may include aggregated data fields such as surgical instrument **7012** manufacture dates, times of use, initial parameters, final state/parameters, and surgical instrument **7012** numbers of uses. Thus, both individual and aggregated data may be used. In other words, the cloud **7004** may analyze both individual data corresponding to the malfunctioning surgical instrument **7012** as well as aggregated data, collected from all related surgical instruments **7012** to the malfunctioning surgical instrument **7012**, for example. Initial and final parameters may be, for example, an initial and final frequency of an applied RF signal of the surgical instrument. CAPA can also involve analysis of the previous time period from when the malfunction occurred or was detected. Such a time period can be, for example, one to two minutes. Based on this CAPA analysis, the cloud **7004** may diagnose the root cause of the malfunction and recommend or execute any suitable corrective action (e.g., readjusting miscalibrated parameters). The automated product inquiry algorithm can also involve a longer follow up of patient outcomes for patients treated with the specific surgical instrument **7012**.

[1555] For example, the cloud **7004** may determine a priority status of watch list for the surgical instrument **7012** so that the surgical instrument **7012** may be monitored for a period of time after the malfunction is detected and addressed. Moreover, the malfunction may cause the cloud **7004** to expand a list of medical items to be tracked (e.g., the integrity of tissue seals made during surgery). This list of items to be tracked may be performed in conjunction with the patient outcome monitoring by the patient outcome analysis module **7028**. The cloud **7004** may also respond to an irregularity indicated by the malfunction by monitoring patient outcomes corresponding to the irregularity. For example, the cloud **7004** can monitor whether the irregularity corresponds to unsuccessful surgical operations for a predetermined amount of time such as 30 days. Any corrective action also can be assessed by the cloud **7004**. Other data fields can also be monitored in addition to the fields discussed above. In this way, the cloud may timely diagnose and respond to surgical instrument **7012** malfunctions using

individual and aggregate data in a manner that an individual healthcare facility could not achieve.

[1556] In one aspect, if the answer to any of steps **11016**, **11018**, **11020**, **11022** (i.e. trigger conditions) is affirmative (i.e. the trigger condition is activated), then additional data associated or pertinent to the device data **11002** is sent to the cloud **7004**, as can be seen in FIG. **198**. This additional data may be handled by the data sorting and prioritization module **7032** while the patient outcome analysis module **7028** may analyze the data, for example. In contrast, if the answer to all of steps **11016**, **11018**, **11020**, **11022** is negative, then the respective data is stored **11026** within the corresponding surgical hub **7006**. Thus, when the answer at step **11022** is no, the device data **11002** may be stored locally within the hub **7006** and no additional data is requested of the hub **7006**. Alternatively, the device data may be sent to the cloud **7006** for storage within the memory devices **7010**, for example, without any triggered requests **11014** by the cloud **7004** for additional data. Steps **11016**, **11018**, **11020**, **11022** could also be used for identifying critical data or flagging the surgical instrument (if the specific surgical device has not already been flagged based on steps **11006**, **11008**) as part of the screening criteria applied at step **11004**. Other trigger conditions aside from steps **11016**, **11018**, **11020**, **11022** are also possible for triggering the request **11014** for additional data. The request can be sent to all surgical hubs **7006** or a subset thereof. The subset can be geographically specific such that, for example, if surgical hub **7006** used in healthcare facilities located in Illinois and Iowa have malfunctioned in a similar manner, only surgical hub **7006** corresponding to healthcare facilities in the Midwestern United States are requested **11014** for additional information. The requested additional data can be different or supporting data concerning the particular use of surgical instruments **7012** so that the cloud **7004** may gain additional insight into the source of the irregularity, as represented by steps **11016**, **11018**, **11020**, **11022**. For example, if malfunctioning surgical instruments **7012** are causing undesirable patient bleeding, the cloud **7004** may request timing information regarding this bleeding for help in potentially diagnosing why the malfunction is causing the bleeding.

[1557] The criticality of data can be identified based on the screening criteria as described above, or by any other suitable data analysis technique. In one aspect, as shown in FIG. **199**, when the critical data is determined, an internal analytic response **11102** of the cloud **7004** may commence. The internal analytic response **11102** can advantageously be made in a timely manner such as in real time or near real time. As discussed above, the criticality of data can be identified based on the severity of an event, the unexpected nature of the data, the suspiciousness of the data, or some other screening criteria (e.g., an internal business flag). The determination of critical data can involve a request generated by a surgical hub **7006** based on the surgical hub **7006** detecting an irregularity or failure of a corresponding surgical instrument **7012** or of a component of the surgical hub **7006** itself. The request by the surgical hub **7006** may comprise a request for a particular prioritization or special treatment of critical data by the cloud **7004**. In various aspects, the cloud internal analytic response **11102** could be to escalate an alarm or response based on the frequency of the event associated with the critical device data **11002**, route the device data **11002** to different locations within the cloud computing system, or exclude the device data **11002**

from the aggregated medical data databases 7011. In addition, the cloud 7004 could also automatically alter a parameter of a malfunctioning surgical instrument 7012 so that modifications for addressing the malfunction can be implemented in real time or near real time. In this manner, even malfunctions that are not readily detected by a clinician in a healthcare facility, for example, may still be advantageously addressed in a timely manner by the cloud 7004.

[1558] FIG. 199 is a flow diagram of an aspect of responding to critical data by the computer-implemented interactive surgical system, according to one aspect of the present disclosure. In particular, the internal analytic response 11102 by the cloud 7004 can include handling critical data which includes determining a priority status to determine a time component or prioritization of the response. The response 11102 itself may be based on an operational characteristic indicated by the critical data, such as the characteristics described above in connection with the screening criteria or the trigger conditions of FIG. 198. The internal response 11102 may be implemented by the data sorting and prioritization module 7032 as well as the data collection and aggregation module 7022. As shown in FIG. 199, in the prioritization branch of the flow diagram (labeled as Q1 in FIG. 199) the cloud may incorporate the binary decision of whether to exclude the critical data from the aggregated medical data databases 7011 with a priority escalation decision framework. At step 11104 of FIG. 199, the cloud 7004 determines whether the critical data should be excluded from the aggregated medical data databases 7011. The exclusion determination may be considered a threshold determination.

[1559] It can be desirable to exclude critical data from the aggregated medical data databases 7011 for verification purposes. For example, critical data that is flagged or designated for special routing may be placed on a hold list maintained by the cloud 7004. The hold list is maintained at a separate storage location in the memory 7010 relative to the aggregated medical data databases 7011 within the cloud 7004, such as the caches 7018. The excluded critical data could also be stored in a more permanent storage location in the memory 7010. Accordingly, if the answer to step 11104 is yes, the cloud 7004 stores 11118 the critical data in the hold list. The cloud 7004 may then validate or verify that the critical device data 11002 is accurate. For example, the cloud 7004 may analyze whether the device data 11002 is logical in light of a corresponding patient outcome or analyze additional associated data of the device data 11002. Upon proper verification, the device data 11002 may also be stored within the aggregated medical data databases 7011. But if the device data 11002 is not verified, the cloud 7004 may not include the unverified device data 11002 in the priority escalation decision framework. That is, before verification, the device data 11002 may not be assigned a priority status according to the priority status classification 11106 for the internal cloud response 11102.

[1560] However, if the device data 11002 is verified, the flow diagram may proceed to the priority status classification 11106. Accordingly, if the answer to the exclusion determination at step 11104 is no, the device data 11002 is prioritized according to the priority escalation decision framework, which can define a predetermined escalation method for handling critical data. As shown in FIG. 199, a predetermined escalation prioritization system 11106 (i.e., priority escalation decision framework) can comprise four

categories, including watch list, automated response, notification, and urgent action required. This predetermined escalation prioritization system 11106 can be considered a form of triage based on classifying critical data according a priority status and escalating between statuses based on particular thresholds. For example, priority can be escalated based on a frequency of event threshold such as the number of misaligned staples fired by a stapling surgical instrument 7012 over a predetermined number of surgical operations.

[1561] Multiple staggered frequency or other thresholds could also be used. The lowest priority level of the priority status classification 11106 is the watch list level designated at level A. As discussed above, the watch list may be a black list maintained in the memory 7010 as a set of database records of identifiers corresponding to prohibited surgical hubs 7006. Surgical hubs 7006 can be prohibited to different extents depending on the nature of the critical device data 11002 or additional data. For example, surgical hubs 7006 may be partially locked out such that only the device components experiencing problems are prevented from functioning. Alternatively, surgical hub 7006 on the watch list may not be restricted from functioning in any way. Instead, the surgical hubs 7006 may be monitored by the cloud 7004 for any additional irregularities that occur. Accordingly, the watch list is designated at level A, the least urgent priority status. As shown in the priority status classification 11106, the automated response at level B is the next most urgent priority status.

[1562] An automated response could be, for example, an automated initial analysis of the device data 11002 by the patient outcome analysis module 7028 of the cloud 7004 via a set of predefined diagnostic tests.

[1563] The third most urgent priority status is notification, which is designated at level C of the priority status classification 11106. In this situation, the cloud 7004 transmits a wireless signal to a healthcare facility employee, clinician, healthcare facility department, or other responsible party depending on the nature of the device data 11002.

[1564] The notification signal can be received at a receiver device located at a suitable location within the healthcare facility, for example. Receiving the notification signal can be indicated by a vibration or sound to notify the responsible party at the healthcare facility. The holder of the receiver device (e.g., a healthcare facility clinician) may then conduct further analysis of the critical device data 11002 or additional data or other analysis for resolving an indicated irregularity. If a solution to the irregularity is known, the solution may be timely implemented. The most urgent priority status as depicted in the priority status classification 11106 is urgent action required, which is designed at level D. Urgent action required indicates that a responsible party, device or instrument should immediately analyze and diagnose the problem implicated by the critical data. Upon proper diagnosis, an appropriate response should immediately be performed. In this way, the cloud 7004 may implement a comprehensive approach to critical data prioritization and triaging that no individual medical facility could achieve on its own. Critical data may be handled in a timely manner according to suitable priority levels which can address solving time sensitive problems that arise in the healthcare field. Moreover, the cloud 7004 can prioritize aggregated critical data from all healthcare facilities categorized within a particular region. Accordingly, the time sensitive prioritized approach to handling critical data can be applied

system wide, such as to a group of healthcare facilities. Furthermore, the cloud 7004 can generate an alert for a responsible party to respond to critical data (and associated issues implicated by such critical data) in a timely way such as in real time or in near real time according to a corresponding priority status. This alert can be received by a suitable receiver of the responsible party. The priority status of the device data 11002 could also be determined based on the severity of the surgical issue implicated by the device data 11002. As discussed above, the cloud 7004 may receive additional data from surgical hubs 7006 or surgical instruments 7012 (via the hubs 7006) which causes the cloud 7004 to elevate the priority status of the device data 11002.

[1565] In one aspect, based on a priority status, the device data 11002 may be subject to the flagging screening at a specific time depending on priority. For example, the device data 11002 may be indicated as critical data but not yet flagged. Additionally, the device data 11002 may first receive an automated response level of priority according to the priority status classification 11106. In this situation, the severity determination at step 11108 may be relatively quickly in accordance with the level B of priority. Specifically, step 11108 may be reached without first placing the surgical instrument 7012 on a watch list. The severity threshold used at step 11108 can be the same or different from the severity threshold used in 11006. Aside from the severity determination at step 11108, other determinations pertinent to the irregularity indicated by the critical device data 11002 or additional data may be made. These determinations may be used to diagnose the occurrence of a critical event. Accordingly, if the answer at step 11108 is yes, the frequency of the event may be assessed at step 11110. Conversely, if the answer at step 11108 is no, the device data 11002 or additional data can be stored 11118 in the hold list. Additionally or alternatively, the device data 11002 or additional data can be routed to different storage locations within the cloud 7004 according to the routing branch of the flow diagram (labeled as Q2 in FIG. 199). The cloud 7004 may wait for a request from the hub 7006 for alternative routing 11120 of the device data 11002 or additional data. At step 11110, the cloud 7004 determines the frequency that the critical event is occurring. Based on this frequency, the priority status assigned according to the priority status classification 11106 can be escalated (see step 11116). For example, the critical event may be the generator of the surgical instrument 7012 is applying an insufficient sealing temperature to therapeutically treat tissue. In other words, the inquiry of step 11110 inquires whether the medical event implicated by the critical data is occurring at an increasing frequency after the problem was initially identified.

[1566] An increase in the number of times this insufficient sealing temperature occurs can be monitored to escalate priority status at step 11116, based on frequency thresholds (see step 11112), for example. If at step 11110, the event is not increasing infrequency, the data can be stored 11118 in the hold list. If the answer at step 11110 is yes (i.e., the event is increasing in frequency), the flow diagram proceeds to step 11112. At step 11112, another data verification inquiry is made. In particular, specific thresholds such as the frequency thresholds described above may be applied to determine whether the combination of device data 11002 or additional data is sufficiently correct to ensure that the critical data should be added to the aggregated medical data databases 7011. Furthermore, the data verification inquiry at

step 11112 may comprise a decision regarding whether the sample size of the critical data is sufficiently large (i.e., reached critical mass). Additionally or alternatively, the sample size is analyzed for whether there is sufficient information to determine an appropriate internal response 11102 of the cloud 7004. The data verification inquiry can also comprise verifying the accuracy of the data by comparison to predetermined standards or verification tests. If the answer to the inquiry at step 11112 is negative, then the critical data is stored within the separate storage location (e.g., hold list) in the cloud 7004. If the answer to the inquiry at step 11110 is affirmative, the device data 11002 or additional data is added to the aggregated medical data databases 7011. At step 11116, the priority status of the device data 11002 or additional data is increased according to the priority status classification 11106. However, besides the event frequency determination, the addition to the aggregated medical data databases 7011 may itself be an action that results in an elevation of the priority status of the critical data at step 7. In any case, the priority status of the device data 11002 or additional data may be escalated or deescalated as appropriate based on additional analysis or data, for example. An internal response 11102 of the cloud 7004 may be made according to the current priority status (i.e., one of levels A-D) of the critical data.

[1567] In addition to prioritizing critical data, the internal response 11102 of the cloud 7004 can also involve advantageously routing, grouping, or sorting critical data the aggregated critical data in a timely manner. In particular, the data may be routed to different storage locations within the cloud 7004, such as in the memory devices 7010. This routing is illustrated by routing branch of the flow diagram labeled as Q2 in FIG. 199 at step 11120. As such, the memory devices 7010 of the central servers 7013 of the cloud 7004 can be organized into various locations that correspond to a characteristic of the critical data or a response corresponding to the critical data. For example, the total memory capability of the memory devices 7010 may be divided into portions that only store data according to individual data routing categories, such as those used at steps 11122, 11124, 11126. As shown at step 11120 of FIG. 199, the critical data may be routed to different various cloud storage locations. Step 11120 can occur in conjunction with or separately from the prioritization branch of the flow diagram. Step 11120 may be triggered by a request generated by a hub 7006. The hub 7006 may transmit such a request because of detecting a failure or irregularity associated with a surgical instrument 7012, for example. The associated critical data may then receive alternative routing 11120 by the cloud 7004 to different cloud storage locations. At step 11122, the alternative routing 11120 can comprise geographical location based routing. That is, the different cloud storage locations may correspond to location based categorization of the cloud memory devices 7010. Various subsets of the cloud memory devices 7010 can correspond to various geographical regions. For example, surgical instruments produced from a manufacturing plant in Texas could be grouped together in storage within the cloud memory devices 7010. In another example, surgical instruments produced from a specific manufacturing company can be categorized together in the cloud memory devices 7010. Therefore, location based categorization can comprise the cloud 7004 routing critical data based on associations with different manufacturing sites or operating companies.

[1568] At step 11124, the alternative routing 11120 can comprise routing for device data 11002 or additional data that requires a rapid internal response 11102 of the cloud 7004. This alternative routing 11120 at step 11124 could be integrated with the priority status classification 11106. For example, escalated or urgent priority critical data, such as those at priority level C and D, may be routed by the cloud 7004 to rapid response portions of the memory devices 7010 to enable a rapid response. For example, such critical data may be routed to rapid response caches 7018 which signifies that a rapid response is necessary. At step 11126, device data 11002 or additional data that implicates a failure of a type that requires special processing are routed to a special processing portion of the memory devices 7010. For example, a surgical instrument 7012 may be determined to have experienced a failure or malfunction during operation based on a control program deficiency common to a whole group of surgical instruments 7012. In this situation, special processing may be required to transmit a collective control program update to the group of surgical instruments 7012. Accordingly, the cloud may route the critical data to the special processing portion of the memory devices 7010 to trigger this special processing. Subsequently, the special processing could also include the patient outcome analysis data analytics module 7028 analyzing and monitoring the effect of the control program update on patient outcomes. The patient outcome analysis module 7028 may also execute an automated product inquiry algorithm as discussed above if necessary.

[1569] FIG. 200 is a flow diagram of an aspect of data sorting and prioritization by the computer-implemented interactive surgical system, according to one aspect of the present disclosure. This sorting and prioritization may be implemented by the data sorting and prioritization module 7032, the data collection and aggregation module 7022, and patient outcome analysis module 7028. As discussed above, critical device data 11002 or additional data can implicate or correspond to various medical events, such as events 1 through 3 as depicted in FIG. 200. An event may be for example, a shift from a phase of tissue treatment to another phase such as a shift from a phase corresponding to cutting with the specific surgical instrument to a phase corresponding to coagulation. In FIG. 200, critical data associated with a first medical event 11202 is detected by the surgical hub 7006 and transmitted to the cloud 7004. Upon receiving the critical data, the cloud 7004 analyzes the critical data at step 11208 to determine that it is comparable to an expected value of the critical data, as described above for example at step 11016. When the critical data is determined as comparable (i.e., the value of the critical data is expected), the critical data may be aggregated within a large data set in the aggregated medical data databases 7011, for example. That is, at step 11216, the critical data is stored within the aggregated databases of the cloud. As shown in FIG. 200, the critical data is also subject to a binary classification at steps 11218, 11220. For example, the critical data can be distinguished by good properties and bad properties. The data sorting and prioritization modules can classify the critical data as associated with a bleeding or a non-bleeding event, for example. In this way, the patient outcome analysis module 7028 may classify critical data as corresponding to a positive patient outcome at step 11218 or a negative patient outcome at step 11210.

[1570] FIG. 200 also shows the critical data associated with a second medical event 11204 is detected by the surgical hub 7006 and transmitted to the cloud 7004. The critical data associated with the second medical event 11204 is determined by the cloud to be suspicious or unusual data at step 11210, which is a trigger condition as described above with reference to step 11118. Accordingly, the cloud 7004 is triggered to request 11114 additional data from the surgical hub 7006 at step 11212 by transmitting a push message to the surgical hub 7006. As discussed above, the additional data may enable the patient outcome analysis module 7028 of the cloud 7004 to gain additional insight into the source of the irregularity implicated by the critical data. If the patient outcome analysis module 7028 sufficiently diagnoses the cause of the second medical event 11214, the critical data or associated additional data is aggregated into the aggregated medical data databases 7011 at step 11216 (see also step 11114). Subsequently, the critical data or additional data is classified according to the good/bad binary classification at steps 11218, 11220. If the cloud 7004 cannot sufficiently diagnose the cause of the second medical event 11204, the process may proceed to step 11224, in which the critical data is evaluated by a suitable person or department of the corresponding medical facility. Step 11224 can include the threshold data exclusion determination at step 11104. That is, because a good reason cannot be readily determined for the suspicious or unusual data, the data may be stored in a hold list in accordance with step 11118. Additionally, the device data 11002 or additional data may be designated at priority status level C, which triggers the evaluation at step 11224 (i.e., healthcare facility employee, clinician, healthcare facility department, or other responsible party evaluates the data).

[1571] As illustrated in FIG. 200, the critical data associated with a third medical event 11206 is detected by the surgical hub 7006 and transmitted to the cloud 7004. The critical data associated with the third medical event 11206 is determined by the cloud 7004 to indicate that the corresponding surgical instrument 7012 is experiencing a failure or malfunction at step 11220. As discussed above, severity thresholds can be used to determine whether the failure is severe. The failure or malfunction may refer back to the trigger condition at step 11022 in FIG. 198 such that the surgical instrument malfunction results in an automated product inquiry through the surgical hub 7006. As discussed above, the automated product inquiry algorithm may comprise the patient outcome analysis module 7028 searching for data of related incidents stored within the cloud 7004 (e.g., the memory devices 7010). The data of related incidents can include video, manufacturer, temporal, and other suitable types of data. Depending on the results of the automated product inquiry, the third medical event 11206 critical data can be prioritized according to priority status classification 11106. Thus, for example, the inquiry may result in a suspicious or unusual result without a sufficient reason, so the critical data is designated at priority level C. In this connection, a suitable person or department of the corresponding medical facility evaluates the critical data and the results of the automated product inquiry at step 11224. The results of the evaluation could be, for example, that the results constitute an error to be disregarded at step 11226 or that the results require additional special processing via the patient outcome analysis module 7028 at step 11228 (see also step 11126). Such special processing at step 11228 can

be the CAPA portion of the automated product inquiry algorithm, as described above. Thus, the cloud-based analytics system may generate timely alerts for triggering a response by the suitable person or department in real time or near real time.

[1572] In general, the cloud-based analytics system described herein may determine critical data and perform timely data handling, sorting, and prioritizing based on priority status and specific thresholds as described above. Accordingly, the cloud-based analytics system advantageously handles critical data in a timely, systematic, and holistic manner over multiple health care facilities. The critical data handling comprises internal responses by the cloud 7004 based on assigned priority levels. Moreover, based on requests by surgical hubs 7006, special routing of data within the memory device 7010 of the cloud 7004 may be achieved. The rerouting, prioritizing, confirming, or requesting supporting as described above may be used to improve analysis of the data by the cloud 7004.

#### Cloud Interface for Client Care Institutions

[1573] All client care institutions require some level of control in a treatment environment. For example, an institution may wish to control inventory that is present within an operating room. Inventory items within an operating room may include not only medical devices to be used during surgery (e.g., scalpels, clamps, surgical tools, etc.) but also medical supplies to be used during surgery in conjunction with such medical devices (e.g., gauze, sutures, staples, etc.). Heretofore, inventory control for many institutions comprises a simple manual count of inventory items on a periodic basis (e.g., daily, weekly, monthly, etc.). Similarly, other institutions utilize a barcode scanner to count and/or document inventory items on a periodic basis.

[1574] Aspects of the present disclosure are presented for a cloud interface accessible by participating client care institutions via a cloud-based analytics system. In order to monitor and/or control inventory items to be utilized or being utilized by an institution, each institution adopts its own practice of documenting inventory item usage. For example, an institution may manually count and/or scan inventory items on a periodic basis. Additional example details are disclosed in U.S. Patent Application Publication No. 2016/0249917, titled SURGICAL APPARATUS CONFIGURED TO TRACK AN END-OF-LIFE PARAMETER, which published on Sep. 1, 2016, U.S. Patent Application Publication No. 2014/0110453, titled SURGICAL INSTRUMENT WITH RAPID POST EVENT DETECTION, which issued on Feb. 23, 2016 as U.S. Pat. No. 9,265,585, U.S. Patent Application Publication No. 2016/0310134, titled HANDHELD ELECTROMECHANICAL SURGICAL SYSTEM, which published on Oct. 27, 2017, and U.S. Patent Application Publication No. 2015/0317899, titled SYSTEM AND METHOD FOR USING RFID TAGS TO DETERMINE STERILIZATION OF DEVICES, which published on Nov. 5, 2015, the entire disclosures of which are hereby incorporated by reference herein. Information regarding counted and/or scanned inventory items may then be stored in a local computer system to track inventory item usage. Such a manual process is not only labor intensive and inefficient, but also prone to human error. As a result, an institution may be unable to perform a surgical procedure(s) and/or the surgical procedure(s) may be unnecessarily delayed because one or more inventory items, required for

the surgical procedure(s), are not available for use for various reasons (e.g., out of stock, in stock but expired, in stock but no longer considered sterile, in stock but defective, etc.). Knowing this, some institutions are forced to carry and/or hold an overstock of inventory items. This, of course, may result in increase expense (e.g., more inventories) and ultimately unnecessary waste (e.g., expired inventory items).

[1575] To help institutions control inventory items, it would be desirable for institutions to have access, via a cloud interface, to a cloud-based analytics system configured to automate inventory control by automatically receiving data associated with inventory items of the institutions, deriving information based on the received data, and conveying, via the cloud interface, real-time knowledge back to the institutions regarding inventory items. Referring to FIG. 201, according to one aspect of the present disclosure, a client care institution system 8000 may transmit (e.g., periodically, in real-time, in batches, etc.) inventory data to a cloud-based analytics system 8002 and the cloud-based analytics system 8002 may derive/extract information from that inventory data. In such an aspect, a cloud-interface 8004 may be accessed/queried by the client care institution system 8000 and the cloud-based analytics system 8002 may transmit its derived/extracted information to the cloud-interface 8004. Further, in such an aspect, the cloud-interface 8004 may convey/package/structure the derived/extracted information to the client care institution system 8000 to reveal knowledge about the client care institution's inventory. In one aspect, the client care institution system may comprise a surgical system 102 (e.g., FIG. 1), the cloud-based analytics system may comprise the cloud-based system 105 (e.g., FIG. 1) and the cloud-interface may comprise at least one of a visualization system 108/208 (e.g., FIGS. 1-2) or a display 135/177 associated with the surgical hub 106 (e.g., FIGS. 1-3, 7, etc.).

[1576] Referring to FIG. 1, in some aspects of the present disclosure, a cloud-based system 105 is communicatively coupled to one or more than one surgical hub of an institution (e.g., one or more than one surgical hub 106 of a surgical system 102). Here, each surgical hub is in communication (e.g., wirelessly) with one or more than one inventory item (e.g., intelligent instrument 112). The cloud-based system 105 may be configured to aggregate data associated with each inventory item of each institution, analyze that data with respect to system-defined constraints, and generate or facilitate a cloud interface for each institution to monitor and control inventory items. In one example, the cloud-based system 105 may be configured to compute a current availability of each inventory item (e.g., an indication of real-time usage and/or scheduled usage for each inventory item in a surgical system 102), a current usage associated with each inventory item (e.g., based on data received from one or more than one surgical hub 106 that has read usage data from a chip/memory associated with each inventory item), irregularities, if any, associated with each inventory item (e.g., defects, etc.), current possible medical device combinations that utilize each inventory item (e.g., various shafts, staple cartridges, end effectors, etc. combinable to form numerous medical device combinations), and available alternatives to each inventory item (e.g., available shaft B and/or shaft C may be substituted for unavailable shaft A for a desired/input surgical procedure(s)). Referring to FIGS. 202-203, in such an exemplification, after input of a desired



surgical procedure(s) (e.g., “cholecystectomy”) by an institution in its cloud interface **8104**, the cloud-based system **105** may provide up-to-date, real-time and/or near real-time knowledge regarding the availability and/or usability of inventory items (e.g., associated with and/or needed to perform the input surgical procedure(s) based on the system-defined constraints. Referring to FIG. **203**, in one example, the institution’s cloud interface **8104** may display an inventory item **8106** (e.g., Handles A, B, and C) in association with its current **8108** and/or remaining usage **8110**. If the remaining usage is not adequate (e.g., based on anticipated usage necessary for the desired surgical procedure, etc.), the cloud interface may further display a warning or alert regarding the inadequacy (e.g., **8112**, highlighting, blacked out, etc.). Such a warning or alert may indicate that the surgical procedure(s) input at the cloud interface cannot be performed based on current inventory items. In one aspect, a same or similar warning or alert may be communicated to the inventory item itself for display on a user interface of the inventory item itself (e.g., a user interface of Handle C). In another aspect, the cloud interface may further display available alternatives to the inventory item (e.g., Handle B). Here, anticipated usage and/or available alternatives may be determined at the surgical hub **106** (e.g., based on local data) and/or the cloud-based analytics system **105** (e.g., based on local data of the surgical hub **106** and/or global data from multiple surgical hubs **106** of multiple institutions). In one example, the surgical hub **106** may infer anticipated usage and/or available alternatives from local data associated with the same or similar surgical procedure (e.g., average number of uses to perform the same or similar surgical procedure, alternative inventory items used to perform the same or similar surgical procedure, etc.). In another example, the cloud-based analytics system **105** may similarly infer anticipated usage and/or available alternatives from local data of the surgical hub **106** and/or global data from multiple surgical hubs **106** of multiple institutions (e.g., average number of uses to perform the same or similar surgical procedure, alternative inventory items used to perform the same or similar surgical procedure, etc.).

[1577] In other aspects of the present disclosure, a cloud-based system **105** is communicatively coupled to one or more than one surgical hub **106** of an institution, each surgical hub **106** in communication (e.g., wirelessly) with one or more than one inventory item (e.g., intelligent instrument **112**). The cloud-based system **105** may be configured to create a list of inventory items not authorized to perform surgical procedures due to one or more system-defined constraints. In one exemplification, after input of a desired surgical procedure(s) by an institution into its cloud interface (e.g., FIG. **202**), the cloud-based system **105** may determine that one or more inventory items of the institution (e.g., detected by and associated with and/or needed to perform the input surgical procedure(s)) are not authorized to perform the input surgical procedure(s) based on system-defined constraints. In such an exemplification, it may be determined that an identifier (e.g., serial number, unique ID, etc.) associated with an inventory item is not authorized to perform the input surgical procedure(s) (e.g., inventory item exceeds usable life, inventory item is counterfeit, inventory item is defective, etc.). In one example, the institution’s cloud interface may display an inventory item in association with its unauthorized status **8114**. In such an aspect, the cloud interface may further display a warning or alert

regarding the unauthorized status (e.g., highlighting, blacked out, etc.). Such a warning or alert may indicate that the surgical procedure(s) input at the cloud interface cannot be performed based on current inventory items. In one aspect, a same or similar warning or alert may be communicated to the inventory item itself for display on a user interface of the inventory item itself (e.g., a user interface of Handle D). Similar to above, the cloud interface **8104** may display available alternatives to the unauthorized inventory item (e.g., Handle B).

[1578] In yet other aspects of the present disclosure, a cloud-based system **105** is communicatively coupled to one or more than one surgical hub **106** of an institution, each surgical hub **106** in communication (e.g., wirelessly) with one or more than one inventory item (e.g., intelligent instrument **112**). The cloud-based system **105** may be configured to create a list of inventory items no longer authorized to perform surgical procedures due to one or more system-defined constraints. In one exemplification, after input of a desired surgical procedure(s) by an institution in its cloud interface (e.g., FIG. **202**), the cloud-based system may determine that one or more inventory items are no longer authorized to perform the input surgical procedure(s) based on system-defined constraints. In such an exemplification, it may be determined that an identifier (e.g., serial number, unique ID, etc.) associated with an inventory item is unusable (e.g., expired, no longer sterile, defective, etc.). In one example, the institution’s cloud interface may display an inventory item in association with its unusable status **8116**. In such an aspect, the cloud interface may further display a warning or alert regarding the unusable status (e.g., highlighting, blacked out, etc.). Such a warning or alert may indicate that the surgical procedure(s) input at the cloud interface cannot be performed based on current inventory items. In one aspect, a same or similar warning or alert may be communicated to the inventory item itself for display on a user interface of the inventory item itself (e.g., a user interface of Handle E). Similar to above, the cloud interface may display available alternatives to the unusable inventory item (e.g., Handle B).

[1579] In this way, the cloud-based system **105** of the present disclosure may provide up-to-date, real-time, and/or near real-time knowledge regarding the availability of inventory items pertinent to the surgical procedure(s) input to the cloud interface of the participating institutions. Such a system goes well-beyond conventional processes of manually counting and/or scanning inventory items.

[1580] FIG. **204** illustrates an example multi-component surgical tool (e.g., a wireless surgical device/instrument **235**) comprising a plurality of modular components **8204**, **8206**, **8208**, **8210**, wherein each modular component is associated with an identifier **8214**, **8216**, **8218**, **8220** respectively (e.g., a serial number). In particular, the surgical tool **235** of FIG. **204** includes a handle **8204**, a modular adapter **8206**, and end effector **8208** (e.g., a disposable loading unit and/or a reloadable disposable loading unit in various aspects), and a staple cartridge **8210**. In this example, the handle **8204** is associated with serial number “SN135b”, the modular adapter **8206** is associated with serial number “SN33b”, the end effector **8208** is associated with serial number “SN1a” and the staple cartridge **8210** is associated with serial number SN121b. In such an aspect, each modular component (e.g., **8204**, **8206**, **8208**, **8210**, etc.) is configured to request a communication link to a surgical hub **106** of an

institution. In other aspects, the surgical hub **106** may be configured to request a communication link with each modular component. Nonetheless, the surgical hub **106** is positioned within a communicative distance from each modular component (e.g., in an operating room). In one aspect of the present disclosure, a requested communication link is established via BLUETOOTH pairing. In other aspects of the present disclosure, other forms of wireless communication (e.g., WiFi, RFID, etc.) or wired communication are contemplated. Referring again to FIG. **204**, each modular component (e.g., handle **8204**, modular adapter **8206**, end effector **8208**, staple cartridge **8210**, etc.) may comprise a processor and a memory unit (not shown) that stores its respective serial number. Here, according to one aspect, once a communication link is established between the surgical hub **106** and each modular component, the identifier (e.g., serial number) associated with each modular component is transmitted by each modular component to the surgical hub **106** (e.g., via the same form or different forms of wired/wireless communication). In one alternative aspect, in light of FIG. **204**, a modular component (e.g., modular adapter **8206**, end effector **8208**, and/or staple cartridge **8210**, etc.) may transmit its respective identifier (e.g., serial number) to another modular component (e.g., handle **8204**) that transmits/relays all identifier(s) to the surgical hub **106**. Here, similar to above, the same form or different forms of wired/wireless communication may be used. For example, each of the modular adapter **8206**, the end effector **8208** and the staple cartridge **8210** may transmit its respective identifier (e.g., **8216**, **8218**, **8220**) to the handle **8204** via RFID and the handle **8204** may relay such identifiers (e.g., **8216**, **8218**, **8220**) along with its own identifier **8214**, via BLUETOOTH, to the surgical hub **106**. In one aspect, once the surgical hub **106** has received all identifiers for all modular components, the surgical hub **106** may transmit the identifiers to the cloud-based analytics system (e.g., comprising cloud-based system **105**).

[1581] In various aspects of the present disclosure, the memory unit of each modular component may be configured to store more than its identifier. In one aspect of the present disclosure, each modular component (e.g., **8204**, **8206**, **8208**, **8210**, etc.) may further comprise a counter (not shown) configured to track a usage parameter of the modular component and its memory unit may be configured to store that usage parameter. In another aspect, the memory unit of each respective modular component may be further configured to store a usable life metric. Such a usable life metric may be stored during manufacture of the modular component. For example, in view of FIG. **204**, the memory unit of the handle **8204** may store both the usage parameter (e.g., **235**) and the usable life metric (e.g., **400**). In such an aspect, the handle **8204** has been used 235 times out of its usable life of 400 uses. Similarly, in view of FIG. **204**, the modular adapter has been used 103 times out of its usable life of 100 uses, and the end effector has been used 5 times out of its usable life of 12 uses. Here, similar to above, once a communication link is established with the surgical hub **106**, the identifier, usage parameter and/or usable life metric stored in the memory unit of each modular component may be transmitted directly from each modular component to the surgical hub **106** or indirectly via another modular component. In addition, similar to above, the same form or different forms of wired/wireless communication may be used. In one aspect, once the surgical hub **106** has received all identifiers

for all modular components, the surgical hub **106** may transmit the identifiers to the cloud-based analytics system (e.g., comprising cloud-based system **105**).

[1582] In an alternative aspect of the present disclosure, the memory unit of each modular component may not store its usage parameter and/or the usable life metric. In such an aspect, the usage parameter and/or the usable life metric may be stored in a database or other memory (see FIG. **10**, e.g., **248/249**) at the surgical hub **106/206**. In such an aspect, the surgical hub **106** may comprise a counter configured to track a usage parameter of each modular component in inventory. Furthermore, the surgical hub **106** may be configured to download usable life metrics (e.g., from a manufacturer server) based on the identifier (e.g., serial number) received from each modular component. In various aspects, storage at the surgical hub **106** may be preferred to minimize memory unit requirements in each modular component and/or to avoid any concerns regarding the tampering with and/or the alteration of usage parameters and/or usable life metrics stored at the modular component level (e.g., altering a memory unit of a modular component to reset a usage parameter and/or increase a usable life metric, etc.).

[1583] In one example, in aspects where the memory unit of each modular component stores its usage parameter and/or usable life metric, the surgical hub **106** may also store/track the usage parameter and/or usable life metric associated with each modular component in its inventory. In such an example, if a usage parameter and/or a usable life metric transmitted from a modular component differs from a usage parameter and/or a usable life metric stored/tracked at the surgical hub **106**, the surgical hub **106** may flag the discrepancy and modify the status of that modular component (e.g., to unavailable, to unauthorized, to unusable, etc.).

[1584] In another alternative aspect, the memory unit of each modular component may not store its usage parameter and/or the usable life metric. In such an aspect, the usage parameter and/or the usable life metric may be stored in a database (e.g., aggregated medical data database **7012** in FIG. **180**) at a cloud-based analytics system. In such an aspect, the cloud-based analytics system may comprise a counter configured to track a usage parameter of each modular component in inventory at each surgical hub. Furthermore, the cloud-based analytics system may be configured to download usable life metrics (e.g., from a manufacturer server) based on the identifier (e.g., a serial number) received from each modular component (e.g., via a surgical hub). Alternatively, the cloud-based analytics system may download a file comprising all identifiers for all modular components (e.g., from a plurality of manufacturers) wherein each identifier is associated with a usable life metric. Here, the cloud-based analytics system may be configured to look-up a received identifier to determine each respective usable life metric. In various aspects, storage at the cloud-based analytics system may be preferred to minimize memory requirements in each modular component and/or to avoid any concerns regarding the tampering with and/or the alteration of usage parameters and/or usable life metrics at the modular component level and/or at the surgical hub level (e.g., altering memory unit of a modular component to reset a usage parameter and/or increase a usable life metric, modifying the database/memory of the surgical hub to reset a usage parameter and/or increase a usable life metric). Such an aspect gives the cloud-based

analytics system of the present disclosure more control over modular component use in the interactive surgical system.

[1585] Looking again to FIG. 204, the illustrated multi-component surgical tool 235 comprises four modular components (e.g., handle 8204, modular adapter 8206, end effector 8208, and staple cartridge 8210). Such modular devices may comprise reusable and/or reprocessed components. In various aspects, each modular component must satisfy system-defined constraints for the combined multi-component surgical tool 235 to be available/usable/authorized for use by the cloud-based analytics system. Notably, system-defined constraints may include restrictions other than and/or in addition to the usable life metric discussed above. Such system-defined constraints may be established at the manufacturer level, at the surgical hub level, and/or at the cloud-based analytics system level. One aspect of the present disclosure comprises a user interface at the surgical hub and/or cloud-based analytics system to create system-defined constraints.

[1586] In one aspect, the surgical hub 106 may be configured to enforce system-defined constraints (e.g., lockout at the hub level). In such an aspect, this may be preferred so that the surgical hub 106 is a local gateway to accessing the cloud-based analytics system. In another aspect, the cloud-based analytics system (e.g., comprising cloud-based system 105) may be configured to enforce system-defined constraints (e.g., lockout at the cloud-based analytics system level). In such an aspect, this may be preferred to maintain control over all surgical hubs communicatively coupled to the cloud-based analytics system (e.g., at one institution or at multiple institutions). System-defined constraints, similar to the usable life metric, may be associated with the identifier of each modular component. For example, a system-defined constraint associated with a modular component may include an expiration date, a requirement that an identifier (e.g., serial number) is a system-recognizable identifier (e.g., not counterfeit), and/or flexible system-defined constraints (e.g., constraints deemed non-critical until a threshold is met and the constraint is deemed critical). In one aspect of the present disclosure, if one system-defined constraint is not met, a modular component (e.g., 8204, 8206, 8208, 8210, etc.) may be deemed unavailable/unusable/unauthorized despite being available/usable/authorized based on other system-defined constraint(s) (e.g., having remaining usable life). In various aspects, one or more predetermined system-defined constraints are non-critical system-defined constraints. Such non-critical system-defined constraints may be waived (see FIG. 204, e.g., 8274, manual override) to render the modular component available/usable/authorized and/or may produce in a warning indicator/message (see FIG. 204, e.g., 8244). Critical system-defined constraints cannot be waived.

[1587] In view of FIG. 204, an example non-critical system-defined constraint is applied (e.g., by the surgical hub 106 and/or the cloud-based analytics system) to the handle 8204. Here, although the handle 8204 has 165 remaining uses (usable life metric less determined usage parameter, e.g., 400-235) an expiration date associated with its identifier 8214 (e.g., SN135b) indicates that the handle's control program is out-of-date. In such an aspect, an interface 8200 may be displayed to show a current status of the handle 8204 (see FIG. 204, e.g., "Count 235/400" and/or "Out-of-Date"). More specifically, the interface 8200 may comprise a grid including fields defined by columns and

rows. In one example, the modular components of a proposed multi-component surgical tool 235 may be presented (e.g., in an exploded, unassembled view) across the columns of the grid in a first row 8201 and a current/updated status associated with each modular component may be presented across corresponding columns of the grid in a second row 8202. As such, in accordance with the example, status field 8224 of the interface 8200 corresponds to the handle 8204 and indicates its current status as "COUNT: 235/400" and "OUT-OF-DATE". According to other aspects, the status field 8224 of the interface 8200 may further show the usage remaining, remaining capabilities, and/or compatibility with other connected modular components, etc.

[1588] According to one aspect, the interface 8200 may comprise a cloud-based interface (see FIG. 203, e.g., 8104) accessible on a cloud-access terminal of the surgical hub (via at least one of a visualization system 108/208 (e.g., FIGS. 1-2) or a display 135/177 associated with the surgical hub 106 (e.g., FIGS. 1-3, 7, etc.)). According to another aspect, the interface 8200 may comprise only a portion(s) of the grid (e.g., status field 8224, modular component field 8234, etc.) accessible on the physical handle 8204 itself via a user interface positioned on the handle 8204. Further, in the context of a non-critical system-defined constraint, the interface 8200 may visually indicate a warning associated with a modular component (e.g., warning indicator 8244, e.g., box associated with identifier 8214 highlighted and/or encircled and/or comprises a link 8254 (e.g., "A") in association with modular component field 8234 of the interface 8200). In one aspect, the link 8254 (e.g., "A") may key to a corresponding "Description of Problem" section of the interface 8200 (e.g., "A" "Handle Serial Number Indicates OUT OF DATE Control Program"). In another aspect, the link 8254 (e.g., "A") may be a hyperlink to present the corresponding description (e.g., "A" "Handle Serial Number Indicates OUT OF DATE Control Program") in the interface 8200. According to such aspects, a portion of the descriptive text (e.g., "OUT OF DATE"), keyed/hyperlinked via link 8254, may be a hyperlink/button 8264. Upon/After selection of the hyperlink/button 8264 a bypass interface 8274 may be presented in the interface 8200. According to another aspect, a portion of descriptive text (e.g., OUT-OF-DATE) in status field 8224 may be a hyperlink/button 8284 to, upon/after selection, directly present the bypass interface 8274 in the interface 8200. Such an aspect may be beneficial/more efficient if the interface 8200 is being presented via a (e.g., smaller) user interface of a modular component (e.g., handle 8204). Further, according to such aspects, the interface 8200 may be configured to receive user input to waive (e.g., manually bypass) a predetermined, non-critical system-defined constraint (e.g., the expiration date constraint). In the context of a non-critical system-defined constraint, the bypass interface 8274 may instruct "USER INPUT NEEDED" and present a first user-interface element (e.g., "Y" button) selectable to bypass the non-critical system-defined constraint (e.g., to permit use of the handle 8204) and a second user-interface element (e.g., "N" button) selectable to not bypass the non-critical system-defined constraint (e.g., to inhibit use of the handle 8204). Here, a selection in the bypass interface 8274 may be transmitted to update the surgical hub 206 and/or the cloud-based system 205.

[1589] Next, in view of FIG. 204, an example flexible system-defined constraint is applied (e.g., by the surgical

hub **106** and/or the cloud-based analytics system) to the modular adapter **8206**. Here, the modular adapter **8206** associated with identifier **8216** (e.g., SN33b) has a usage parameter of **103** (e.g., already 3 times over its suggested usable life metric of 100 uses). In this example, the exceeding use is deemed non-critical until a 10% overage threshold is met (e.g., 110% of the suggested 100 uses, or 110 uses) and the exceeding use is deemed critical. In such an aspect an interface **8200** may be displayed to show a current status of the modular adapter **8206** (see FIG. **204**, e.g., “COUNT: **103/100**” “EXCEEDS”). More specifically, in accordance with the example described above, status field **8226** corresponds to the modular adapter **8206** and indicates its current status as “COUNT: **103/100**” and “EXCEEDS”. According to other aspects the status field **8226** of the interface **8200** may further show overage remaining, remaining capabilities, and/or compatibility with other connected modular components.

[1590] Again, according to one aspect the interface **8200** may comprise a cloud-based interface (see FIG. **203**, e.g., **8104**) accessible on a cloud-access terminal of the surgical hub (via at least one of a visualization system **108/208** (e.g., FIGS. **1-2**) or a display **135/177** associated with the surgical hub **106** (e.g., FIGS. **1-3**, **7**, etc.)). According to another aspect, the interface **8200** may comprise only a portion(s) of the grid (e.g., the status field **8226**, modular component field **8236**, etc.) accessible directly on the physical modular adapter **8206** itself via a user interface positioned on the modular adapter **8206** and/or indirectly on the physical handle **8204** itself via a user interface positioned on the handle **8204**. Further, in the context of a flexible system-defined constraint, the interface **8200** may visually indicate a warning associated with a modular component (e.g., warning indicator **8246**, e.g., description of current status encircled and/or comprises a link **8256** (e.g., “B”) in association with status field **8226** of the interface **8200**). In one aspect, the link **8256** (e.g., “B”) may key to a corresponding “Description of Problem” section of the interface **8200** (e.g., “B” “Modular Adapter EXCEEDS Suggested Life Limit”). In another aspect, the link **8256** (e.g., “B”) may be a hyperlink to present the corresponding description (e.g., “B” “Modular Adapter EXCEEDS Suggested Life Limit”) in the interface **8200**. According to such aspects, a portion of the descriptive text (e.g., “EXCEEDS”), keyed/hyperlinked via link **8256**, may be a hyperlink/button **8266**. Upon/After selection of the hyperlink/button **8266** a warning interface **8276** may be presented in the interface **8200**. According to another aspect, a portion of descriptive text (e.g., EXCEEDS) in status field **8226** may be a hyperlink/button **8286** to, upon/after selection, directly present the warning interface **8276** in the interface **8200**. Such an aspect may be beneficial/more efficient if the interface **8200** is being presented via a (e.g., smaller) user interface of a modular component (e.g., modular adapter **8206** and/or handle **8204**). Further, according to such aspects, the interface **8200** may be configured to present a warning that the modular adapter **8206** is approaching its overage threshold. In one aspect, the warning interface **8276** may instruct “NO INPUT NEEDED” and present a warning indicating that the overage threshold is being approached (e.g., “Approaching 10% Limit Warning”). In other aspects, the warning may indicate how many uses remain until the overage threshold is met (e.g., “7 Uses Until 10% Overage Limit Is Met”).

[1591] Next, in view of FIG. **204**, an example system-defined constraint is applied (e.g., by the surgical hub **106** and/or the cloud-based analytics system) to the end effector **8208**. Here, the end effector **8208** associated with identifier **8218** (e.g., SN1a) has a usage parameter of **5** (e.g., 7 uses under its suggested usable life metric of 12 uses remain). As such, in accordance with this example, the system-defined constraint is deemed satisfied and the end effector **8208** is rendered available/usable/authorized. In such an aspect, an interface **8200** may be displayed to show a current status of the end effector **8208** (see FIG. **204**, e.g., “COUNT: **5/12**”). More specifically, in accordance with the example described above, status field **8228** corresponds to the modular adapter **8208** and indicates its current status as “COUNT: **5/12**”. According to other aspects the status field **8228** of the interface **8200** may further show usage remaining, remaining capabilities, and/or compatibility with other connected modular components.

[1592] Yet again, according to one aspect, the interface **8200** may comprise a cloud-based interface (see FIG. **203**, e.g., **8104**) accessible on a cloud-access terminal of the surgical hub (via at least one of a visualization system **108/208** (e.g., FIGS. **1-2**) or a display **135/177** associated with the surgical hub **106** (e.g., FIGS. **1-3**, **7**, etc.)). According to another aspect, the interface **8200** may comprise only a portion(s) of the grid (e.g., the status field **8228**, modular component field **8238**, etc.) accessible directly on the physical end effector **8208** itself via a user interface positioned on the end effector **8208** and/or indirectly on the physical handle **8204** itself via a user interface positioned on the handle **8204**. Here, since the system-defined constraint is satisfied, no warning interface and/or bypass interface is displayed.

[1593] Lastly, still in view of FIG. **204**, an example critical system-defined constraint is applied (e.g., by the surgical hub **106** and/or the cloud-based analytics system) to the staple cartridge **8210**. Here, identifier **8220** (e.g., SN121b), associated with the staple cartridge **8210**, is not a system-recognizable identifier. According to one aspect, this may occur when the surgical hub **206** and/or the cloud-based analytics system (e.g., comprising cloud-based system **205**) is unable to match an identifier (e.g., serial number) received from a modular component with identifiers (e.g., serial numbers) downloaded from the manufacturer(s) of the modular component(s). As such, continuing the example, the system-defined constraint is critical, the system-defined constraint is deemed not satisfied, and the staple cartridge **8210** is rendered unavailable/unusable/unauthorized. Further, as a result, since the critical system-defined constraint cannot be waived, any combined multi-component surgical tool comprising the staple cartridge **8210** may be similarly rendered unavailable/unusable/unauthorized. In such an aspect, an interface **8200** may be displayed to show a current status of the staple cartridge **8210** (see FIG. **204**, e.g., “LOADED” “COUNTERFEIT”). More specifically, in accordance with the example described above, status field **8230** corresponds to the staple cartridge **8210** and indicates its current status as “LOADED” and “COUNTERFEIT”.

[1594] Yet again, according to one aspect, the interface **8200** may comprise a cloud-based interface (see FIG. **203**, e.g., **8104**) accessible on a cloud-access terminal of the surgical hub (via at least one of a visualization system **108/208** (e.g., FIGS. **1-2**) or a display **135/177** associated with the surgical hub **106** (e.g., FIGS. **1-3**, **7**, etc.)). Accord-

ing to another aspect, the interface **8200** may comprise only a portion(s) of the grid (e.g., the status field **8230**, modular component field **8240**, etc.) accessible directly on the physical staple cartridge **8210** itself via a user interface positioned on the staple cartridge **8210** and/or indirectly on the physical handle **8204** itself via a user interface positioned on the handle **8204**. Further, in the context of a critical system-defined constraint, the interface **8200** may visually indicate a warning associated with a modular component (e.g., warning indicator **8250**, e.g., box associated with identifier **8220** highlighted and/or encircled and/or comprises a link **8260** (e.g., “C”) in association with modular component field **8240** of the interface **8200**). In one aspect, the link **8260** (e.g., “C”) may key to a corresponding “Description of Problem” section of the interface **8200** (e.g., “C” “Serial Number of Cartridge Indicates COUNTERFEIT Cartridge”). In another aspect, the link **8260** (e.g., “C”) may be a hyperlink to present the corresponding description (e.g., “C” “Serial Number of Cartridge Indicates COUNTERFEIT Cartridge”) in the interface **8200**. According to such aspects, a portion of the descriptive text (e.g., “COUNTERFEIT”), keyed/hyperlinked via link **8260**, may be a hyperlink/button **8270**. Upon/After selection of the hyperlink/button **8270** an action interface **8280** may be presented in the interface **8200**. According to another aspect, a portion of descriptive text (e.g., COUNTERFEIT) in status field **8230** may be a hyperlink/button **8290** to, upon/after selection, directly present the action interface **8280** in the interface **8200**. Such an aspect may be beneficial/more efficient if the interface **8200** is being presented via a (e.g., smaller) user interface of a modular component (e.g., staple cartridge **8210** and/or handle **8204**). Further, according to such aspects, the interface **8200** may be configured to instruct a user to perform an action (e.g., to remove the staple cartridge **8210** associated with the identifier **8220** (e.g., SN121b) and reload with a staple cartridge associated with a system-recognizable identifier. In one aspect, the action interface **8280** may instruct “ACTION REQUIRED” and present a directive “Remove & Reload”. Here, since the system-defined constraint is critical, no warning interface and/or bypass interface is displayed. In one further aspect, a list of available and/or alternative modular components (e.g., staple cartridges) may be displayed.

[1595] In a similar manner, a list (e.g., black-listed devices) of surgical tools (e.g., wireless surgical devices/instruments **235**) and/or modular components (e.g., handles, modular adapters, end effectors, staple cartridges, etc.) may be declared unavailable/unusable/unauthorized to communicate with and/or access the surgical hub **206** and/or cloud-based analytics system (e.g., comprising cloud-based system **205**). In one aspect of the present disclosure, such black-listed devices may comprise inventory items that are known and/or established to be counterfeit, defective, damaged, beyond their usable life, expired, unsterile, etc. In such an aspect, black-listed devices may be used as critical system-defined constraints (e.g., if the device is on the “black-list,” it cannot communicate with and/or access the surgical hub and/or cloud-based analytics system). In line with above, critical system-defined constraints cannot be waived/bypassed. Creating and/or maintaining such a “black-list” of devices at the surgical hub level and/or the cloud-based analytics level, may improve safety and reliability in the operating room. In one aspect, a database (e.g., aggregated medical data database **7012** in FIG. **180**) at the cloud-based

analytics system may be updated each time a counterfeit device is detected via a surgical hub **206** (e.g., similar to the staple cartridge in FIG. **204**). Since a plurality of surgical hubs associated with a plurality institutions may communicate with the cloud-based analytics system, such a database, and associated “black-list”, builds rather quickly. Such a database at the cloud-based analytics system would prevent a black-listed device from being used at a different surgical hub (e.g., a surgical hub other than the surgical hub at which the counterfeit was initially detected) communicatively coupled to the cloud-based analytics system.

[1596] In another aspect of the present disclosure, black-listed devices may include surgical tools (e.g., wireless surgical devices/instruments **235**) and/or modular components (e.g., handles, modular adapters, end effectors, staple cartridges, etc.) developed by third-parties wishing to take advantage of benefits provided by the surgical hub and/or cloud-based analytics system (e.g., various inventory control aspects discussed herein). In such an aspect of the present disclosure, black-listed devices may be used as non-critical system-defined constraints and/or flexible system-defined constraints (e.g., if the device is on the “black-list,” it cannot communicate with and/or access the surgical hub and/or cloud-based analytics system). However, contrary to the previously disclosed aspect, such non-critical system-defined constraints and/or flexible system-defined constraints may be waived/bypassed. In one aspect of the present disclosure, such a black-listed device (e.g., a third-party device) may be granted access to the surgical hub and/or cloud-based analytics system for a fee. In one example a competitor product may be initially declared counterfeit. However, once an agreed upon fee is paid, that competitor product may be granted access to the surgical hub and/or cloud-based analytics system. In another aspect, such a black-listed device may be granted partial access to the surgical hub and/or cloud-based analytics system but may be subject to established secondary system-defined constraints. In another aspect, such a black-listed device may be granted access to the surgical hub and/or cloud-based analytics system but may not be able to fully function (e.g., limited functionality) when paired with the surgical hub. Similar to above, a database (e.g., aggregated medical data database **7012** in FIG. **180**) at the cloud-based analytics system may be updated each time a previously black-listed device is granted access, partial access with secondary system-defined constraints and/or access with limited functionality. Since a plurality of surgical hubs associated with a plurality institutions may communicate with the cloud-based analytics system, such a database, and its associated access levels, can be implemented across all communicatively coupled surgical hubs. In all such aspects, the surgical hub and/or cloud-based analytics system maintains complete control over devices seeking access.

[1597] In yet another aspect of the present disclosure a database of the surgical hub (see FIG. **10**, e.g., **248/249**) and/or a database (e.g., aggregated medical data database **7012** in FIG. **180**) of the cloud-based analytics system may record each modular component and/or surgical tool identifier (e.g., serial number) in a “used identifier list” when first used in the system. As such, each time a new modular component and/or a new surgical tool is plugged in and/or requests communication with the surgical hub and/or cloud-based analytics system, an identifier of the new modular component and/or surgical tool is cross-checked with the

“used identifier list.” In such an aspect, if the identifier of the new modular component and/or the new surgical tool matches an identifier already in the “used identifier list,” that identifier may be automatically placed on a “black-list” (e.g., critical system-defined constraint). Here, identifiers (e.g., serial numbers) should be unique. If an already used identifier is presented at first use multiple times, this may evidence fraud and/or counterfeit activity.

**[1598]** As discussed herein, various aspects of the present disclosure are directed to the application of system-defined constraints. For example, as discussed with reference to FIG. 204 above, each modular component of a surgical tool may be associated with an identifier and each identifier may be associated with one or more than one parameter (e.g., usage parameter, expiration date, flexible parameter, etc.). In another aspect of the present disclosure, a surgical tool may be associated with an identifier wherein that identifier is associated with one or more than one parameter. In such an aspect, either the surgical tool does not comprise modular components or the surgical tool comprises modular components associated with the same identifier (e.g., serial number, activation code). Here, system-defined constraints, as discussed herein, may be applied to such a surgical tool in a similar manner.

**[1599]** Further, as discussed herein, various aspects of the present disclosure pertain to the identification of reusable/reprocessed devices (e.g., modular components, surgical tools, etc.) and the display of each reusable device’s availability/readiness for a next/proposed surgical procedure and its operational status on a screen other than the screen of the reusable device (e.g., a screen of a cloud-access terminal of the surgical hub). In one aspect of the present disclosure the status of each reusable device (e.g., status of each modular component, status of a surgical tool, and/or overall status of combined modular components and/or subassemblies) is queried and/or determined when the reusable device connects to the system or as the reusable device connects to the system (e.g., to the surgical hub and/or the cloud-based analytics system). In another aspect of the present disclosure, once/after the reusable device is used, the surgical hub and/or cloud-based analytics system time-stamps the use and updates the usage of each modular component and/or surgical tool in its respective database.

**[1600]** In further various aspects of the present disclosure, a modular component and/or a surgical tool may be flagged by the surgical hub and/or cloud based analytics system based on predetermined criteria. For example, if a modular component is incompatible with other modular components, its identifier (e.g., serial number) is known to be fake, and/or it is subject to a recall, a database of the surgical hub and/or the cloud-based analytics system may be updated to not allow use of the modular component and/or surgical tool in the system (e.g., creation of critical system-defined constraints). Such created system-defined constraints may be applied as discussed herein.

**[1601]** In yet further aspects of the present disclosure, a modular component and/or a surgical tool may be flagged by the surgical hub and/or cloud based analytics system based on a previous use. For example, the surgical hub and/or the cloud based analytics system may track performance of the modular component and/or the surgical tool. Here, performance results may be analyzed by the cloud-based analytics system to inform future uses of the modular component and/or surgical tool. For example, if the end effector did not

clamp properly or jammed in a previous use, the end effector may be flagged in a database of the surgical hub and/or the cloud-based analytics system (e.g., black-listed) so that the end effector cannot be used again in the system.

**[1602]** Various aspects of the present disclosure are also directed to a cloud-based analytics system that generates a cloud interface for a client care institution. More specifically, aspects of the present disclosure pertain to a cloud-based system including a client care institution surgical hub coupleable with a plurality of inventory items (e.g., handles, modular adapters, end effectors, staple cartridges, etc.) and a cloud-based analytics system. The surgical hub may include a processor programmed to communicate with the plurality of inventory items and the cloud-based analytics system. The cloud-based analytics system may include a processor programmed to i) receive, via the surgical hub, data associated with the plurality of inventory items, wherein the received data comprises a unique identifier for each inventory item, ii) determine whether each inventory item is available for use based on its respective unique identifier and system-defined constraints, wherein the system-defined constraints comprise at least one use restriction, iii) generate a cloud interface for the institution, wherein the institution’s cloud interface comprises a plurality of user-interface elements, wherein at least one user-interface element enables the institution to select one or more than one surgical procedure to be performed, and wherein after selection of a surgical procedure, via the at least one user-interface element, the availability of each inventory item associated with the selected surgical procedure is dynamically generated on the institution’s cloud interface, and iv) display an alert for each inventory item determined as not available based on the system-defined constraints, wherein the alert is displayable on at least one of the institution’s cloud interface or the inventory item. Here, in line with the disclosure herein, alternative inventory items for unavailable items may also be displayed. Such a cloud interface enables an institution to evaluate whether a desired/proposed surgical procedure can proceed based on current inventories. Here, data at the surgical hub level (e.g., historical local usage) and/or the cloud-based analytics system level (e.g., historical local and/or global usage) may be used to determine combinations of modular components and/or surgical tools usable for the surgical procedure selected via the user-interface element. Furthermore, alternative and/or preferred modular components and/or surgical tools may be recommended for the surgical procedure selected via the user-interface element. Such a recommendation (e.g., best practices) may be based on a statistical analysis of data at the surgical hub level and/or the cloud-based analytics system level. Such a recommendation may or may not be based on current inventory of the institution.

**[1603]** In yet another aspect of the present disclosure, a modular component and/or surgical tool may be a single-use device rather than a reusable and/or reprocessed device. In such an aspect, packaging associated with the single-use device may include a one-time use activation code. In such an aspect, the one-time use activation code may be entered into an activation input field on a cloud interface via the cloud-access terminal of the surgical hub and transmitted to the cloud-based analytics system. Here, upon receipt, the cloud-based analytics system may cross-check the one-time use activation code with a database of one-time use activation codes (e.g., downloaded from a manufacturer) to autho-

size use with the system. If the one-time use activation code matches an unused activation code, the modular component and/or surgical tool is authorized. However, if the one-time use activation code does not match an activation code in the database or the one-time use activation code matches an already used activation code, that one-time use activation code may be placed on a black-list such that the single-use modular component and/or surgical tool is not authorized (e.g., critical system-defined constraint).

#### Examples

**[1604]** Various aspects of the subject matter described herein are set out in the following numbered examples.

**[1605]** Example 1: A method of displaying an operational parameter of a surgical system, the method comprising: receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system; analyzing, by the cloud computing system, the first and the second usage data to correlate the first and the second usage data with surgical outcome data; determining, by the cloud computing system, based on the correlation, a recommended medical resource usage configuration; and displaying, on respective displays on the first and the second subset of surgical hubs, indications of the recommended medical resource usage configuration.

**[1606]** Example 2: The method of Example 1, wherein the surgical outcome data comprises positive outcome data that are determined based on a comparison to an expected outcome.

**[1607]** Example 3: The method of any one of Examples 1-2, wherein the surgical outcome data comprises bleeding event data.

**[1608]** Example 4: The method of any one of Examples 1-3, further comprising: determining, by the cloud computing system, medical product waste data based on the first and the second usage data; and adjusting the recommended medical resource usage configuration based on the medical product waste data, wherein the recommended medical resource usage configuration includes a reduced quantity of a first product.

**[1609]** Example 5: The method of any one of Examples 1-4, further comprising: identifying, by the cloud computing system, a missing medical product required for a surgical procedure to be performed by a surgical instrument of the surgical system, wherein the surgical instrument is communicatively coupled to a surgical hub of the first subset of surgical hubs; and displaying, by the surgical hub, an alert transmitted by the cloud computing system indicating the missing medical product.

**[1610]** Example 6: The method of any one of Examples 1-5, wherein the recommended medical resource usage configuration comprises a recommended staple cartridge type and a staple cartridge color.

**[1611]** Example 7: The method of any one of Examples 1-6, wherein the recommended medical resource usage configuration comprises a recommended adjustment to a procedural practice.

**[1612]** Example 8: A method of improving a surgical system, the method comprising: receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system, wherein the first subset of surgical hubs correspond to a first medical

facility; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system, wherein the second subset of surgical hubs correspond to a second medical facility; analyzing, by the cloud computing system, the first and the second usage data to correlate the first and the second usage data with surgical outcome data; determining, by the cloud computing system, based on the correlation, that the first usage data is correlated with a first number of positive surgical outcomes and the second usage data is correlated with a second number of positive surgical outcomes, wherein the first number of positive surgical outcomes is greater than the second number of positive surgical outcomes; transmitting, by the cloud computing system, the first usage data to the second subset of surgical hubs; and determining, by the second subset of surgical hubs, an improved medical resource usage configuration based on the first usage data.

**[1613]** Example 9: The method of Example 8, wherein the usage data comprises one or more of resources utilized, time, and procedural cost.

**[1614]** Example 10: The method of any one of Examples 8-9, further comprising: transmitting, by the second subset of surgical hubs, a request for additional information to the cloud computing system.

**[1615]** Example 11: The method of any one of Examples 8-10, wherein the surgical outcome data comprises positive outcome data that are determined based on a comparison to an expected outcome.

**[1616]** Example 12: The method of any one of Examples 8-11, wherein the surgical outcome data comprises bleeding event data.

**[1617]** Example 13: The method of any one of Examples 8-12, further comprising: identifying, by the cloud computing system, an error impacting the second usage data.

**[1618]** Example 14: The method of any one of Examples 8-13, further comprising: comparing, by the cloud computing system, the first and the second usage data to a predetermined threshold.

**[1619]** Example 15: The method of any one of Examples 8-14, further comprising: transmitting, by the cloud computing system, a control program update to the second subset of surgical hubs.

**[1620]** Example 16: A method of controlling security of a surgical system, the method comprising: receiving, by a cloud computing system of the surgical system, first usage data, from a first subset of surgical hubs of the surgical system; receiving, by the cloud computing system, second usage data, from a second subset of surgical hubs of the surgical system; analyzing, by the cloud computing system, the first and the second usage data to compare the first and the second usage data with security data; determining, by the cloud computing system, based on the comparison, that the first usage data is indicative of a security irregularity and the second usage data is indicative of an acceptable security status; determining, by the cloud computing system, a change in a security parameter based on the indicated security irregularity; transmitting, by the cloud computing system, the change in the security parameter to the second subset of surgical hubs; and changing, by the second subset of surgical hubs, the security parameter.

**[1621]** Example 17: The method of Example 16, further comprising: comparing, by the cloud computing system, the first usage data to a predetermined threshold to determine the security irregularity; and generating, by the cloud com-

puting system, a security flag and transmitting the security flag to the second subset of surgical hubs.

**[1622]** Example 18: The method of any one of Examples 16-17, wherein the security irregularity is one or more of: a duplicate serial number, incorrect digital signature, and a number of data requests that exceeds a request number threshold.

**[1623]** Example 19: The method of any one of Examples 16-18, further comprising: transmitting, by the cloud computing system, an operational lockout signal to a plurality of surgical instruments of the surgical system, wherein the plurality of surgical instruments are prevented from connecting to the first subset of surgical hubs based on the operational lockout signal.

**[1624]** Example 20: The method of any one of Examples 16-19, further comprising: verifying, by the cloud computing system, an accuracy of a control program update previously transmitted to the second subset of surgical hubs.

**[1625]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1626]** Example 1: A surgical hub configured to transmit generator data associated with a surgical procedure from a generator of the surgical hub to a cloud-based system communicatively coupled to a plurality of surgical hubs, the surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive generator data from the generator, wherein the generator data is structured into a data packet comprising at least two of the following fields: a field that indicates a source of the data; a unique time stamp; a field indicating an energy mode of the generator; a field indicating a power output of the generator; and a field indicating a duration of the power output of the generator; encrypt the generator data; generate a message authentication code based on the generator data; generate a datagram comprising the encrypted generator data, the generated message authentication code, a source identifier and a destination identifier; and transmit the datagram to a cloud-based system, wherein the datagram allows for the cloud-based system to: decrypt the encrypted generator data of the transmitted datagram; verify the integrity of the generator data based on the message authentication code; authenticate the surgical hub as the source of the datagram; and validate a transmission path followed by the datagram between the surgical hub and the cloud-based system.

**[1627]** Example 2: The surgical hub of Example 1, wherein generating the datagram comprises: generating a datagram header, wherein the datagram header is structured to comprise: a field indicating an IP address associated with the surgical hub; and a field indicating an IP address associated with the cloud-based system; and generating a datagram payload, wherein the datagram payload is structured to comprise the encrypted generator data and the generated message authentication code.

**[1628]** Example 3: The surgical hub of Examples 2, wherein the datagram header is further structured to comprise: a field indicating a transmission path designating at least one IP address associated with at least one intermediate network component through which the datagram is to pass as the datagram is transmitted from the IP address associated with the surgical hub to the IP address associated with the cloud-based system.

**[1629]** Example 4: The surgical hub of any one of Examples 1-3, wherein the instructions are further executable by the processor to: receive a receipt message from the cloud-based system in response to the transmitted datagram, wherein the receipt message indicates at least one of: the integrity of the generator data, decrypted from the transmitted datagram, has been verified by the cloud-based system; the surgical hub has been authenticated as the source of the datagram by the cloud-based system; or the transmission path followed by the transmitted datagram between the surgical hub and the cloud-based system has been validated by the cloud-based system.

**[1630]** Example 5: The surgical hub of any one of Examples 1-4, wherein the instructions are further executable by the processor to: send a message to the cloud-based system, wherein the message requests recommendation generator data associated with a particular surgical procedure; receive a response datagram from the cloud-based system, wherein the response datagram comprises encrypted recommendation generator data and a response message authentication code; decrypt the encrypted recommendation generator data of the response datagram, wherein the recommendation generator data is structured into a response data packet comprising at least one of the following fields: a field indicating an energy mode of the generator for the particular surgical procedure; a field indicating a power output of the generator for the particular surgical procedure; or a field indicating a duration of the power output of the generator for the particular surgical procedure; verify the integrity of the recommendation generator data based on the response message authentication code; and send the recommendation generator data to the generator for implementation, via a generator module, during the particular surgical procedure.

**[1631]** Example 6: The surgical hub of Example 5, wherein the recommendation generator data is based on generator data associated with the particular surgical procedure as securely transmitted by the plurality of surgical hubs to the cloud-based system over time.

**[1632]** Example 7: The surgical hub of Example 1, wherein generating the message authentication code comprises: calculating the message authentication code based on a key, a hash function and one of the received generator data or the encrypted generator data.

**[1633]** Example 8: The surgical hub of Example 7, wherein the key is a secret key and the hash algorithm is a message authentication code algorithm, and wherein calculating the message authentication code comprises processing the encrypted generator data through the message authentication code algorithm using the secret key.

**[1634]** Example 9: The surgical hub of any one of Examples 7-8, wherein the key is a secret key and the hash algorithm is a message authentication code algorithm, and wherein calculating the message authentication code comprises processing the received generator data through the message authentication code algorithm using the secret key.

**[1635]** Example 10: The surgical hub of Example 1, wherein encrypting the generator data comprises encrypting the received generator data using a shared secret or a public key associated with the cloud-based system.

**[1636]** Example 11: A surgical hub configured to transmit generator data associated with a surgical procedure from a generator of the surgical hub to a cloud-based system communicatively coupled to a plurality of surgical hubs, the



surgical hub, comprising: a control circuit configured to: receive generator data from the generator, wherein the generator data is structured into a data packet comprising at least two of the following fields: a field that indicates a source of the data; a unique time stamp; a field indicating an energy mode of the generator; a field indicating a power output of the generator; and a field indicating a duration of the power output of the generator; encrypt the generator data; generate a message authentication code based on the generator data; generate a datagram comprising the encrypted generator data, the generated message authentication code, a source identifier and a destination identifier; and transmit the datagram to a cloud-based system, wherein the datagram allows for the cloud-based system to: decrypt the encrypted generator data of the transmitted datagram; verify the integrity of the generator data based on the message authentication code; authenticate the surgical hub as the source of the datagram; and validate a transmission path followed by the datagram between the surgical hub and the cloud-based system.

**[1637]** Example 12: The surgical hub of Example 11, wherein the control circuit is further configured to: send a message to the cloud-based system, wherein the message requests recommendation generator data associated with a particular surgical procedure; receive a response datagram from the cloud-based system, wherein the response datagram comprises encrypted recommendation generator data and a response message authentication code; decrypt the encrypted recommendation generator data of the response datagram, wherein the recommendation generator data is structured into a response data packet comprising at least one of the following fields: a field indicating an energy mode of the generator for the particular surgical procedure; a field indicating a power output of the generator for the particular surgical procedure; or a field indicating a duration of the power output of the generator for the particular surgical procedure; verify the integrity of the recommendation generator data based on the response message authentication code; and send the recommendation generator data to the generator for implementation, via a generator module, during the particular surgical procedure.

**[1638]** Example 13: The surgical hub of any one of Examples 11-12, wherein the recommendation generator data is based on generator data associated with the particular surgical procedure as securely transmitted by the plurality of surgical hubs to the cloud-based system over time.

**[1639]** Example 14: A surgical hub configured to prioritize surgical data associated with a surgical procedure from a surgical device of the surgical hub to a cloud-based system communicatively coupled to a plurality of surgical hubs, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: capture surgical data, wherein the surgical data comprises data associated with the surgical device; time-stamp the captured surgical data; identify a failure event; identify a time period associated with the failure event; isolate failure event surgical data from surgical data not associated with the failure event based on the identified time period; chronologize the failure event surgical data by time-stamp; encrypt the chronologized failure event surgical data; generate a datagram comprising the encrypted failure event surgical data, wherein the datagram is structured to include a field which includes a flag that prioritizes the encrypted failure event surgical data over

other encrypted data of the datagram; transmit the datagram to the cloud-based system, wherein the datagram allows for the cloud-based system to: decrypt the encrypted failure event surgical data; focus analysis on the failure event surgical data rather than surgical data not associated with the failure event; and flag the surgical device associated with the failure event for at least one of: removal from an operating room; return to a manufacturer; future inoperability in the cloud-based system; or a download update to prevent failure events.

**[1640]** Example 15: The surgical hub of Example 14, wherein the surgical device comprises an end effector including a staple cartridge, wherein the captured surgical data comprises snapshots taken via an endoscope of the surgical hub during a stapling portion of a surgical procedure, and wherein identifying the failure event comprises comparing, via an imaging module of the surgical hub, the snapshots to images conveying correctly fired staples to detect at least one of a misfired staple or evidence of a misfired staple.

**[1641]** Example 16: The surgical hub of any one of Examples 14-15, wherein the instructions are further executable by the processor to: download a program from the cloud-based system for execution by the surgical device, wherein execution of the program modifies the surgical device to prevent misfired staples.

**[1642]** Example 17: The surgical hub of any one of Examples 14-16, wherein the surgical device comprises an end effector including a temperature sensor, wherein the captured surgical data comprises at least one temperature detected by the temperature sensor during a tissue sealing portion of a surgical procedure, and wherein identifying the failure event comprises comparing the at least one detected temperature to a temperature or a range of temperatures associated with that surgical procedure to detect an inadequate sealing temperature.

**[1643]** Example 18: The surgical hub of Example 17, wherein the instructions are further executable by the processor to: download a program from the cloud-based system for execution by the surgical device, wherein execution of the program modifies the surgical device to prevent inadequate sealing temperatures.

**[1644]** Example 19: The surgical hub of Example 14, wherein the identified time period includes a period of time prior to the failure event being identified.

**[1645]** Example 20: The surgical hub of any one of Examples 14-18, wherein the instructions are further executable by the processor to: receive an action message from the cloud-based system, wherein the action message indicates the surgical device as flagged for at least one of: removal from the operating room; return to the manufacturer; future inoperability in the cloud-based system; or the download update to prevent failure events; and provide a notification, via at least one of a user interface of the surgical hub or a user interface of the surgical device, to perform an action associated with the action message.

**[1646]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1647]** Example 1: A surgical hub configured to authenticate data communications with surgical devices, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: detect that a surgical device is communica-

tively coupled to the surgical hub; transmit a public key associated with the surgical hub to the surgical device; receive a message from the surgical device, wherein the message is encrypted using the public key associated with the surgical hub, wherein the encrypted message comprises a shared secret associated with the surgical device and a checksum function associated with the shared secret, and wherein the shared secret comprises an identifier assigned to the surgical device; decrypt the encrypted message, using a private key associated with the transmitted public key, to reveal the shared secret and the checksum function; receive data communications from the surgical device, wherein each data communication is encrypted using the shared secret received from the surgical device, and wherein each data communication comprises a checksum value, derived via the checksum function, based on the data of each received communication; and decrypt each data communication using the shared secret until the surgical device is decoupled from the surgical hub, wherein the integrity of each data communication is verifiable based on its associated checksum value.

**[1648]** Example 2: The surgical hub of Example 1, wherein the identifier assigned to the surgical device comprises a unique serial number of the surgical device.

**[1649]** Example 3: The surgical hub of any one of Examples 1-2, wherein the instructions are further executable by the processor to: transmit a message to a cloud-based system communicatively coupled to a plurality of surgical hubs, wherein the message is encrypted using the public key associated with the cloud-based system, wherein the encrypted message comprises the shared secret associated with the surgical device, and wherein the shared secret comprises the identifier assigned to the surgical device; and transmit each data communication received from the surgical device to the cloud-based system, wherein each data communication is encrypted using the shared secret received from the surgical device to allow the cloud-based system to decrypt each data communication using the shared secret until the surgical device is decoupled from the surgical hub.

**[1650]** Example 4: The surgical hub of any one of Examples 1-3, wherein the instructions are further executable by the processor to: detect that a multi-component surgical device comprising a plurality of sub-components is communicatively coupled to the surgical hub, wherein each sub-component is associated with an identifier; transmit a public key associated with the surgical hub to the multi-component surgical device; receive a message from the multi-component surgical device, wherein the message is encrypted using the public key associated with the surgical hub, wherein the encrypted message comprises a shared secret associated with the multi-component surgical device and a checksum function associated with the shared secret, and wherein the shared secret comprises a unique string of the identifiers associated with the plurality of sub-components that combine to form the multi-component surgical device; decrypt the encrypted message, using a private key associated with the transmitted public key, to reveal the shared secret and the checksum function; receive data communications from the multi-component surgical device, wherein each data communication is encrypted using the shared secret received from the multi-component surgical device, and wherein each data communication comprises a checksum value, derived via the checksum function, based on the data of each received communication; and decrypt

each data communication using the shared secret until the multi-component surgical device is decoupled from the surgical hub, wherein the integrity of each data communication is verifiable based on its associated checksum value.

**[1651]** Example 5: The surgical hub of Example 4, wherein the unique string of the identifiers associated with the plurality of sub-components that combine to form the multi-component surgical device comprises a random ordering of the identifiers associated with the plurality of sub-components.

**[1652]** Example 6: The surgical hub of Example 5, wherein each identifier of the unique string of identifiers comprises a unique serial number associated with each respective sub-component of the multi-component surgical device.

**[1653]** Example 7: A surgical hub configured to authenticate data communications with surgical devices, the surgical hub comprising a control circuit configured to: detect that a surgical device is communicatively coupled to the surgical hub; transmit a public key associated with the surgical hub to the surgical device; receive a message from the surgical device, wherein the message is encrypted using the public key associated with the surgical hub, wherein the encrypted message comprises a shared secret associated with the surgical device and a checksum function associated with the shared secret, and wherein the shared secret comprises an identifier assigned to the surgical device; decrypt the encrypted message, using a private key associated with the transmitted public key, to reveal the shared secret and the checksum function; receive data communications from the surgical device, wherein each data communication is encrypted using the shared secret received from the surgical device, and wherein each data communication comprises a checksum value, derived via the checksum function, based on the data of each received communication; and decrypt each data communication using the shared secret until the surgical device is decoupled from the surgical hub, wherein the integrity of each data communication is verifiable based on its associated checksum value.

**[1654]** Example 8: A surgical hub configured to authenticate surgical devices coupled to the surgical hub, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: detect that a surgical device is communicatively coupled to the surgical hub; receive an encrypted identifier and a source ID from the surgical device; transmit a first message from the surgical hub to a server of a surgical device manufacturer associated with the source ID, wherein the first message comprises the encrypted identifier, and wherein the first message is encrypted using a public key associated with the surgical device manufacturer; receive a second message from the server of the surgical device manufacturer, wherein the second message is encrypted using a public key associated with the surgical hub, and wherein the encrypted second message comprises a shared secret associated with the encrypted identifier of the surgical device; decrypt the encrypted second message using a private key associated with the public key used to encrypt the second message to reveal the shared secret associated with the encrypted identifier of the surgical device; and decrypt the encrypted identifier of the surgical device using the shared secret to reveal the identifier to authenticate the surgical device and its manufacturer.

[1655] Example 9: The surgical hub of any one of Example 8, wherein the identifier comprises a unique serial number of the surgical device.

[1656] Example 10: The surgical hub of any one of Examples 8-9, wherein the instructions are further executable by the processor to: compare the decrypted identifier to a list of authorized identifiers; and permit use of the surgical device based on a match of the decrypted identifier to an authorized identifier in the list.

[1657] Example 11: The surgical hub of Example 10, wherein the instructions are further executable by the processor to: download the list of authorized identifiers from a cloud-based system communicatively coupled to a plurality of surgical hubs.

[1658] Example 12: The surgical hub of any one of Examples 8-11, wherein receiving the encrypted identifier and the source ID from the surgical device comprises: reading the encrypted identifier and the source ID from a memory device of the surgical device.

[1659] Example 13: The surgical hub of any one of Examples 8-12, wherein the instructions are further executable by the processor to: read usage data from a memory device of the coupled surgical device; store at least a portion of the read usage data each time the surgical device is coupled to the surgical hub; compare the read usage data to previously stored usage data to identify a discrepancy in the usage data; and prevent usage of the surgical device with the surgical hub based on an identified discrepancy.

[1660] Example 14: The surgical hub of any one of Examples 8-13, wherein the instructions are further executable by the processor to: transmit a record of the coupling of the surgical device and the surgical hub to at least one of a cloud-based system or a plurality of other surgical hubs communicatively coupled to the cloud-based system in a surgical system, wherein the record links the unique identifier assigned to the surgical device to a unique identifier assigned to the surgical hub.

[1661] Example 15: The surgical hub of any one of Examples 8-14, wherein the unique identifier assigned to the surgical device comprises a serial number.

[1662] Example 16: The surgical hub of any one of Examples 8-15, wherein the instructions are further executable by the processor to: store the record of the coupling of the surgical device and the surgical hub as a genesis record, wherein the genesis record comprises a timestamp.

[1663] Example 17: The surgical hub of any one of Examples 8-16, wherein the instructions are further executable by the processor to: store a new record for each subsequent coupling of the surgical device to the surgical hub, wherein each new record in a chain of records associated with the surgical device comprises a cryptographic hash of the most recent record, the linkage of the unique identifier assigned to the surgical device to the unique identifier assigned to the surgical hub, and a timestamp.

[1664] Example 18: The surgical hub of any one of Examples 8-17, wherein the instructions are further executable by the processor to: receive a record of a coupling of the surgical device to one of the plurality of other surgical hubs communicatively coupled to the cloud-based system; and store a new record for the coupling of the surgical device to the one of the plurality of other surgical hubs, wherein the new record in a chain of records associated with the surgical device comprises a cryptographic hash of the most recent record, a linkage of the unique identifier assigned to the

surgical device to a unique identifier assigned to the one of the plurality of other surgical hubs, and a timestamp.

[1665] Example 19: The surgical hub of Example 18, wherein the instructions are further executable by the processor to: trace couplings of the surgical device to the surgical hub and the plurality of other surgical hubs in the surgical system back to the genesis record.

[1666] Example 20: The surgical hub of Example 19, wherein the instructions are further executable by the processor to: analyze the traced couplings to determine whether past usage of the surgical device contributed to or caused a failure event.

[1667] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1668] Example 1: A surgical hub for use with a surgical system in a surgical procedure performed in an operating room, wherein the surgical hub comprises a control circuit configured to: pair the surgical hub with a first device of the surgical system; assign a first identifier to the first device; pair the surgical hub with a second device of the surgical system; assign a second identifier to the second device; and selectively pair the first device with the second device based on perioperative data.

[1669] Example 2: The surgical hub of Example 1, further comprising a storage medium, wherein the control circuit is further configured to store a record indicative of the pairing between the first device and the second device in the storage medium.

[1670] Example 3: The surgical hub of any one of Examples 1-2, wherein the pairing between the first device and the second device defines a communication pathway therebetween.

[1671] Example 4: The surgical hub of any one of Examples 1-3, wherein the pairing between the first device and the second device defines a control pathway for transmitting control actions from the second device to the first device.

[1672] Example 5: The surgical hub of any one of Examples 1-4, wherein the control circuit is further configured to: pair the surgical hub with a third device of the surgical system; assign a third identifier to the third device; sever the pairing between the first device and the second device; and selectively pair the first device with the third device.

[1673] Example 6: The surgical hub of any one of Examples 1-5, wherein the control circuit is further configured to store a record indicative of the pairing between the first device and the third device in the storage medium.

[1674] Example 7: The surgical hub of any one of Examples 1-6, wherein the pairing between the first device and the third device defines a communication pathway therebetween.

[1675] Example 8: The surgical hub of any one of Examples 1-7, wherein the pairing between the first device and the third device defines a control pathway for transmitting control actions from the third device to the first device.

[1676] Example 9: A surgical hub for use with a surgical system in a surgical procedure performed in an operating room, wherein the surgical hub comprises: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: pair the surgical hub with a first device of the surgical system; assign a first identifier to the first device; pair the surgical hub with a

second device of the surgical system; assign a second identifier to the second device; and selectively pair the first device with the second device based on perioperative data.

**[1677]** Example 10: The surgical hub of Example 9, a record indicative of the pairing between the first device and the second device is stored in the memory.

**[1678]** Example 11: The surgical hub of any one of Examples 9-10, wherein the pairing between the first device and the second device defines a communication pathway therebetween.

**[1679]** Example 12: The surgical hub of any one of Examples 9-11, wherein the pairing between the first device and the second device defines a control pathway for transmitting control actions from the second device to the first device.

**[1680]** Example 13: The surgical hub of any one of Examples 9-12, wherein the control circuit is further configured to: pair the surgical hub with a third device of the surgical system; assign a third identifier to the third device; sever the pairing between the first device and the second device; and selectively pair the first device with the third device.

**[1681]** Example 14: The surgical hub of any one of Examples 9-13, wherein a record indicative of the pairing between the first device and the third device is stored in the memory.

**[1682]** Example 15: The surgical hub of any one of Examples 9-14, wherein the pairing between the first device and the third device defines a communication pathway therebetween.

**[1683]** Example 16: The surgical hub of any one of Examples 9-15, wherein the pairing between the first device and the third device defines a control pathway for transmitting control actions from the third device to the first device.

**[1684]** Example 17: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: pair a surgical hub with a first device of a surgical system; assign a first identifier to the first device; pair the surgical hub with a second device of the surgical system; assign a second identifier to the second device; and selectively pair the first device with the second device based on perioperative data.

**[1685]** Example 18: The non-transitory computer readable medium of Example 17, wherein the pairing between the first device and the second device defines a control pathway for transmitting control actions from the second device to the first device.

**[1686]** Example 19: The non-transitory computer readable medium of any one of Examples 17-18, wherein the computer readable instructions, when executed, further cause a machine to: pair the surgical hub with a third device of the surgical system; assign a third identifier to the third device; sever the pairing between the first device and the second device; and selectively pair the first device with the third device.

**[1687]** Example 20: The non-transitory computer readable medium of any one of Examples 17-19, wherein the pairing between the first device and the third device defines a control pathway for transmitting control actions from the third device to the first device.

**[1688]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1689]** Example 1: A surgical hub for use with a surgical system in a surgical procedure performed in an operating room, wherein the surgical hub comprises a control circuit configured to: determine bounds of the operating room; determine devices of the surgical system located within the bounds of the operating room; and pair the surgical hub with the devices of the surgical system located within the bounds of the operating room.

**[1690]** Example 2: The surgical hub of any one of Example 1, wherein the step of determining devices of the surgical system comprises: detecting a potential device of the surgical system; and assessing whether the potential device of the surgical system is within the bounds of the operating room or outside the bounds of the operating room.

**[1691]** Example 3: The surgical hub of any one of Examples 1-2, wherein the control circuit is configured to determine the bounds of the operating room after activation of the surgical hub.

**[1692]** Example 4: The surgical hub of any one of Examples 1-3, wherein the control circuit is configured to redetermine the bounds of the operating room after determining that the surgical hub has been moved.

**[1693]** Example 5: The surgical hub of any one of Examples 1-4, wherein the control circuit is configured to redetermine the bounds of the operating room after a potential device of the surgical system is detected.

**[1694]** Example 6: The surgical hub of any one of Examples 1-5, wherein the control circuit is configured to periodically determine the bounds of the operating room.

**[1695]** Example 7: The surgical hub of any one of Examples 1-6, comprising non-contact sensors configured to measure the bounds of the operating room.

**[1696]** Example 8: A surgical hub for use with a surgical system in a surgical procedure performed in an operating room, wherein the surgical hub comprises: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: determine bounds of the operating room; determine devices of the surgical system located within the bounds of the operating room; and pair the surgical hub with the devices of the surgical system located within the bounds of the operating room.

**[1697]** Example 9: The surgical hub of Example 8, wherein the step of determining devices of the surgical system comprises: detecting a potential device of the surgical system; and assessing whether the potential device of the surgical system is within the bounds of the operating room or outside the bounds of the operating room.

**[1698]** Example 10: The surgical hub of any one of Examples 8-9, wherein the memory further stores instructions executable by the processor to determine the bounds of the operating room after activation of the surgical hub.

**[1699]** Example 11: The surgical hub of any one of Examples 8-10, wherein the memory further stores instructions executable by the processor to redetermine the bounds of the operating room after determining that the surgical hub has been moved.

**[1700]** Example 12: The surgical hub of any one of Examples 8-11, wherein the memory further stores instructions executable by the processor to redetermine the bounds of the operating room after a potential device of the surgical system is detected.

**[1701]** Example 13: The surgical hub of any one of Examples 8-12, wherein the memory further stores instruc-

tions executable by the processor to periodically determine the bounds of the operating room.

**[1702]** Example 14: The surgical hub of any one of Examples 8-13, comprising non-contact sensors configured to measure the bounds of the operating room.

**[1703]** Example 15: A non-transitory computer readable medium storing computer readable instructions which, when executed, cause a machine to: determine bounds of an operating room; determine devices of a surgical system located within the bounds of the operating room; and pair a surgical hub with the devices of the surgical system located within the bounds of the operating room.

**[1704]** Example 16: The non-transitory computer readable medium of Example 15, wherein the step of determining devices of the surgical system comprises: detecting a potential device of the surgical system; and assessing whether the potential device of the surgical system is within the bounds of the operating room or outside the bounds of the operating room.

**[1705]** Example 17: The non-transitory computer readable medium of any one of Examples 15-16, wherein the computer readable instructions, when executed, further cause a machine to determine the bounds of the operating room after activation of the surgical hub.

**[1706]** Example 18: The non-transitory computer readable medium of any one of Examples 15-17, wherein the computer readable instructions, when executed, further cause a machine to redetermine the bounds of the operating room after determining that the surgical hub has been moved.

**[1707]** Example 19: The non-transitory computer readable medium of any one of Examples 15-18, wherein the computer readable instructions, when executed, further cause a machine to redetermine the bounds of the operating room after a potential device of the surgical system is detected.

**[1708]** Example 20: The non-transitory computer readable medium of any one of Examples 15-19, wherein the computer readable instructions, when executed, further cause a machine to periodically determine the bounds of the operating room.

**[1709]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1710]** Example 1: A surgical hub for use with a medical imaging device at a remote surgical site in a surgical procedure, wherein the surgical hub comprises a circuit configured to: receive a livestream of the surgical site from the medical imaging device; capture an image frame of a surgical step of the surgical procedure from the livestream; derive information relevant to the surgical step from data extracted from the image frame; and overlay the information onto the livestream.

**[1711]** Example 2: The surgical hub of Example 1, wherein the information is regarding completion of the surgical step.

**[1712]** Example 3: The surgical hub of any one of Examples 1-2, wherein the surgical step comprises deploying staples from a staple cartridge into tissue at the surgical site.

**[1713]** Example 4: The surgical hub of any one of Examples 1-3, wherein the information identifies the staple cartridge.

**[1714]** Example 5: The surgical hub of any one of Examples 1-4, wherein the information comprises a serial number of the staple cartridge.

**[1715]** Example 6: The surgical hub of any one of Examples 1-5, wherein the information identifies a leak at the surgical site.

**[1716]** Example 7: The surgical hub of any one of Examples 1-7, wherein the information identifies the surgical step.

**[1717]** Example 8: A surgical hub for use with a medical imaging device at a remote surgical site in a surgical procedure including surgical steps, wherein the surgical hub comprises a circuit configured to: receive a livestream of the surgical site from the medical imaging device; capture image frames of the surgical steps of the surgical procedure from the livestream; and differentiate among the surgical steps based on data extracted from the image frames.

**[1718]** Example 9: The surgical hub of Example 8, derive information regarding completion of the surgical steps from the data extracted from the image frames.

**[1719]** Example 10: The surgical hub of any one of Examples 8-9, wherein one of the surgical steps comprises deploying staples from a staple cartridge into tissue at the surgical site.

**[1720]** Example 11: The surgical hub of any one of Examples 8-10, wherein the information identifies the staple cartridge.

**[1721]** Example 12: The surgical hub of any one of Examples 8-11, wherein the information comprises a serial number of the staple cartridge.

**[1722]** Example 13: The surgical hub of any one of Examples 8-12, wherein the information identifies a leak at the surgical site.

**[1723]** Example 14: The surgical hub of any one of Examples 8-10, wherein another one of the surgical steps comprises applying energy to tissue at the surgical site.

**[1724]** Example 15: A surgical hub for use with a medical imaging device at a remote surgical site in a surgical procedure, wherein the surgical hub comprises a circuit configured to: receive a livestream of the surgical site from the medical imaging device; capture an image frame from the livestream; detect a staple pattern in the image frame, wherein the staple pattern is defined by staples deployed from a staple cartridge into tissue at the surgical site; and identify the staple cartridge based on the staple pattern.

**[1725]** Example 16: The surgical hub of Example 15, wherein the staple pattern corresponds to a serial number of the staple cartridge.

**[1726]** Example 17: The surgical hub of any one of Examples 15-16, wherein the staples comprise a first staple and a second staple different than the first staple.

**[1727]** Example 18: The surgical hub of any one of Examples 15-17, wherein the first staple is comprised of a non-ionized material.

**[1728]** Example 19: The surgical hub of any one of Examples 15-18, wherein the second staple is comprised of an ionized material.

**[1729]** Example 20: The surgical hub of any one of Examples 15-19, wherein the staple pattern is defined in a plurality of rows of staples.

**[1730]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1731]** Example 1: A surgical hub for use with a surgical system in a surgical procedure performed in an operating room, wherein the surgical hub comprises: non-contact sensors; and a control circuit configured to: determine

bounds of the operating room based on measurements performed by the non-contact sensors; and establish a control arrangement with a detected surgical hub located within the bounds of the operating room.

[1732] Example 2: The surgical hub of Example 1, wherein the control arrangement is a master-slave arrangement.

[1733] Example 3: The surgical hub of any one of Examples 1-2, wherein the control circuit is configured to select one of a master mode of operation or a slave mode of operation in the master-slave arrangement.

[1734] Example 4: The surgical hub of any one of Examples 1-3, wherein the control circuit is configured to surrender control of at least one surgical instrument to the detected surgical hub in the slave mode of operation.

[1735] Example 5: The surgical hub of any one of Examples 1, wherein the control arrangement is a peer-to-peer arrangement.

[1736] Example 6: The surgical hub of Example 1-5, wherein the non-contact sensors are ultrasonic sensors.

[1737] Example 7: The surgical hub of Example 1-5, wherein the non-contact sensors are laser sensors.

[1738] Example 8: A surgical hub for use with a surgical system in a surgical procedure performed in an operating room, wherein the surgical hub comprises: non-contact sensors; a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: determine bounds of the operating room based on measurements performed by the non-contact sensors; and establish a control arrangement with a detected surgical hub located within the bounds of the operating room.

[1739] Example 9: The surgical hub of Example 8, wherein the control arrangement is a master-slave arrangement.

[1740] Example 10: The surgical hub of Example 9, wherein the control circuit is configured to select one of a master mode of operation or a slave mode of operation in the master-slave arrangement.

[1741] Example 11: The surgical hub of Example 10, wherein the control circuit is configured to surrender control of at least one surgical instrument to the detected surgical hub in the slave mode of operation.

[1742] Example 12: The surgical hub of Example 11, wherein the control arrangement is a peer-to-peer arrangement.

[1743] Example 13: The surgical hub of any one of Examples 8-12, wherein the non-contact sensors are ultrasonic sensors.

[1744] Example 14: The surgical hub of Example 8, wherein the non-contact sensors are laser sensors.

[1745] Example 15: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: determine bounds of an operating room; and establish a control arrangement with a detected surgical hub located within the bounds of the operating room.

[1746] Example 16: The non-transitory computer readable medium of Example 15, wherein the control arrangement is a master-slave arrangement.

[1747] Example 17: The non-transitory computer readable medium of any one of Examples 15-16, wherein the computer readable instructions, when executed, further causes the machine to select one of a master mode of operation or a slave mode of operation in the master-slave arrangement.

[1748] Example 18: The non-transitory computer readable medium of any one of Examples 15-17, wherein the computer readable instructions, when executed, further causes the machine to surrender control of at least one surgical instrument to the detected surgical hub in the slave mode of operation.

[1749] Example 19: The non-transitory computer readable medium of any one of Examples 15-18, wherein the control arrangement is a peer-to-peer arrangement.

[1750] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1751] Example 1: A surgical hub, comprising: a processor; a memory coupled to the processor, the memory storing instructions executable by the processor to: interrogate a modular device coupled to the processor via a modular communication hub, wherein the modular device is a source of data sets that include patient identity data and surgical procedure data; receive a data set from the modular device; discard the patient identity data and any portion of the surgical procedure data that identifies the patient from the data set; extract anonymous data from the data set and create an anonymized data set; and configure operation of the surgical hub or the modular device based on the anonymized data set.

[1752] Example 2: The surgical hub of Example 1, wherein the anonymized data set includes catastrophic failure of a modular device, and wherein the memory stores instructions executable by the processor to initiate automatic archive and submission of data for implications analysis based on the catastrophic failure of the modular device.

[1753] Example 3: The surgical hub of any one of Examples 1-2, wherein the memory stores instructions executable by the processor to detect counterfeit component information from the anonymized data set.

[1754] Example 4: The surgical hub of any one of Examples 1-3, wherein the memory stores instructions executable by the processor to derive implications of the modular device from the anonymized data set.

[1755] Example 5: The surgical hub of any one of Examples 1-4, wherein the memory stores instructions executable by the processor to configure the modular device to operate based on the derived implications.

[1756] Example 6: The surgical hub of any one of Examples 1-5, wherein the memory stores instructions executable by the processor to configure the surgical hub based on the derived implications.

[1757] Example 7: The surgical hub of any one of Examples 1-6, wherein the memory stores instructions executable by the processor to conglomerate the anonymized data.

[1758] Example 8: The surgical hub of any one of Examples 1-7, wherein the memory stores instructions executable by the processor to extract the anonymized data prior to storing the received data in a storage device coupled to the surgical hub.

[1759] Example 9: The surgical hub of any one of Examples 1-8, wherein the memory stores instructions executable by the processor to: transmit the anonymized data to a remote network outside of the surgical hub; compile the anonymized data at the remote network; and store a copy of the data set from the modular device in a patient electronic medical records database.

[1760] Example 10: The surgical hub of any one of Examples 1-9, comprising a modular communication hub coupled to the processor, the modular communication hub configured to connect modular devices located in one or more operating theaters to the surgical hub.

[1761] Example 11: A method of stripping data originating from a modular device coupled to a surgical hub by a communication hub, the surgical hub comprising a processor and a memory coupled to the processor, the memory storing instructions executable by the processor, the method comprising: interrogating, by a processor, a modular device coupled to the processor via a modular communication hub, wherein the modular communication hub is configured to connect modular devices located in one or more operating theaters to a surgical hub, wherein the modular device is a source of data sets that include patient identity data and surgical procedure data; receiving, by the processor, a data set from the modular device by the processor via the communication hub; discarding, by the processor, the patient identity data and any portion of the surgical procedure data that identifies the patient from the data set; extracting, by the processor, anonymous data from the data set and create an anonymized data set; and configuring, by the processor, operation of the surgical hub or the modular device based on the anonymized data set.

[1762] Example 12: The method of Example 11, comprising: initiating, by the processor, automatic archive and submission of data for implications analysis based on the catastrophic failure of the modular device wherein the anonymized data set includes catastrophic failure of a modular device.

[1763] Example 13: The method of any one of Examples 11-12, comprising by the processor, detecting counterfeit component information from the anonymized data set.

[1764] Example 14: The method of any one of Examples 11-13, comprising deriving implications of the modular device from the anonymized data set.

[1765] Example 15: The method of Example 14, comprising configuring, by the processor, the modular device to operate based on the derived implications.

[1766] Example 16: The method of any one of Examples 14-15, comprising configuring, by the processor, the surgical hub based on the derived implications.

[1767] Example 17: The method of any one of Examples 14-16, comprising, conglomerating by the processor, the anonymized data.

[1768] Example 18: The method of Example 11, extracting, by the processor, the anonymized data prior to storing the received data in a storage device coupled to the surgical hub.

[1769] Example 19: The method of Example 11, comprising: transmitting, by the processor, the anonymized data to a remote network outside of the surgical hub; compiling, by a server at the remote network, the anonymized data at the remote network; and storing, by the processor, a copy of the data set from the modular device in a patient electronic medical records database.

[1770] Example 20: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: interrogate a modular device coupled to the processor via the modular communication hub, wherein the modular device is a source of data sets that include patient identity data and surgical procedure data; receive a data set from the modular device; discard the

patient identity data and any portion of the surgical procedure data that identifies the patient from the data set; extract anonymous data from the data set and create an anonymized data set; and configure operation of the surgical hub or the modular device based on the anonymized data set.

[1771] Example 21: The non-transitory computer readable medium of Example 20, storing computer readable instructions which, when executed, causes a machine to derive implications of the modular device from the anonymized data set.

[1772] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1773] Example 1: A surgical hub comprising: a storage device; a processor coupled to the storage device; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive data from a surgical instrument coupled to the surgical hub; and determine a rate at which to transfer the data from the surgical hub to a remote cloud-based medical analytics network based on available storage capacity of the storage device.

[1774] Example 2: The surgical hub of Example 1, wherein the memory stores instructions executable by the processor to determine a frequency at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on the available storage capacity of the storage device.

[1775] Example 3: The surgical hub of any one of Examples 1-2, wherein the memory stores instructions executable by the processor to: detect surgical hub network down time; and determine a frequency at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on the detected surgical hub network down time.

[1776] Example 4: The surgical hub of any one of Examples 1-3, wherein the memory stores instructions executable by the processor to determine a type of data to transfer from the surgical hub to the remote cloud-based medical analytics network based on inclusion or exclusion of data associated with a users, patient, or surgical procedure.

[1777] Example 5: The surgical hub of any one of Examples 1-4, wherein the memory stores instructions executable by the processor to determine when to transfer data from the surgical hub to the remote cloud-based medical analytics network.

[1778] Example 6: The surgical hub of any one of Examples 1-5, wherein the memory stores instructions executable by the processor to receive new operational parameters for the surgical hub from the remote cloud-based medical analytics network.

[1779] Example 7: The surgical hub of any one of Examples 1-6, wherein the memory stores instructions executable by the processor to receive new operational parameters for the surgical instrument from the remote cloud-based medical analytics network.

[1780] Example 8: A method of transmitting data from a surgical hub to a remote cloud-based medical analytics network, the surgical hub comprising a storage device, a processor coupled to the storage device, and a memory coupled to the processor, the memory storing instructions executable by the processor, the method comprising: receiving, by a processor, data from a surgical instrument coupled

to the surgical hub; and determining, by the processor, a rate at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on available storage capacity of a storage device coupled to the surgical hub.

[1781] Example 9: The method of Example 8, comprising determining, by the processor, a frequency at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on the available storage capacity of the storage device

[1782] Example 10: The method of any one of Examples 8-9, comprising: detecting, by the processor, surgical hub network down time; and determining, by the processor, a frequency at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on the detected surgical hub network down time.

[1783] Example 11: The method of any one of Examples 8-10, comprising determining, by the processor, a type of data to transfer from the surgical hub to the remote cloud-based medical analytics network based on inclusion or exclusion of data associated with a users, patient, or surgical procedure.

[1784] Example 12: The method of any one of Examples 8-11, comprising determining, by the processor, when to transfer the data from the surgical hub to the remote cloud-based medical analytics network.

[1785] Example 13: The method of any one of Examples 8-12, comprising receiving, by the processor, new operational parameters for the surgical hub from the remote cloud-based medical analytics network.

[1786] Example 14: The method of any one of Examples 8-13, comprising receiving, by the processor, new operational parameters for the surgical instrument from the remote cloud-based medical analytics network.

[1787] Example 15: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: receive data from a surgical instrument coupled to the surgical hub; and determine a rate at which to transfer the data from the surgical hub to a remote cloud-based medical analytics network based on available storage capacity of the storage device.

[1788] Example 16: The non-transitory computer readable medium of Example 15, storing computer readable instructions which, when executed, causes a machine to determine a frequency at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on the available storage capacity of the storage device.

[1789] Example 17: The non-transitory computer readable medium of any one of Examples 15-16, storing computer readable instructions which, when executed, causes a machine to: detect surgical hub network down time; and determine a frequency at which to transfer the data from the surgical hub to the remote cloud-based medical analytics network based on the detected surgical hub network down time.

[1790] Example 18: the non-transitory computer readable medium of any one of Examples 15-17, storing computer readable instructions which, when executed, causes a machine to determine a type of data to transfer from the surgical hub to the remote cloud-based medical analytics network based on inclusion or exclusion of data associated with a users, patient, or surgical procedure.

[1791] Example 19: The non-transitory computer readable medium of any one of Examples 15-18, storing computer

readable instructions which, when executed, causes a machine to determine when to transfer data from the surgical hub to the remote cloud-based medical analytics network.

[1792] Example 20: The non-transitory computer readable medium of any one of Examples 15-19, storing computer readable instructions which, when executed, causes a machine to receive new operational parameters for the surgical hub from the remote cloud-based medical analytics network.

[1793] Example 21: The non-transitory computer readable medium of any one of Examples 15-20, storing computer readable instructions which, when executed, causes a machine to receive new operational parameters for the surgical instrument from the remote cloud-based medical analytics network.

[1794] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1795] Example 1: A surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive a first self-describing data packet from a first data source, the first self-describing data packet comprising a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate, wherein the first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet; parse the received first preamble; and interpret the first data payload based on the first preamble.

[1796] Example 2: The surgical hub of Example 1, wherein the memory stores instructions executable by the processor to: receive a second self-describing data packet from a second data source, the second self-describing data packet comprising a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate, wherein the second preamble defines the second data payload and the second encryption certificate verifies the authenticity of the second data packet; parse the received second preamble; interpret the second data payload based on the second preamble; associate the first and second self-describing data packets; and generate a third self-describing data packet comprising the first and second data payloads.

[1797] Example 3: The surgical hub of any one of Examples 1-2, wherein the memory stores instructions executable by the processor to anonymize the data payload of the third self-describing data packet.

[1798] Example 4: The surgical hub of any one of Examples 1-3, wherein the memory stores instructions executable by the processor to: determine that a data payload was generated by a new data source; verify the new data source of the data payload; and alter a data collection process at the surgical hub to receive subsequent data packets from the new data source.

[1799] Example 5: The surgical hub of any one of Examples 1-4, wherein the memory stores instructions executable by the processor to associate the first and second self-describing data packets based on a key.

[1800] Example 6: The surgical hub of any one of Examples 1-5, wherein the memory stores instructions executable by the processor to: receive an anonymized third self-describing data packet; and reintegrate the anonymized third self-describing data packet into the first and second self-describing data packets using the key.



**[1801]** Example 7: A surgical hub, comprising: a control circuit configured to: receive a first self-describing data packet from a first data source, the first self-describing data packet comprising a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate, wherein the first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet; parse the received first preamble; and interpret the first data payload based on the first preamble.

**[1802]** Example 8: The surgical hub of Example 7, wherein the control circuit is further configured: receive a second self-describing data packet from a second data source, the second self-describing data packet comprising a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate, wherein the second preamble defines the second data payload and the second encryption certificate verifies the authenticity of the second data packet; parse the received second preamble; interpret the second data payload based on the second preamble; associate the first and second self-describing data packets; and generate a third self-describing data packet comprising the first and second data payloads.

**[1803]** Example 9: The surgical hub of any one of Examples 7-8, wherein the control circuit is further configured to anonymize the data payload of the third self-describing data packet.

**[1804]** Example 10: The surgical hub of any one of Examples 7-9, wherein the control circuit is further configured to: determine that a data payload was generated by a new data source; verify the new data source of the data payload; and alter a data collection process at the surgical hub to receive subsequent data packets from the new data source.

**[1805]** Example 11: The surgical hub of any one of Examples 7-10, wherein the control circuit is further configured to associate the first and second self-describing data packets based on a key.

**[1806]** Example 12: The surgical hub of any one of Examples 7-11, wherein the control circuit is further configured to: receive an anonymized third self-describing data packet; and reintegrate the anonymized third self-describing data packet into the first and second self-describing data packets using the key.

**[1807]** Example 13: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: receive a first self-describing data packet from a first data source, the first self-describing data packet comprising a first preamble, a first data payload, a source of the first data payload, and a first encryption certificate, wherein the first preamble defines the first data payload and the first encryption certificate verifies the authenticity of the first data packet; parse the received first preamble; and interpret the first data payload based on the first preamble.

**[1808]** Example 14: The non-transitory computer-readable medium of Example 13, storing computer readable instructions which, when executed, causes a machine to: receive a second self-describing data packet from a second data source, the second self-describing data packet comprising a second preamble, a second data payload, a source of the second data payload, and a second encryption certificate, wherein the second preamble defines the second data payload and the second encryption certificate verifies the

authenticity of the second data packet; parse the received second preamble; interpret the second data payload based on the second preamble; associate the first and second self-describing data packets; and generate a third self-describing data packet comprising the first and second data payloads.

**[1809]** Example 15: The non-transitory computer-readable medium of any one of Examples 13-14, storing computer readable instructions to anonymize the data payload of the third self-describing data packet.

**[1810]** Example 16: The non-transitory computer-readable medium of any one of Examples 13-15, storing computer readable instructions to: determine that a data payload was generated by a new data source; verify the new data source of the data payload; and alter a data collection process at the surgical hub to receive subsequent data packets from the new data source.

**[1811]** Example 17: The non-transitory computer-readable medium of any one of Examples 13-16, storing computer readable instructions to associate the first and second self-describing data packets based on a key.

**[1812]** Example 18: The non-transitory computer-readable medium of any one of Examples 13-17, storing computer readable instructions to: receive an anonymized third self-describing data packet; and reintegrate the anonymized third self-describing data packet into the first and second self-describing data packets using the key.

**[1813]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1814]** Example 1: A surgical hub configured to communicate with a surgical instrument, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive a first data set associated with a surgical procedure, wherein the first data set is generated at a first time; receive a second data set associated with the efficacy of the surgical procedure, wherein the second data set is generated at a second time, wherein the second time is separate and distinct from the first time; anonymize the first and second data sets by removing information that identifies a patient, a surgery, or a scheduled time of the surgery; and store the first and second anonymized data sets to generate a data pair grouped by surgery.

**[1815]** Example 2: The surgical hub of Example 1, wherein the memory stores instructions executable by the processor to reconstruct a series of chronological events based on the data pair.

**[1816]** Example 3: The surgical hub of any one of Examples 1-2, wherein the memory stores instructions executable by the processor to reconstruct a series of coupled but unconstrained data sets based on the data pair.

**[1817]** Example 4: The surgical hub of any one of Examples 1-3, wherein the memory stores instructions executable by the processor to: encrypt the data pair; define a backup format for the data pair; and mirror the data pair to a cloud storage device.

**[1818]** Example 5: A surgical hub configured to communicate with a surgical instrument, the surgical hub comprising: a control circuit configured to: receive a first data set associated with a surgical procedure, wherein the first data set is generated at a first time; receive a second data set associated with the efficacy of the surgical procedure, wherein the second data set is generated at a second time, wherein the second time is separate and distinct from the

first time; anonymize the first and second data sets by removing information that identifies a patient, a surgery, or a scheduled time of the surgery; and store the first and second anonymized data sets to generate a data pair grouped by surgery.

[1819] Example 6: The surgical hub of Example 5, wherein the control circuit is further configured to reconstruct a series of chronological events based on the data pair.

[1820] Example 7: The surgical hub of any one of Examples 5-6, wherein the control circuit is further configured to reconstruct a series of coupled but unconstrained data sets based on the data pair.

[1821] Example 8: The surgical hub of any one of Examples 5-7, wherein the control circuit is further configured to: encrypt the data pair; define a backup format for the data pair; and mirror the data pair to a cloud storage device.

[1822] Example 9: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: receive a first data set associated with a surgical procedure, wherein the first data set is generated at a first time; receive a second data set associated with the efficacy of the surgical procedure, wherein the second data set is generated at a second time, wherein the second time is separate and distinct from the first time; anonymize the first and second data sets by removing information that identifies a patient, a surgery, or a scheduled time of the surgery; and store the first and second anonymized data sets to generate a data pair grouped by surgery.

[1823] Example 10: The non-transitory computer-readable medium of Example 9, storing computer readable instructions which, when executed, causes a machine to reconstruct a series of chronological events based on the data pair.

[1824] Example 11: The surgical hub of any one of Examples 9-10, storing computer readable instructions which, when executed, causes a machine to reconstruct a series of coupled but unconstrained data sets based on the data pair.

[1825] Example 12: The surgical hub of any one of any one of Examples 9-11, storing computer readable instructions which, when executed, causes a machine to: encrypt the data pair; define a backup format for the data pair; and mirror the data pair to a cloud storage device.

[1826] Example 13: A surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: interrogate a surgical instrument, wherein the surgical instrument is a first source of patient data; retrieve a first data set from the surgical instrument, wherein the first data set is associated with a patient and a surgical procedure; interrogate a medical imaging device, wherein the medical imaging device is a second source of patient data; retrieve a second data set from the medical imaging device, wherein the second data set is associated with the patient and an outcome of the surgical procedure; associate the first and second data sets by a key; and transmit the associated first and second data sets to remote network outside of the surgical hub.

[1827] Example 14: The surgical hub of Example 13, wherein the memory stores instructions executable by the processor to: retrieve the first data set using the key; anonymize the first data set by removing patient information from the first data set; retrieve the second data set using the key; anonymize the second data set by removing patient information from the second data set; pair the anonymized

first and second data sets; and determine success rates of surgical procedures grouped by the surgical procedure based on the anonymized paired first and second data sets.

[1828] Example 15: The surgical hub of any one of Examples 13-14, wherein the memory stores instructions executable by the processor to: retrieve the anonymized first data set; retrieve the anonymized second data set; and reintegrate the anonymized first and second data sets using the key.

[1829] Example 16: The surgical hub of any one of Examples 13-15, wherein the first and second data sets define first and second data payloads in respective first and second data packets.

[1830] Example 17: The surgical hub of any one of Examples 13-16, wherein the memory stores instructions executable by the processor to retrieve information from an electronic medical records database.

[1831] Example 18: The surgical hub of any one of Examples 13-17, wherein the memory stores instructions executable by the processor to anonymize the information retrieved from the electronic medical records database by removing patient information from the information retrieved from the electronic medical records database.

[1832] Various additional aspects of the subject matter described herein are set out in the following numbered examples:

[1833] Example 1: A surgical hub configured to communicably couple to a data source and a modular device, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions that, when executed by the processor, cause the surgical hub to: receive perioperative data from the data source, wherein the perioperative data comprises data detected by the data source during the course of a surgical procedure; determine contextual information regarding the surgical procedure according to the perioperative data; determine control adjustments for the modular device according to the contextual information; and control the modular device according to the control adjustments.

[1834] Example 2: The surgical hub of any one of Example 1, wherein the data source comprises a first modular device and the modular device comprises a second modular device.

[1835] Example 3: The surgical hub of any one of Examples 1-2, wherein the data source comprises a patient monitoring device.

[1836] Example 4: The surgical hub of any one of Examples 1-3, wherein the contextual information comprises a procedural type of the surgical procedure.

[1837] Example 5: The surgical hub of any one of Examples 1-4, wherein the contextual information comprises a procedural step of the surgical procedure.

[1838] Example 6: The surgical hub of any one of Examples 1-5, wherein the perioperative data comprises a parameter associated with the modular device.

[1839] Example 7: The surgical hub of any one of Examples 1-6, wherein the perioperative data comprises a parameter associated with a patient.

[1840] Example 8: A surgical hub configured to communicably couple to a data source and a modular device, the surgical hub comprising a control circuit configured to receive perioperative data from the data source, wherein the perioperative data comprises data detected by the data source during the course of a surgical procedure; determine

contextual information regarding the surgical procedure according to the perioperative data; determine control adjustments for the modular device according to the contextual information; and control the modular device according to the control adjustments.

**[1841]** Example 9: The surgical hub of any one of Example 8, wherein the data source comprises a first modular device and the modular device comprises a second modular device.

**[1842]** Example 10: The surgical hub of any one of Examples 8-9, wherein the data source comprises a patient monitoring device.

**[1843]** Example 11: The surgical hub of any one of Examples 8-10, wherein the contextual information comprises a procedural type of the surgical procedure.

**[1844]** Example 12: The surgical hub of any one of Examples 8-11, wherein the contextual information comprises a procedural step of the surgical procedure.

**[1845]** Example 13: The surgical hub of any one of Examples 8-12, wherein the perioperative data comprises a parameter associated with the modular device.

**[1846]** Example 14: The surgical hub of any one of Examples 8-13, wherein the perioperative data comprises a parameter associated with a patient.

**[1847]** Example 15: A non-transitory computer readable medium storing computer readable instructions thereon that, when executed by a surgical hub configured to communicably couple to a data source and a modular device, causes the surgical hub to receive perioperative data from the data source, wherein the perioperative data comprises data detected by the data source during the course of a surgical procedure; determine contextual information regarding the surgical procedure according to the perioperative data; determine control adjustments for the modular device according to the contextual information; and control the modular device according to the control adjustments.

**[1848]** Example 16: The surgical hub of any one of Examples 15, wherein the data source comprises a first modular device and the modular device comprises a second modular device.

**[1849]** Example 17: The surgical hub of any one of Examples 15-16, wherein the data source comprises a patient monitoring device.

**[1850]** Example 18: The surgical hub of any one of Examples 15-17, wherein the contextual information comprises a procedural type of the surgical procedure.

**[1851]** Example 19: The surgical hub of any one of Examples 15-18, wherein the contextual information comprises a procedural step of the surgical procedure.

**[1852]** Example 20: The surgical hub of any one of Examples 15-19, wherein the perioperative data comprises a parameter associated with the modular device.

**[1853]** Example 21: The surgical hub of any one of Examples 15-20, wherein the perioperative data comprises a parameter associated with a patient.

**[1854]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1855]** Example 1: A system comprising a surgical hub configured to communicably couple to a modular device comprising a sensor configured to detect data associated with the modular device and a device processor, the surgical hub comprising: a hub processor; and a hub memory coupled to the hub processor; and a distributed control

system executable at least in part by each of the device processor and the hub processor, the distributed control system configured to: receive the data detected by the sensor; determine control adjustments for the modular device according to the data; and control the modular device according to the control adjustments; wherein in a first mode the distributed control system is executed by both the hub processor and the device processor, and in a second mode the distributed control system is executed solely by the device processor.

**[1856]** Example 2: The system of any one of Examples 1, wherein the distributed control system is configured to shift from the first mode to the second mode when a sampling rate of the sensor is greater than a data transmission rate from the modular device to the surgical hub.

**[1857]** Example 3: The system of any one of Examples 1-2, wherein the distributed control system is configured to shift from the second mode to the first mode when a sampling rate of the sensor is less than a data transmission rate from the modular device to the surgical hub.

**[1858]** Example 4: The system of any one of Examples 1-3, wherein the modular device comprises a radiofrequency (RF) electrosurgical instrument and the distributed control system is configured to control an energy level of the RF electrosurgical instrument.

**[1859]** Example 5: The system of any one of Examples 1-4, wherein the modular device comprises a surgical cutting and stapling instrument and the distributed control system is configured to control a rate at which a motor of the surgical cutting and stapling instrument drives a knife.

**[1860]** Example 6: A system comprising: a modular device configured to communicably couple to a surgical hub comprising a hub processor, the modular device comprising: a sensor configured to detect data associated with the modular device; a device memory; and a device processor coupled to the device memory and the sensor; and a distributed control system executable at least in part by each of the device processor and the hub processor, the distributed control system configured to: receive the data detected by the sensor; determine control adjustments for the modular device according to the data; and control the modular device according to the control adjustments; wherein in a first mode the distributed control system is executed by both the hub processor and the device processor, and in a second mode the distributed control system is executed solely by the device processor.

**[1861]** Example 7: The system of any one of Examples 6, wherein the distributed control system is configured to shift from the first mode to the second mode when a sampling rate of the sensor is greater than a data transmission rate from the modular device to the surgical hub.

**[1862]** Example 8: The system of any one of Examples 6-7, wherein the distributed control system is configured to shift from the second mode to the first mode when a sampling rate of the sensor is less than a data transmission rate from the modular device to the surgical hub.

**[1863]** Example 9: The system of any one of Examples 6-8, wherein the modular device comprises a radiofrequency (RF) electrosurgical instrument and the distributed control system is configured to control an energy level of the RF electrosurgical instrument.

**[1864]** Example 10: The system of any one of Examples 6-9, wherein the modular device comprises a surgical cutting and stapling instrument and the distributed control system is

configured to control a rate at which a motor of the surgical cutting and stapling instrument drives a knife.

**[1865]** Example 11: A system configured to control a modular device comprising a sensor configured to detect data associated with the modular device, the system comprising: a first surgical hub configured to communicably couple to the modular device and to a second surgical hub comprising a second processor, the first surgical hub comprising: a memory; and a first processor coupled to the memory; and a distributed control system executable at least in part by each of the first processor and the second processor, the distributed control system configured to: receive the data detected by the sensor; determine control adjustments for the modular device according to the data; and control the modular device according to the control adjustments.

**[1866]** Example 12: The system of any one of Examples 11, wherein the distributed control system is transitionable between a first mode, where the distributed control system is executed by both the first processor and the second processor, and a second mode, where the distributed control system is executed solely by the first processor.

**[1867]** Example 13: The system of any one of Examples 11-12, wherein the distributed control system is configured to shift between the first mode and the second mode upon receiving a command.

**[1868]** Example 14: The system of any one of Examples 11-13, wherein the modular device comprises a radiofrequency (RF) electro-surgical instrument and the distributed control system is configured to control an energy level of the RF electro-surgical instrument.

**[1869]** Example 15: The system of any one of Examples 11-14, wherein the modular device comprises a surgical cutting and stapling instrument and the distributed control system is configured to control a rate at which a motor of the surgical cutting and stapling instrument drives a knife.

**[1870]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1871]** Example 1: A surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive first image data from a first image sensor, wherein the first image data represents a first field of view; receive second image data from a second image sensor, wherein the second image data represents a second field of view; and display, on a display coupled to the processor, a first image rendered from the first image data corresponding to the first field of view and a second image rendered from the second image data corresponding to the second field of view.

**[1872]** Example 2: The surgical hub of Example 1, wherein the first field of view is a narrow angle field of view.

**[1873]** Example 3: The surgical hub of any one of Examples 1-2, wherein the first field of view is a wide angle field of view.

**[1874]** Example 4: The surgical hub of any one of Examples 1-3, wherein the memory stores instructions executable by the processor to augment the first image with the second image on the display.

**[1875]** Example 5: The surgical hub of any one of Examples 1-4, wherein the memory stores instructions executable by the processor to fuse the first image and the second image into a third image and display a fused image on the display.

**[1876]** Example 6: The surgical hub of any one of Examples 1-5, wherein the fused image data comprises status information associated with a surgical device, an image data integration landmark to interlock a plurality of images, and at least one guidance parameter.

**[1877]** Example 7: The surgical hub of any one of Examples 1-6, wherein the first image sensor is the same as the second image sensor and wherein the first image data is captured as a first time by the first image sensor and the second image data is captured at a second time by the first image sensor.

**[1878]** Example 8: The surgical hub of any one of Examples 1-7, wherein the memory stores instructions executable by the processor to: receive third image data from a third image sensor, wherein the third image data represents a third field of view; generate composite image data comprising the second and third image data; display the first image in a first window of the display, wherein the first image corresponds to the first image data; and display a third image in a second window of the display, wherein the third image corresponds to the composite image data.

**[1879]** Example 9: The surgical hub of any one of Examples 1-8, wherein the memory stores instructions executable by the processor to: receive third image data from a third image sensor, wherein the third image data represents a third field of view; fuse the second and third image data to generate fused image data; display the first image in a first window of the display, wherein the first image corresponds to the first image data; and display a third image in a second window of the display, wherein the third image corresponds to the fused image data.

**[1880]** Example 10: A surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: detect a surgical device connection to the surgical hub; transmit a control signal to the detected surgical device to transmit to the surgical hub surgical parameter data associated with the detected surgical device; receive the surgical parameter data from the detected surgical device; receive image data from an image sensor; and display, on a display coupled to the surgical hub, an image rendered based on the image data received from the image sensor in conjunction with the surgical parameter data received from the surgical device.

**[1881]** Example 11: The surgical hub of Example 10, wherein the surgical device comprises a local display that is separate from the display coupled to the surgical hub.

**[1882]** Example 12: The surgical hub of any one of Examples 10-11, wherein the surgical device connected to the surgical hub is configured to reconfigure the local display to present information that is different from information presented when the surgical device is not connected to the surgical hub.

**[1883]** Example 13: The surgical hub of any one of Examples 10-12, wherein a portion of information displayed on the local display is displayed on the display coupled to the surgical hub.

**[1884]** Example 14: The surgical hub of any one of Examples 10-13, wherein information displayed on the display coupled to the surgical hub is mirrored on the local display of the surgical device.

**[1885]** Example 15: A surgical hub, comprising: a control circuit configured to: detect a surgical device connection to the surgical hub; transmit a control signal to the detected surgical device to transmit to the surgical hub surgical

parameter data associated with the detected surgical device; receive the surgical parameter data from the detected surgical device; receive image data from an image sensor; and display, on a display coupled to the surgical hub, an image received from the image sensor in conjunction with the surgical parameter data received from the surgical device.

[1886] Example 16: The surgical hub of Example 15, wherein the surgical device comprises a local display that is separate from the display coupled to the surgical hub.

[1887] Example 17: The surgical hub of any one of Examples 15-16, wherein the surgical device connected to the surgical hub is configured to reconfigure the local display to present information that is different from information presented when the surgical device is not connected to the surgical hub.

[1888] Example 18: The surgical hub of any one of Examples 15-17, wherein a portion of information displayed on the local display is displayed on the display coupled to the surgical hub.

[1889] Example 19: The surgical hub of any one of Examples 15-18, wherein information displayed on the display coupled to the surgical hub is mirrored on the local display of the surgical device.

[1890] Example 20: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: detect a surgical device connection to the surgical hub; transmit a control signal to the detected surgical device to transmit to the surgical hub surgical parameter data associated with the detected surgical device; receive the surgical parameter data from the detected surgical device; receive image data from an image sensor; and display, on a display coupled to the surgical hub, an image received from the image sensor in conjunction with the surgical parameter data received from the surgical device.

[1891] Example 21: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: receive first image data from a first image sensor, wherein the first image data represents a first field of view; receive second image data from a second image sensor, wherein the second image data represents a second field of view; and display, on a display coupled to the surgical hub, a first image corresponding to the first field of view and a second image corresponding to the second field of view.

[1892] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1893] Example 1: A surgical hub, comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive image data from an image sensor; generate a first image based on the image data; display the first image on a surgical hub display coupled to the processor; receive a signal from a non-contact sensor, the signal indicative of a position of a surgical device; generate a second image based on the signal indicative of the position of the surgical device; and display the second image on the surgical hub display coupled to the processor.

[1894] Example 2: The surgical hub of Example 1, wherein the first image data represents a center of a staple line.

[1895] Example 3: The surgical hub of any one of Examples 1-2, wherein the first image represents a target corresponding to the center of the staple line.

[1896] Example 4: The surgical hub of any one of Examples 1-3, wherein the signal is indicative of the position of the surgical device relative to the center of the staple line.

[1897] Example 5: The surgical hub of any one of Examples 1-4, wherein the second image represents the position of the surgical device along a projected path of the surgical device toward the center of the staple line.

[1898] Example 6: The surgical hub of Example 1, wherein the staple line is a double staple line defining a staple overlap portion.

[1899] Example 7: The surgical hub of Example 6, wherein the surgical device is a circular stapler comprising an anvil trocar and the non-contact sensor is configured to detect the location of the anvil trocar relative to the staple overlap portion.

[1900] Example 8: The surgical hub of Example 1, wherein the staple line is a linear staple line formed using a linear transection technique.

[1901] Example 9: The surgical hub of Example 8, wherein a center of the linear staple line is located halfway between one end of the linear staple line and an opposite end of the linear staple line.

[1902] Example 10: The surgical hub of any one of Examples 1-9, wherein the image sensor is coupled to a medical imaging device.

[1903] Example 11: The surgical hub of any one of Examples 1-10, wherein the image sensor and the surgical device are separate devices.

[1904] Example 12: The surgical hub of Example 1, wherein the non-contact sensor is an inductive sensor.

[1905] Example 13: The surgical hub of Example 1, wherein the non-contact sensor is a capacitive sensor.

[1906] Example 14: A method of aligning a surgical instrument coupled to a surgical hub, the method comprising: receiving image data by a processor from an image sensor; generating a first image by the processor based on the image data; displaying the first image on a surgical hub display coupled to the processor; receiving a signal by the processor from a non-contact sensor, the signal indicative of a position of a surgical device; generating a second image by the processor based on the signal indicative of the position of the surgical device; and displaying the second image on the surgical hub display coupled to the processor.

[1907] Example 15: The method of Example 14, comprising displaying, on the surgical hub display coupled to the processor, an indication when the second image is not aligned with the first image.

[1908] Example 16: The method of any one of Examples 14-15, comprising displaying, on the surgical hub display coupled to the processor, an indication when the second image is aligned with the first image.

[1909] Example 17: The method of any one of Examples 14-16, comprising displaying, on the surgical hub display coupled to the processor, a projected path of the surgical device as the second image moves towards the first image.

[1910] Example 18: The method of any one of Examples 14-17, comprising displaying, on the surgical hub display coupled to the processor, the position of the surgical device along the projected path of the surgical device toward the center of the staple line.

[1911] Example 19: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: receive image data by a processor from an image sensor; generate a first image by the processor based on the image data; display the first image on a surgical hub display coupled to the processor; receive a signal by the processor from a non-contact sensor, the signal indicative of a position of a surgical device; generate a second image by the processor based on the signal indicative of the position of the surgical device; and display the second image on the surgical hub display coupled to the processor.

[1912] Example 20: The non-transitory computer readable medium of any one of Example 19, storing computer readable instructions which, when executed, causes a machine to display, on the surgical hub display coupled to the processor, an indication when the second image is not aligned with the first image.

[1913] Example 21: The non-transitory computer readable medium of any one of Examples 19-20, storing computer readable instructions which, when executed, causes a machine to display, on the surgical hub display coupled to the processor, an indication when the second image is aligned with the first image.

[1914] Example 22: The non-transitory computer readable medium of any one of Examples 19-21, storing computer readable instructions which, when executed, causes a machine to display, on the surgical hub display coupled to the processor, a projected path of the surgical device as the second image moves towards the first image.

[1915] Example 23: The non-transitory computer readable medium of any one of Examples 19-22, storing computer readable instructions which, when executed, causes a machine to display, on the surgical hub display coupled to the processor, the position of the surgical device along the projected path of the surgical device toward the center of the staple line.

[1916] Example 24: A surgical hub for aligning a surgical instrument, the surgical hub comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive image data from an image sensor, wherein the first image data represents a center of a staple line; generate a first image based on the image data; display the first image on a monitor coupled to the processor, wherein the first image represents a target corresponding to the center of the staple line; receive a signal from a non-contact sensor, the signal indicative of a position of a surgical device relative to the center of the staple line; and generate a second image based on the position of the surgical device; display the second image on the monitor, wherein the second image represents the position of the surgical device along a projected path of the surgical device toward the center of the staple line.

[1917] Example 25: The surgical hub of Example 24, wherein the center of the staple line is a double-staple overlap portion zone.

[1918] Example 26: The surgical hub of any one of Examples 24-25, wherein the image sensor receives an image from a medical imaging device.

[1919] Example 27: The surgical hub of any one of Examples 24-26, wherein the surgical device is a circular stapler comprising an anvil trocar and the non-contact sensor is configured to detect the location of the anvil trocar relative to the center of the staple line.

[1920] Example 28: The surgical hub of Example 24, wherein the non-contact sensor is an inductive sensor.

[1921] Example 29: The surgical hub of Example 24, wherein the non-contact sensor is a capacitive sensor.

[1922] Example 30: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: receive image data from an image sensor, wherein the first image data represents a center of a staple line; generate a first image based on the image data; display the first image on a monitor coupled to the processor, wherein the first image represents a target corresponding to the center of the staple line; receive a signal from a non-contact sensor, wherein the signal is indicative of a position of a surgical device relative to the center of the staple line; generate a second image based on the position of the surgical device; and display the second image on the monitor, wherein the second image represents the position of the surgical device along a projected path of the surgical device toward the center of the staple line.

[1923] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1924] Example 1: An interactive control unit, comprising: an interactive touchscreen display; an interface configured to couple the control unit to a surgical hub; a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive input commands from the interactive touchscreen display located inside a sterile field; and transmit the input commands to the surgical hub to control devices coupled to the surgical hub located outside the sterile field.

[1925] Example 2: The interactive control unit of Example 1, wherein the processor is configured to receive an image array from a scanning device and display the image on the interactive touchscreen display.

[1926] Example 3: The interactive control unit of any one of Examples 1-2, wherein the processor is configured to display on the interactive touchscreen display an image of a virtual anatomy based on the received image array.

[1927] Example 4: The interactive control unit of any one of Examples 1-3, wherein the processor is configured to receive an image array from a laser Doppler scanning device.

[1928] Example 5: The interactive control unit of any one of Examples 1-4, wherein the processor is configured to re-configure wireless devices coupled to the surgical hub from control inputs received via the interactive touchscreen display.

[1929] Example 6: The interactive control unit of any one of Examples 1-5, wherein the interactive touchscreen display comprises multiple input and output zones.

[1930] Example 7: An interactive control unit, comprising: an interactive touchscreen display; an interface configured to couple the control unit to a first surgical hub; a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive input commands from the interactive touchscreen display located inside a sterile field; transmit the input commands to the first surgical hub to control devices coupled to the first surgical hub located outside the sterile field; receive a consult request from a second surgical hub; and configure a portion of the interactive touchscreen display to display information received from the second surgical hub after receiving the consult request.

[1931] Example 8: The interactive control unit of Example 7, wherein the processor is configured to temporarily store data associated with the interactive touchscreen display.

[1932] Example 9: The interactive control unit of any one of Examples 7-8, wherein the processor is configured to back up the data in time.

[1933] Example 10: The interactive control unit of any one of Examples 7-9, wherein the processor is configured to view the information received from the second surgical hub.

[1934] Example 11: The interactive control unit of any one of Examples 7-10, wherein the processor is configured to delete the information received from the second surgical hub.

[1935] Example 12: The interactive control unit of any one of Examples 7-11, wherein the processor is configured to return control to the interactive surgical touchscreen in the first surgical hub.

[1936] Example 13: An interactive control unit, comprising: an interactive touchscreen display; an interface configured to couple the control unit to a surgical hub; and a control circuit to: receive input commands from the interactive touchscreen display located inside a sterile field; and transmit the input commands to the surgical hub to control devices coupled to the surgical hub located outside the sterile field.

[1937] Example 14: The interactive control unit of Example 13, wherein the control circuit is configured to receive an image array from a scanning device and display the image on the interactive touchscreen display.

[1938] Example 15: The interactive control unit of any one of Examples 13-14, wherein the control circuit is configured to display on the interactive touchscreen display an image of a virtual anatomy based on the received image array.

[1939] Example 16: The interactive control unit of any one of Examples 13-15, wherein the control circuit is configured to receive an image array from a laser Doppler scanning device.

[1940] Example 17: The interactive control unit of any one of Examples 13-16, wherein the control circuit is configured to re-configure wireless devices coupled to the surgical hub from control inputs received via the interactive touchscreen display.

[1941] Example 18: The interactive control unit of any one of Examples 13-17, wherein the interactive touchscreen display comprises multiple input and output zones.

[1942] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1943] Example 1: A surgical hub for use with a surgical instrument configured to deliver therapeutic energy to tissue at a surgical site of a surgical procedure, wherein the surgical hub comprises: a hub enclosure, comprising a docking station including a docking port comprising data and power contacts; and a combo generator module removably retainable in the docking station, wherein the combo generator module comprises: an ultrasonic energy generator component; a radio frequency (RF) energy generator component; a smoke evacuation component; a connection port, wherein at least one of the ultrasonic energy generator component and the radio frequency (RF) generator component is coupleable to the surgical instrument through the connection port; and at least one smoke evacuation component configured to evacuate smoke generated by an application of therapeutic energy to the tissue by the surgical instrument.

[1944] Example 2: The surgical hub of Example 1, wherein the docking station is a first docking station, wherein the docking port is a first docking port, and wherein the hub enclosure comprises a second docking station comprising a second docking port that has data and power contacts.

[1945] Example 3: The surgical hub of Example 2, further comprising a suction and irrigation module removably retainable in the second docking station.

[1946] Example 4: The surgical hub of Example 3, wherein the combo generator module comprises a third docking port connectable to the first docking port of the first docking station.

[1947] Example 5: The surgical hub of Example 4, wherein the suction and irrigation module comprises a fourth docking port connectable to the second docking port of the second docking station.

[1948] Example 6: The surgical hub of Example 5, wherein the hub enclosure comprises a communication link between the second docking port and the first docking port.

[1949] Example 7: The surgical hub of any of Examples 1-6, wherein the combo generator module comprises a fluid line extendable to the remote surgical site for passing the smoke evacuated from the remote surgical site to the combo generator module.

[1950] Example 8: The surgical hub of any one of Examples 1-7, wherein the docking station comprises brackets configured to slidably receive and guide the combo generator module into a working connection with the power and data contacts of the docking port.

[1951] Example 9: The surgical hub of any one of Examples 1-8, wherein the combo generator module comprises side brackets configured to movably engage the brackets of the docking station.

[1952] Example 10: A modular surgical hub for use with a surgical instrument configured to deliver therapeutic energy to tissue at a surgical site of a surgical procedure, wherein the modular surgical enclosure comprises: a first energy-generator module configured to generate a first therapeutic energy for application to the tissue; a first docking station comprising a first docking port that includes first data and power contacts, wherein the first energy-generator module is slidably movable into an electrical engagement with the first data and power contacts, and wherein the first energy-generator module is slidably movable out of the electrical engagement with the first data and power contacts; a second energy-generator module configured to generate a second therapeutic energy, different than the first therapeutic energy, for application to the tissue; a second docking station comprising a second docking port that includes second data and power contacts, wherein the second energy-generator module is slidably movable into an electrical engagement with the second data and power contacts, and wherein the second energy-generator module is slidably movable out of the electrical engagement with the second data and power contacts; and a communication bus between the first docking port and the second docking port configured to facilitate communication between the first energy-generator module and the second energy-generator module.

[1953] Example 11: The modular surgical hub of Example 10, wherein the first docking station comprises brackets configured to slidably receive and guide the first energy-generator module into the electrical engagement with the first data and power contacts.

[1954] Example 12: The modular surgical hub of Example 11, wherein the second docking station comprises brackets configured to slidably receive and guide the second energy-generator module into the electrical engagement with the second data and power contacts.

[1955] Example 13: The modular surgical hub of any one of Examples 10-12, wherein the first therapeutic energy is an ultrasonic energy.

[1956] Example 14: The modular surgical hub of any one of Exemplified 10-12, wherein the second therapeutic energy is a radio frequency (RF) energy.

[1957] Example 15: The modular surgical hub of any one of Examples 10-14, further comprising a smoke evacuation module configured to evacuate smoke generated at the remote surgical site by application of the first therapeutic energy to the tissue.

[1958] Example 16: The modular surgical hub of Example 15, further comprising a third docking station comprising a third docking port that includes third data and power contacts.

[1959] Example 17: The modular surgical hub of Example 16, further comprising a suction and irrigation module slidably movable into an electrical engagement with the third data and power contacts, and wherein the suction and irrigation module is slidably movable out of the electrical engagement with the third data and power contacts.

[1960] Example 18: A surgical hub for use with a surgical instrument configured to deliver therapeutic energy to tissue at a surgical site of a surgical procedure, wherein the surgical hub comprises: a hub enclosure, comprising docking stations including docking ports comprising data and power contacts; a combo generator module slidably receivable in a first of the docking stations, wherein the combo generator module comprises: an ultrasonic energy generator component; a radio frequency (RF) energy generator component; and a connection port, wherein at least one of the ultrasonic energy generator component and the radio frequency (RF) generator component is couplable to the surgical instrument through the connection port; a smoke evacuation module slidably receivable in a second one of the docking stations, wherein the smoke evacuation module is configured to evacuate smoke generated by an application of the therapeutic energy to the tissue by the surgical instrument; a processing module slidably receivable in a third one of the docking stations; a memory module slidably receivable in a fourth one of the docking stations; and an operating-room mapping module slidably receivable in a fifth one of the docking stations.

[1961] Example 19: The surgical hub of Example 18, wherein the docking stations comprise brackets configured to slidably guide the modules into electrical engagements with the power and data contacts of the docking ports.

[1962] Example 20: The surgical hub of any one of Examples 18-19, comprising a display.

[1963] Various additional aspects of the subject matter described herein are set out in the following numbered examples.

[1964] Example 1: A surgical image acquisition system comprising: a plurality of illumination sources wherein each illumination source is configured to emit light having a specified central wavelength; a light sensor configured to receive a portion of the light reflected from a tissue sample when illuminated by the one or more of the plurality of illumination sources; and a computing system, wherein the

computing system is configured to: receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; determine a depth location of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; and calculate visualization data regarding the structure and the depth location of the structure, wherein the visualization data have a data format that may be used by a display system, and wherein the structure comprises one or more vascular tissues.

[1965] Example 2: The surgical image acquisition system of any one of Example 1, wherein the plurality of illumination sources comprises an illumination source having a central wavelength in a range between 635 nm and 660 nm, inclusive.

[1966] Example 3: The surgical image acquisition system of any one of Examples 1-2, wherein the plurality of illumination sources comprises an illumination source having a central wavelength in a range between 750 nm and 3000 nm.

[1967] Example 4: The surgical image acquisition system of any one of Examples 1-3, wherein the plurality of illumination sources comprises an illumination source configured to emit a broad spectral range of illumination.

[1968] Example 5: The surgical image acquisition system of any one of Examples 1-4, wherein the plurality of illumination sources comprises a laser illumination source.

[1969] Example 6: A surgical image acquisition system comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: control the operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive data from a light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; determine a depth location of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; and calculate visualization data regarding the structure and the depth location of the structure, wherein the visualization data have a data format that may be used by a display system, and wherein the structure comprises one or more vascular tissues.

[1970] Example 7: The surgical image acquisition system of any one of Example 6, wherein the instruction, executable by the processor, to determine a depth location of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources comprises an instruction to determine a depth location of a structure within the tissue sample based on a central wavelength of light emitted by at least one of the plurality of illumination sources.

[1971] Example 8: The surgical image acquisition system of any one of Example 6 through Example 7, wherein the instructions, executable by the processor, further comprise an instruction to calculate a flow of a material through the one or more vascular tissues.

[1972] Example 9: The surgical image acquisition system of any one of Example 8, wherein the instruction, executable by the processor, to calculate visualization data regarding the structure and the depth location of the structure further includes an instruction, executable by the processor, to



calculate visualization data including data representative of the flow of material through the one or more vascular tissues.

**[1973]** Example 10: A surgical image acquisition system comprising: a control circuit configured to: control the operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive data from a light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; determine a depth location of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; and calculate visualization data regarding the structure and the depth location of the structure, wherein the visualization data have a data format that may be used by a display system, and wherein the structure comprises one or more vascular tissues.

**[1974]** Example 11: The surgical image acquisition system of any one of Example 10, wherein the control circuit configured to control the operation of a plurality of illumination sources of a tissue sample comprises a control circuit configured to sequentially actuate each of the plurality of illumination sources to illuminate the tissue sample.

**[1975]** Example 12: The surgical image acquisition system of any one of Examples 10-11, wherein the control circuit configured to determine a depth location of a structure within the tissue sample comprises a control circuit configured to determine the depth location of the structure based on a penetration depth of illumination sourced by each of the plurality of illumination sources.

**[1976]** Example 13: The surgical image acquisition system of any one of Example 12, wherein the structure comprises a surface structure within the tissue sample.

**[1977]** Example 14: The surgical image acquisition system of any one of Examples 10-13, wherein the control circuit configured to control the operation of a plurality of illumination sources of a tissue sample comprises a control circuit configured to operate at least one of a red light illumination source, a green light illumination source, and a blue light illumination source.

**[1978]** Example 15: The surgical image acquisition system of any one of Examples 10-14, wherein the control circuit configured to control the operation of a plurality of illumination sources of a tissue sample comprises a control circuit configured to operate at least one of an infrared light illumination source and an ultraviolet light illumination source.

**[1979]** Example 16: The surgical image acquisition system of any one of Examples 10-15, wherein the control circuit is further configured to determine a flow of material through the one or more vascular tissues.

**[1980]** Example 17: The surgical image acquisition system of any one of Example 16, wherein the control circuit configured to determine a flow of material through the one or more vascular tissues comprises a control circuit configured to analyze the data received from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources for a Doppler shift in wavelength of light emitted by each of the plurality of illumination sources.

**[1981]** Example 18: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: control the operation of a plurality of illumination sources of a tissue sample wherein

each illumination source is configured to emit light having a specified central wavelength; receive data from a light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; determine a depth location of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; and calculate visualization data regarding the structure and the depth location of the structure, wherein the visualization data have a data format that may be used by a display system, and wherein the structure comprises one or more vascular tissues.

**[1982]** Example 19: The non-transitory computer readable medium of any one of Example 18, wherein the computer readable instructions, when executed, further cause the machine to: control the operation of an additional illumination source wherein the additional illumination source is a white light source; and receive data from the light sensor when the tissue sample is illuminated by the white light source.

**[1983]** Example 20: The non-transitory computer readable medium of any one of Example 19, wherein the computer readable instructions, when executed, that cause the machine to calculate visualization data regarding the structure and the depth location of the structure further cause the machine to calculate visualization data based on the data received from the light sensor when the tissue sample is illuminated by the white light source.

**[1984]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[1985]** Example 1: A surgical image acquisition system comprising: a plurality of illumination sources wherein each illumination source is configured to emit light having a specified central wavelength; a light sensor configured to receive a portion of the light reflected from a tissue sample when illuminated by the one or more of the plurality of illumination sources; and a computing system, wherein the computing system is configured to: receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; calculate structural data related to a characteristic of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the illumination sources; and transmit the structural data related to the characteristic of the structure to be received by a smart surgical device, wherein the characteristic of the structure is a surface characteristic or a structure composition.

**[1986]** Example 2: The surgical image acquisition system of any one of Example 1, wherein the plurality of illumination sources comprises at least one of a red light illumination source, a green light illumination source, and a blue light illumination source.

**[1987]** Example 3: The surgical image acquisition system of any one of Examples 1-2, wherein the plurality of illumination sources comprises at least one of an infrared light illumination source and an ultraviolet light illumination source.

**[1988]** Example 4: The surgical image acquisition system of any one of Examples 1-3, wherein the computing system, configured to calculate structural data related to a characteristic of a structure within the tissue, comprises a computing system configured to calculate structural data related to a composition of a structure within the tissue.

**[1989]** Example 5: The surgical image acquisition system of any one of Examples 1-4, wherein the computing system, configured to calculate structural data related to a characteristic of a structure within the tissue, comprises a computing system configured to calculate structural data related to a surface roughness of a structure within the tissue.

**[1990]** Example 6: A surgical image acquisition system comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: control the operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; calculate structural data related to a characteristic of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the illumination sources; and transmit the structural data related to the characteristic of the structure to be received by a smart surgical device, wherein the characteristic of the structure is a surface characteristic or a structure composition.

**[1991]** Example 7: The surgical image acquisition system of any one of Example 6, wherein the instructions executable by the processor to control the operation of a plurality of illumination sources comprise one or more instructions to illuminate the tissue sample sequentially by each of the plurality of illumination sources.

**[1992]** Example 8: The surgical image acquisition system of any one of Example 6 through Example 7 wherein the instructions executable by the processor to calculate structural data related to a characteristic of a structure within the tissue sample based on the data received by the light sensor comprise one or more instructions to calculate structural data related to a characteristic of a structure within the tissue sample based on a phase shift in the illumination reflected by the tissue sample.

**[1993]** Example 9: The surgical image acquisition system of any one of Examples 6-8, wherein the structure composition comprises a relative composition of collagen and elastin in a tissue.

**[1994]** Example 10: The surgical image acquisition system of any one of Examples 6-9, wherein the structure composition comprises an amount of hydration of a tissue.

**[1995]** Example 11: A surgical image acquisition system comprising: a control circuit configured to: control the operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; calculate structural data related to a characteristic of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the illumination sources; and transmit the structural data related to the characteristic of the structure to be received by a smart surgical device, wherein the characteristic of the structure is a surface characteristic or a structure composition.

**[1996]** Example 12: The surgical image acquisition system of any one of Example 11, wherein the control circuit is configured to transmit the structural data related to the

characteristic of the structure to be received by a smart surgical device wherein the smart surgical device is a smart surgical stapler.

**[1997]** Example 13: The surgical image acquisition system of any one of Example 12, wherein the control circuit is further configured to transmit data related to an anvil pressure based on the characteristic of the structure to be received by the smart surgical stapler.

**[1998]** Example 14: The surgical image acquisition system of any one of Examples 11-13, wherein the control circuit is configured to transmit the structural data related to the characteristic of the structure to be received by a smart surgical device wherein the smart surgical device is a smart surgical RF sealing device.

**[1999]** Example 15: The surgical image acquisition system of any one of Example 14, wherein the control circuit is further configured to transmit data related to an amount of RF power based on the characteristic of the structure to be received by the smart RF sealing device.

**[2000]** Example 16: The surgical image acquisition system of any one of Examples 11-15, wherein the control circuit is configured to transmit the structural data related to the characteristic of the structure to be received by a smart surgical device wherein the smart surgical device is a smart ultrasound cutting device.

**[2001]** Example 17: The surgical image acquisition system of any one of Example 16, wherein the control circuit is further configured to transmit data related to an amount of power provided to an ultrasonic transducer or a driving frequency of the ultrasonic transducer based on the characteristic of the structure to be received by the ultrasound cutting device.

**[2002]** Example 18: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: control the operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive data from the light sensor when the tissue sample is illuminated by each of the plurality of illumination sources; calculate structural data related to a characteristic of a structure within the tissue sample based on the data received by the light sensor when the tissue sample is illuminated by each of the illumination sources; and transmit the structural data related to the characteristic of the structure to be received by a smart surgical device, wherein the characteristic of the structure is a surface characteristic or a structure composition.

**[2003]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2004]** Example 1: A minimally invasive image acquisition system comprising: a plurality of illumination sources wherein each illumination source is configured to emit light having a specified central wavelength; a first light sensing element having a first field of view and configured to receive illumination reflected from a first portion of a surgical site when the first portion of the surgical site is illuminated by at least one of the plurality of illumination sources; a second light sensing element having a second field of view and configured to receive illumination reflected from a second portion of the surgical site when the second portion of the surgical site is illuminated by at least one of the plurality of illumination sources, wherein the second field of view overlaps at least a portion of the first field of view; and a

computing system, wherein the computing system is configured to: receive data from the first light sensing element, receive data from the second light sensing element, compute imaging data based on the data received from the first light sensing element and the data received from the second light sensing element, and transmit the imaging data for receipt by a display system.

**[2005]** Example 2: The minimally invasive image acquisition system of any one of Example 1, wherein the first field of view has a first angle and the second field of view has a second angle and the first angle is the same as the second angle.

**[2006]** Example 3: The minimally invasive image acquisition system of any one of Examples 1-2, wherein the first field of view has a first angle and the second field of view has a second angle and the first angle differs from the second angle.

**[2007]** Example 4: The minimally invasive image acquisition system of any one of Examples 1-3, wherein the first light sensing element has an optical component configured to adjust the first field of view.

**[2008]** Example 5: The minimally invasive image acquisition system of any one of Examples 1-4, wherein the second light sensing element has an optical component configured to adjust the second field of view.

**[2009]** Example 6: The minimally invasive image acquisition system of any one of Examples 1-5, wherein the second field of view overlaps all of the first field of view.

**[2010]** Example 7: The minimally invasive image acquisition system of any one of Examples 1-6, wherein the first field of view is completely enclosed by the second field of view.

**[2011]** Example 8: The minimally invasive image acquisition system of any one of Examples 1-7, wherein the first light sensing element and the second light sensing element are at least partially disposed within an elongated camera probe.

**[2012]** Example 9: The minimally invasive image acquisition system of any one of Examples 1-8, wherein each of the plurality of illumination source is configured to emit light having a specified central wavelength within a visible spectrum.

**[2013]** Example 10: The minimally invasive image acquisition system of any one of Examples 1-9, wherein at least one of the plurality of illumination source is configured to emit light having a specified central wavelength outside of a visible spectrum.

**[2014]** Example 11: The minimally invasive image acquisition system of any one of Example 10, wherein the specified central wavelength outside of the visible spectrum is within an ultra-violet range.

**[2015]** Example 12: The minimally invasive image acquisition system of any one of Examples 10-11, wherein the specified central wavelength outside of the visible spectrum is within an infrared range.

**[2016]** Example 13: The minimally invasive image acquisition system of any one of Examples 1-12, wherein the computing system configured to compute imaging data based on the data received from the first light sensing element and the data received from the second light sensing element comprises a computing system configured to perform a first data analysis on the data received from the first light sensing element and a second data analysis on the data received from the second light sensing element.

**[2017]** Example 14: The minimally invasive image acquisition system of any one of Example 13, wherein the first data analysis differs from the second data analysis.

**[2018]** Example 15: A minimally invasive image acquisition system comprising: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: control an operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive, from a first light sensing element, first data related to illumination reflected from a first portion of a surgical site when the first portion of the surgical site is illuminated by at least one of the plurality of illumination source, receive, from a second light sensing element, second data related to illumination reflected from a second portion of the surgical site when the second portion of the surgical site is illuminated by at least one of the plurality of illumination sources, wherein the second field of view overlaps at least a portion of the first field of view, compute imaging data based on the first data received from the first light sensing element and the second data received from the second light sensing element, and transmit the imaging data for receipt by a display system.

**[2019]** Example 16: The minimally invasive image acquisition system of any one of Example 15, wherein the memory coupled to the processor further stores instructions executable by the processor to receive, from a surgical instrument, operational data related to a function or status of the surgical instrument.

**[2020]** Example 17: The minimally invasive image acquisition system of any one of Example 16, wherein the memory coupled to the processor further stores instructions executable by the processor to compute imaging data based on the first data received from the first light sensing element, the second data received from the second light sensing element, and the operational data related to the function or status of the surgical instrument.

**[2021]** Example 18: A minimally invasive image acquisition system comprising: a control circuit configured to: control an operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive, from a first light sensing element, first data related to illumination reflected from a first portion of a surgical site when the first portion of the surgical site is illuminated by at least one of the plurality of illumination source, receive, from a second light sensing element, second data related to illumination reflected from a second portion of the surgical site when the second portion of the surgical site is illuminated by at least one of the plurality of illumination sources, wherein the second field of view overlaps at least a portion of the first field of view, compute imaging data based on the first data received from the first light sensing element and the second data received from the second light sensing element, and transmit the imaging data for receipt by a display system.

**[2022]** Example 19: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a machine to: control an operation of a plurality of illumination sources of a tissue sample wherein each illumination source is configured to emit light having a specified central wavelength; receive, from a first light sensing element, first data related to illumination reflected from a first portion of a surgical site when the first portion

of the surgical site is illuminated by at least one of the plurality of illumination source, receive, from a second light sensing element, second data related to illumination reflected from a second portion of the surgical site when the second portion of the surgical site is illuminated by at least one of the plurality of illumination sources, wherein the second field of view overlaps at least a portion of the first field of view, compute imaging data based on the first data received from the first light sensing element and the second data received from the second light sensing element, and transmit the imaging data for receipt by a display system.

**[2023]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2024]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2025]** Example 1: An analytics system configured to communicably couple to a surgical hub, the surgical hub configured to communicable couple to a modular device that is controlled by a control program, the analytics system comprising: a processor; and a memory coupled to the processor, the memory storing instructions that, when executed by the processor, cause the analytics system to: receive perioperative data indicative of an operational behavior of the modular device, wherein the perioperative data comprises data detected by the modular device during a surgical procedure; receive procedural outcome data associated with the surgical procedure; analyze the perioperative data and the procedural outcome data to determine whether the operational behavior is suboptimal; generate a control program update configured to alter the manner in which the control program operates the modular device during the surgical procedure for the operational behavior; and transmit the control program update to the modular device.

**[2026]** Example 2: The analytics system of Example 1, wherein the memory stores instructions that, when executed by the processor, cause the analytics system to determine whether the operational behavior is suboptimal according to whether the operational behavior correlates to a negative procedural outcome.

**[2027]** Example 3: The analytics system of any one of Examples 1-2, wherein: the operational behavior is a first operational behavior; the perioperative data is further indicative of a second operational behavior; and the memory stores instructions that, when executed by the processor, cause the analytics system to determine whether the first operational behavior is suboptimal according to whether the second operational behavior is more highly correlated to a positive procedural outcome than the first operational behavior.

**[2028]** Example 4: The analytics system of any one of Examples 1-3, wherein the control program update is configured to provide an alert associated with the operational behavior.

**[2029]** Example 5: The analytics system of any one of Examples 1-4, wherein the control program update is configured to change a manually controlled function to a function controlled by the control program.

**[2030]** Example 6: The analytics system of any one of Examples 1-5, wherein the memory stores instructions that, when executed by the processor, cause the analytics system to receive the procedural outcome data from an EMR database.

**[2031]** Example 7: The analytics system of any one of Examples 1-6, wherein the memory stores instructions that, when executed by the processor, cause the analytics system to receive the procedural outcome data from the surgical hub.

**[2032]** Example 8: An analytics system configured to communicably couple to a surgical hub, the surgical hub configured to communicable couple to a modular device that is controlled by a control program, the analytics system comprising: a control circuit configured to: receive perioperative data indicative of an operational behavior of the modular device, wherein the perioperative data comprises data detected by the modular device during a surgical procedure; receive procedural outcome data associated with the surgical procedure; analyze the perioperative data and the procedural outcome data to determine whether the operational behavior is suboptimal; generate a control program update configured to alter the manner in which the control program operates the modular device during the surgical procedure for the operational behavior; and transmit the control program update to the modular device.

**[2033]** Example 9: The analytics system of Example 8, wherein the control circuit is configured to determine whether the operational behavior is suboptimal according to whether the operational behavior correlates to a negative procedural outcome.

**[2034]** Example 10: The analytics system of any one of Examples 8-9, wherein: the operational behavior is a first operational behavior; the perioperative data is further indicative of a second operational behavior; and the control circuit is configured to determine whether the first operational behavior is suboptimal according to whether the second operational behavior is more highly correlated to a positive procedural outcome than the first operational behavior.

**[2035]** Example 11: The analytics system of any one of Examples 8-10, wherein the control program update is configured to provide an alert associated with the operational behavior.

**[2036]** Example 12: The analytics system of any one of Examples 8-11, wherein the control program update is configured to change a manually controlled function to a function controlled by the control program.

**[2037]** Example 13: The analytics system of any one of Examples 8-12, wherein the control circuit is configured to cause the analytics system to receive the procedural outcome data from an EMR database.

**[2038]** Example 14: The analytics system of any one of Examples 8-13, wherein the control circuit is configured to cause the analytics system to receive the procedural outcome data from the surgical hub.

**[2039]** Example 15: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes an analytics system configured to communicably couple to a surgical hub, the surgical hub configured to communicable couple to a modular device that is controlled by a control program, to: receive perioperative data indicative of an operational behavior of the modular device, wherein the perioperative data comprises data detected by the modular device during a surgical procedure; receive procedural outcome data associated with the surgical procedure; analyze the perioperative data and the procedural outcome data to determine whether the operational behavior is suboptimal; generate a control program update configured to alter the manner in which the control program operates the

modular device during the surgical procedure for the operational behavior; and transmit the control program update to the modular device.

**[2040]** Example 16: The non-transitory computer readable medium of Example 15, wherein the non-transitory computer readable medium stores instructions that cause the analytics system to determine whether the operational behavior is suboptimal according to whether the operational behavior correlates to a negative procedural outcome.

**[2041]** Example 17: The non-transitory computer readable medium of any one of Examples 15-16, wherein: the operational behavior is a first operational behavior; the perioperative data is further indicative of a second operational behavior; and the non-transitory computer readable medium stores instructions that cause the analytics system to determine whether the first operational behavior is suboptimal according to whether the second operational behavior is more highly correlated to a positive procedural outcome than the first operational behavior.

**[2042]** Example 18: The non-transitory computer readable medium of any one of Examples 15-17, wherein the control program update is configured to provide an alert associated with the operational behavior.

**[2043]** Example 19: The non-transitory computer readable medium of any one of Examples 15-18, wherein the control program update is configured to change a manually controlled function to a function controlled by the control program.

**[2044]** Example 20: The non-transitory computer readable medium of any one of Examples 15-19, wherein the non-transitory computer readable medium stores instructions that cause the analytics system to receive the procedural outcome data from an EMR database.

**[2045]** Example 21: The non-transitory computer readable medium of any one of Examples 15-20, wherein the non-transitory computer readable medium stores instructions that cause the analytics system to receive the procedural outcome data from the surgical hub.

**[2046]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2047]** Example 1: An analytics system configured to communicably couple to a surgical hub, the surgical hub configured to communicably couple to a modular device that is controlled by a control program, the analytics system comprising: a processor; and a memory coupled to the processor, the memory storing instructions that, when executed by the processor, cause the analytics system to: receive perioperative data indicative of an operational behavior of the modular device, wherein the perioperative data comprises data detected by the modular device during a surgical procedure; receive procedural outcome data associated with the surgical procedure; analyze the perioperative data and the procedural outcome data to determine whether the operational behavior is suboptimal; generate a control program update configured to alter the manner in which the control program operates the modular device during the surgical procedure for the operational behavior; and transmit the control program update to the modular device.

**[2048]** Example 2: The analytics system of Example 1, wherein the memory stores instructions that, when executed by the processor, cause the analytics system to determine

whether the operational behavior is suboptimal according to whether the operational behavior correlates to a negative procedural outcome.

**[2049]** Example 3: The analytics system of any one of Examples 1-2, wherein: the operational behavior is a first operational behavior; the perioperative data is further indicative of a second operational behavior; and the memory stores instructions that, when executed by the processor, cause the analytics system to determine whether the first operational behavior is suboptimal according to whether the second operational behavior is more highly correlated to a positive procedural outcome than the first operational behavior.

**[2050]** Example 4: The analytics system of any one of Examples 1-3, wherein the control program update is configured to provide an alert associated with the operational behavior.

**[2051]** Example 5: The analytics system of any one of Examples 1-4, wherein the control program update is configured to change a manually controlled function to a function controlled by the control program.

**[2052]** Example 6: The analytics system of any one of Examples 1-5, wherein the memory stores instructions that, when executed by the processor, cause the analytics system to receive the procedural outcome data from an EMR database.

**[2053]** Example 7: The analytics system of any one of Examples 1-6, wherein the memory stores instructions that, when executed by the processor, cause the analytics system to receive the procedural outcome data from the surgical hub.

**[2054]** Example 8: An analytics system configured to communicably couple to a surgical hub, the surgical hub configured to communicably couple to a modular device that is controlled by a control program, the analytics system comprising: a control circuit configured to: receive perioperative data indicative of an operational behavior of the modular device, wherein the perioperative data comprises data detected by the modular device during a surgical procedure; receive procedural outcome data associated with the surgical procedure; analyze the perioperative data and the procedural outcome data to determine whether the operational behavior is suboptimal; generate a control program update configured to alter the manner in which the control program operates the modular device during the surgical procedure for the operational behavior; and transmit the control program update to the modular device.

**[2055]** Example 9: The analytics system of Example 8, wherein the control circuit is configured to determine whether the operational behavior is suboptimal according to whether the operational behavior correlates to a negative procedural outcome.

**[2056]** Example 10: The analytics system of any one of Examples 8-9, wherein: the operational behavior is a first operational behavior; the perioperative data is further indicative of a second operational behavior; and the control circuit is configured to determine whether the first operational behavior is suboptimal according to whether the second operational behavior is more highly correlated to a positive procedural outcome than the first operational behavior.

**[2057]** Example 11: The analytics system of any one of Examples 8-10, wherein the control program update is configured to provide an alert associated with the operational behavior.

**[2058]** Example 12: The analytics system of any one of Examples 8-11, wherein the control program update is configured to change a manually controlled function to a function controlled by the control program.

**[2059]** Example 13: The analytics system of any one of Examples 8-12, wherein the control circuit is configured to cause the analytics system to receive the procedural outcome data from an EMR database.

**[2060]** Example 14: The analytics system of any one of Examples 8-13, wherein the control circuit is configured to cause the analytics system to receive the procedural outcome data from the surgical hub.

**[2061]** Example 15: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes an analytics system configured to communicably couple to a surgical hub, the surgical hub configured to communicably couple to a modular device that is controlled by a control program, to: receive perioperative data indicative of an operational behavior of the modular device, wherein the perioperative data comprises data detected by the modular device during a surgical procedure; receive procedural outcome data associated with the surgical procedure; analyze the perioperative data and the procedural outcome data to determine whether the operational behavior is suboptimal; generate a control program update configured to alter the manner in which the control program operates the modular device during the surgical procedure for the operational behavior; and transmit the control program update to the modular device.

**[2062]** Example 16: The non-transitory computer readable medium of Example 15, wherein the non-transitory computer readable medium stores instructions that cause the analytics system to determine whether the operational behavior is suboptimal according to whether the operational behavior correlates to a negative procedural outcome.

**[2063]** Example 17: The non-transitory computer readable medium of any one of Examples 15-16, wherein: the operational behavior is a first operational behavior; the perioperative data is further indicative of a second operational behavior; and the non-transitory computer readable medium stores instructions that cause the analytics system to determine whether the first operational behavior is suboptimal according to whether the second operational behavior is more highly correlated to a positive procedural outcome than the first operational behavior.

**[2064]** Example 18: The non-transitory computer readable medium of any one of Examples 15-17, wherein the control program update is configured to provide an alert associated with the operational behavior.

**[2065]** Example 19: The non-transitory computer readable medium of any one of Examples 15-18, wherein the control program update is configured to change a manually controlled function to a function controlled by the control program.

**[2066]** Example 20: The non-transitory computer readable medium of any one of Examples 15-19, wherein the non-transitory computer readable medium stores instructions that cause the analytics system to receive the procedural outcome data from an EMR database.

**[2067]** Example 21: The non-transitory computer readable medium of any one of Examples 15-20, wherein the non-transitory computer readable medium stores instructions that cause the analytics system to receive the procedural outcome data from the surgical hub.

**[2068]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2069]** Example 1: A cloud based analytics medical system comprising: at least one processor; at least one memory communicatively coupled to the processor; an input/output interface configured for accessing data from a plurality of medical hub communication devices, each communicatively coupled to at least one surgical instrument; and a database residing in the at least one memory and configured to store the data; the at least one memory storing instructions executable by the at least one processor to: aggregate medical resource usage data from the plurality of medical hubs, the medical resource usage data comprising: data pertaining to medical products and an indication of efficiency based on their usage; disposal records of when the medical products were disposed of; and for each description of the medical product: location data describing which medical facility said medical product was allocated to; and outcome data pertaining to an outcome of a patient from a procedure that utilized the medical product; determine a correlation between positive outcomes from the outcome data and location data of the medical product; generate a medical recommendation to change a medical resource usage practice based on the correlation; and display the medical recommendation to at least one medical hub at the local facility.

**[2070]** Example 2: The cloud based analytics medical system of Example 1, wherein the disposal records are derived at least in part from disposal bins configured to automatically record an amount of medical products disposed into the bins.

**[2071]** Example 3: The cloud based analytics medical system of any one of Examples 1-2, wherein the outcome data is derived at least in part from operational data transmitted by a medical device used during the procedure.

**[2072]** Example 4: The cloud based analytics medical system of any one of Examples 1-3, wherein the operational data includes a recordation by the medical device of a number of staple firings that were fired by the medical device during the procedure.

**[2073]** Example 5: The cloud based analytics medical system of any one of Examples 1-4, wherein the recommendation comprises a recommendation to substitute use of a first medical product for user of a second medical product during a specific medical procedure.

**[2074]** Example 6: The cloud based analytics medical system of any one of Examples 1-5, wherein the recommendation comprises a recommendation to reduce a number of staple firings that are fired by a medical device during a specific medical procedure.

**[2075]** Example 7: The cloud based analytics medical system of any one of Examples 1-6, wherein the recommendation comprises a recommendation to reduce a rate of use of the medical product during a specific medical procedure.

**[2076]** Example 8: A method of a cloud based analytics medical system for improving efficiency in a medical environment, the method comprising: aggregating, by the cloud based analytics system, medical resource usage data from a plurality of medical hubs located in different medical facility locations, each communicatively coupled to the cloud based analytics system, the medical resource usage data comprising: data pertaining to medical products and an indication of efficiency based on their usage; disposal records of when the

medical products were disposed of; and for each description of the medical product: location data describing which medical facility said medical product was allocated to; and outcome data pertaining to an outcome of a patient from a procedure that utilized the medical product; determining, by the cloud based analytics medical system, a correlation between positive outcomes from the outcome data and location data of the medical product; generating, by the cloud based analytics medical system, a medical recommendation to change a medical resource usage practice based on the correlation; and causing display in at least one of the medical hubs, by the cloud based analytics medical system, the medical recommendation.

**[2077]** Example 9: The method of Example 8, wherein the disposal records are derived at least in part from disposal bins configured to automatically record an amount of medical products disposed into the bins.

**[2078]** Example 10: The method of any one of Examples 8-9, wherein the outcome data is derived at least in part from operational data transmitted by a medical device used during the procedure.

**[2079]** Example 11: The method of any one of Examples 8-10, wherein the operational data includes a recordation by the medical device of a number of staple firings that were fired by the medical device during the procedure.

**[2080]** Example 12: The method of any one of Examples 8-11, wherein the recommendation comprises a recommendation to substitute use of a first medical product for user of a second medical product during a specific medical procedure.

**[2081]** Example 13: The method of any one of Examples 8-12, wherein the recommendation comprises a recommendation to reduce a number of staple firings that are fired by a medical device during a specific medical procedure.

**[2082]** Example 14: The method of any one of Examples 8-13, wherein the recommendation comprises a recommendation to reduce a rate of use of the medical product during a specific medical procedure.

**[2083]** Example 15: A non-transitory computer readable medium storing computer readable instructions executable by the at least one processor of a cloud-based analytics system to: aggregate medical resource usage data from a plurality of medical hubs located in different medical facility locations, each communicatively coupled to a cloud based analytics system, the medical resource usage data comprising: data pertaining to medical products and an indication of efficiency based on their usage; disposal records of when the medical products were disposed of; and for each description of the medical product: location data describing which medical facility said medical product was allocated to; and outcome data pertaining to an outcome of a patient from a procedure that utilized the medical product; determine a correlation between positive outcomes from the outcome data and location data of the medical product; generate a medical recommendation to change a medical resource usage practice based on the correlation; and cause display of the medical recommendation to at least one medical hub at a local facility.

**[2084]** Example 16: The non-transitory computer readable medium of Example 15, wherein the disposal records are derived at least in part from disposal bins configured to automatically record an amount of medical products disposed into the bins.

**[2085]** Example 17: The non-transitory computer readable medium of any one of Examples 15-16, wherein the outcome data is derived at least in part from operational data transmitted by a medical device used during the procedure.

**[2086]** Example 18: The non-transitory computer readable medium of any one of Examples 15-17, wherein the operational data includes a recordation by the medical device of a number of staple firings that were fired by the medical device during the procedure.

**[2087]** Example 19: The non-transitory computer readable medium of any one of Examples 15-18, wherein the recommendation comprises a recommendation to substitute use of a first medical product for user of a second medical product during a specific medical procedure.

**[2088]** Example 20: The non-transitory computer readable medium of any one of Examples 15-19, wherein the recommendation comprises a recommendation to reduce a number of staple firings that are fired by a medical device during a specific medical procedure.

**[2089]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2090]** Example 1: A cloud-based analytics medical system comprising: at least one processor; at least one memory communicatively coupled to the at least one processor; an input/output interface configured for accessing data from a plurality of medical hub communication devices, each communicatively coupled to at least one surgical instrument; and a database residing in the at least one memory and configured to store the data; the at least one memory storing instructions executable by the at least one processor to: aggregate patient outcome data from the plurality of medical hubs, the patient outcome data comprising: data pertaining to steps performed and corresponding timings for each step in patient procedures; data pertaining to an outcome of each patient procedure performed; data pertaining to medical resources used in the patient procedures; for each data item pertaining to the medical resource: location data describing which medical facility said medical resource was allocated to; and for each data item pertaining to the outcome of the patient procedure: data pertaining to an indication of whether the outcome was a success or failure; aggregate medical resource acquisition data from the plurality of medical hubs; determine a correlation between positive outcomes from the patient outcome data and the resource acquisition data; generate a medical recommendation to change a medical resource acquisition practice based on the correlation; and cause display of the medical recommendation to a plurality of medical hubs located at different medical facilities.

**[2091]** Example 2: The cloud based analytics medical system of Example 1, wherein the at least one memory storing instructions executable by the at least one processor to: evaluate the patient outcome data and the resource acquisition data of a particular medical facility; determine that a level of performance of the particular medical facility is below average compared to other medical facilities, based on a comparison of the evaluated patient outcome data and the resource acquisition data of the particular medical facility to the aggregated patient outcome data; and generate a localized recommendation to change a practice of the particular medical facility.

**[2092]** Example 3: The cloud based analytics medical system of any one of Examples 1-2, wherein the localized

recommendation comprises instructions to revise a medical procedure to account for a surgeon level of experience.

**[2093]** Example 4: The cloud based analytics medical system of any one of Examples 1-3, wherein the localized recommendation comprises instructions to revise resource inventory management to reduce inventory of a first product and increase inventory of a second product.

**[2094]** Example 5: The cloud based analytics medical system of any one of Examples 1-4, wherein the at least one memory storing instructions executable by the at least one processor to: evaluate the patient outcome data and the resource acquisition data of medical facilities belonging to a geographical region; determine that a level of performance of the medical facilities in the geographical region is below average compared to a global average of medical facilities, based on a comparison of the evaluated patient outcome data and the resource acquisition data of the medical facilities in the geographical region to the aggregated patient outcome data; and generate a regionalized recommendation to change a practice of the medical facilities belonging to the geographical region.

**[2095]** Example 6: The cloud based analytics medical system of any one of Examples 1-5, wherein the at least one memory storing instructions executable by the at least one processor to: perform trending analysis indicating an expected change in demographics of a population; and generate a predictive modeling recommendation indicating an instruction to change a medical procedure or inventory of one or more medical products over a period of time, to address the expected change in demographics, based on the trending analysis.

**[2096]** Example 7: The cloud based analytics medical system of any one of Examples 1-6, wherein the at least one memory storing instructions executable by the at least one processor to: compare performance metrics of a first method for conducting a medical procedure with performance metrics of a second method for conducting the same medical procedure; and generate a predictive modeling recommendation indicating an instruction to perform the first method for conducting the medical procedure based on the performance comparison.

**[2097]** Example 8: A non-transitory computer readable medium storing computer readable instructions executable by the at least one processor of a cloud-based analytics system to: aggregate patient outcome data from the plurality of medical hubs, the patient outcome data comprising: data pertaining to steps performed and corresponding timings for each step in patient procedures; data pertaining to an outcome of each patient procedure performed; data pertaining to medical resources used in the patient procedures; for each data item pertaining to the medical resource: location data describing which medical facility said medical resource was allocated to; and for each data item pertaining to the outcome of the patient procedure: data pertaining to an indication of whether the outcome was a success or failure; aggregate medical resource acquisition data from the plurality of medical hubs; determine a correlation between positive outcomes from the patient outcome data and the resource acquisition data; generate a medical recommendation to change a medical resource acquisition practice based on the correlation; and cause display of the medical recommendation to a plurality of medical hubs located at different medical facilities.

**[2098]** Example 9: The non-transitory computer readable medium of Example 8, wherein the instructions are further executable to: evaluate the patient outcome data and the resource acquisition data of a particular medical facility; determine that a level of performance of the particular medical facility is below average compared to other medical facilities, based on a comparison of the evaluated patient outcome data and the resource acquisition data of the particular medical facility to the aggregated patient outcome data; and generate a localized recommendation to change a practice of the particular medical facility.

**[2099]** Example 10: The non-transitory computer readable medium of any one of Examples 8-9, wherein the localized recommendation comprises instructions to revise a medical procedure to account for a surgeon level of experience.

**[2100]** Example 11: The non-transitory computer readable medium of any one of Examples 8-10, wherein the localized recommendation comprises instructions to revise resource inventory management to reduce inventory of a first product and increase inventory of a second product.

**[2101]** Example 12: The non-transitory computer readable medium of any one of Examples 8-11, wherein the instructions are further executable to: evaluate the patient outcome data and the resource acquisition data of medical facilities belonging to a geographical region; determine that a level of performance of the medical facilities in the geographical region is below average compared to a global average of medical facilities, based on a comparison of the evaluated patient outcome data and the resource acquisition data of the medical facilities in the geographical region to the aggregated patient outcome data; and generate a regionalized recommendation to change a practice of the medical facilities belonging to the geographical region.

**[2102]** Example 13: The non-transitory computer readable medium of any one of Examples 8-12, wherein the instructions are further configured to: perform trending analysis indicating an expected change in demographics of a population; and generate a predictive modeling recommendation indicating an instruction to change a medical procedure or inventory of one or more medical products over a period of time, to address the expected change in demographics, based on the trending analysis.

**[2103]** Example 14: The non-transitory computer readable medium of any one of Examples 8-13, wherein the instructions are further configured to: compare performance metrics of a first method for conducting a medical procedure with performance metrics of a second method for conducting the same medical procedure; and generate a predictive modeling recommendation indicating an instruction to perform the first method for conducting the medical procedure based on the performance comparison.

**[2104]** Example 15: A cloud based analytics medical system comprising: at least one processor; at least one memory communicatively coupled to the at least one processor; an input/output interface configured for accessing data from a plurality of medical hub communication devices, each communicatively coupled to at least one surgical instrument; and a database residing in the at least one memory and configured to store the data; wherein the at least one memory storing instructions executable by the at least one processor to: aggregate medical instrument data from the plurality of medical hubs, the medical instrument data comprising: data pertaining to physical and performance parameters of medical devices; for each datum pertaining to the medical device:



usage data pertaining to medical procedures that utilized the medical device; and for each medical procedure; an outcome of the medical procedure; and a status of the condition of the medical device during the medical procedure; determine a correlation between outcomes of the medical procedures and the statuses of the conditions of the medical devices utilized in the respective medical procedures; access live medical procedure data for a live medical procedure, the live medical procedure data comprising a description of the medical devices present in an operating room that is performing the live medical procedure; determine an irregularity in the description of the medical devices present in the live medical procedure, based on the determined correlation between the outcomes and the medical devices utilized; and provide an alert to a medical communication hub that is utilized in the operating room of the live medical procedure.

**[2105]** Example 16: The cloud based analytics medical system of Example 15, wherein the medical devices present in the operating room comprise a manual medical instrument and a robotic medical instrument.

**[2106]** Example 17: The cloud based analytics medical system of any one of Examples 15-16, wherein the at least one processor is further configured to generate a change in firmware or software of a medical device present in the live medical procedure in concert with the provided alert.

**[2107]** Example 18: The cloud based analytics medical system of any one of Examples 15-17, wherein the irregularity comprises use of a medical resource in a medical device present in the live medical procedure that is inconsistent with the aggregated medical instrument data pertaining to the medical procedure.

**[2108]** Example 19: The cloud based analytics medical system of any one of Examples 15-18, wherein the alert comprises an instruction to change a firing or clamping speed of a medical device present in the live medical procedure.

**[2109]** Example 20: The cloud based analytics medical system of any one of Examples 15-19, wherein the alert comprises an instruction to change an ultrasonic blade length of a medical device present in the live medical procedure.

**[2110]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2111]** Example 1: A cloud based analytics medical system comprising: at least one processor; at least one memory communicatively coupled to the at least one processor; an input/output interface configured for accessing data from a plurality of medical hub communication devices, each communicatively coupled to at least one surgical instrument; and a database residing in the at least one memory and configured to store the data; the at least one memory storing instructions executable by the at least one processor to: generate common medical usage patterns of medical devices based on an aggregation of usage data for the medical devices from the plurality of medical hubs; aggregate patient outcome data from the plurality of medical hubs, the patient outcome data comprising: data pertaining to steps performed and corresponding timings for each step in patient procedures; data pertaining to allocation of medical resources used in the patient procedures; for each datum pertaining to the medical resource: location data indicating which medical facility said medical resource was allocated to; and for each datum pertaining to the patient procedure: data indicative of

the outcome of the patient procedure; data indicative of a biographical characterization about the patient; and data indicative of a physiologic characterization about the patient; for data indicative of a positive outcome of the patient procedure, determine a biographical characterization or physiologic difference about the patient compared to biographical or physiologic characterization data in common medical usage patterns; determine a customized change in the medical usage pattern of the medical devices for the medical facility associated with the biographical characterization or physiologic difference; and output a recommendation of the customized change to the medical facility associated with the biographical characterization or physiologic difference.

**[2112]** Example 2: The cloud based analytics medical system of Example 1, wherein the customized change comprises a change to a device setting in a medical device.

**[2113]** Example 3: The cloud based analytics medical system of any one of Examples 1-2, wherein the customized change comprises a change in orientation to how a medical device is handled during a medical procedure.

**[2114]** Example 4: The cloud based analytics medical system of any one of Examples 1-3, wherein the customized change comprises a change of when a medical device is used during a medical procedure.

**[2115]** Example 5: The cloud based analytics medical system of any one of Examples 1-4, wherein the customized change comprises a change in a control algorithm of a medical device.

**[2116]** Example 6: The cloud based analytics medical system of any one of Examples 1-5, wherein the customized change comprises a substitution of a first medical device for a second medical device during a medical procedure.

**[2117]** Example 7: The cloud based analytics medical system of any one of Examples 1-6, wherein the at least one processor is further configured to cause display of quantitative metrics illustrating an estimate of superior results when the recommended change is adopted.

**[2118]** Example 8: A method of a cloud based analytics medical system for improving medical procedures on an individualized basis, the method comprising: generating, by the cloud based analytics medical system, common medical usage patterns of medical devices based on an aggregation of usage data for the medical devices from a plurality of medical hubs communicatively coupled to the cloud based analytics medical system; aggregating, by the cloud based analytics medical system, patient outcome data from the plurality of medical hubs, the patient outcome data comprising: data pertaining to steps performed and corresponding timings for each step in patient procedures; data pertaining to allocation of medical resources used in the patient procedures; for each datum pertaining to the medical resource: location data indicating which medical facility said medical resource was allocated to; and for each datum pertaining to the patient procedure: data indicative of the outcome of the patient procedure; data indicative of a biographical characterization about the patient; and data indicative of a physiologic characterization about the patient; for data indicative of a positive outcome of the patient procedure, determining, by the cloud based analytics medical system, a biographical characterization or physiologic difference about the patient compared to biographical or physiologic characterization data in common medical usage patterns; determining, by the cloud based analytics

medical system, a customized change in the medical usage pattern of the medical devices for the medical facility associated with the biographical characterization or physiologic difference; and outputting, by the cloud based analytics medical system, a recommendation of the customized change to the medical facility associated with the biographical characterization or physiologic difference.

**[2119]** Example 9: The method of Example 8, wherein the customized change comprises a change to a device setting in a medical device.

**[2120]** Example 10: The method of any one of Examples 8-9, wherein the customized change comprises a change in orientation to how a medical device is handled during a medical procedure.

**[2121]** Example 11: The method of any one of Examples 8-10, wherein the customized change comprises a change of when a medical device is used during a medical procedure.

**[2122]** Example 12: The method of any one of Examples 8-11, wherein the customized change comprises a change in a control algorithm of a medical device.

**[2123]** Example 13: The method of any one of Examples 8-12, wherein the customized change comprises a substitution of a first medical device for a second medical device during a medical procedure.

**[2124]** Example 14: The method of any one of Examples 8-13, wherein the at least one processor is further configured to cause display of quantitative metrics illustrating an estimate of superior results when the recommended change is adopted.

**[2125]** Example 15: A non-transitory computer readable medium storing computer readable instructions executable by at least one processor of a cloud-based analytics system to: generate common medical usage patterns of medical devices based on an aggregation of usage data for the medical devices from a plurality of medical hubs communicatively coupled to the cloud-based analytics system; aggregate patient outcome data from the plurality of medical hubs, the patient outcome data comprising: data pertaining to steps performed and corresponding timings for each step in patient procedures; data pertaining to allocation of medical resources used in the patient procedures; for each datum pertaining to the medical resource: location data indicating which medical facility said medical resource was allocated to; and for each datum pertaining to the patient procedure: data indicative of the outcome of the patient procedure; data indicative of a biographical characterization about the patient; and data indicative of a physiologic characterization about the patient compared to biographical or physiologic characterization data in common medical usage patterns; determine a customized change in the medical usage pattern of the medical devices for the medical facility associated with the biographical characterization or physiologic difference; and output a recommendation of the customized change to the medical facility associated with the biographical characterization or physiologic difference.

**[2126]** Example 16: The non-transitory computer readable medium of Example 15, wherein the customized change comprises a change to a device setting in a medical device.

**[2127]** Example 17: The non-transitory computer readable medium of any one of Examples 15-16, wherein the cus-

tomized change comprises a change in orientation to how a medical device is handled during a medical procedure.

**[2128]** Example 18: The non-transitory computer readable medium of any one of Examples 15-17, wherein the customized change comprises a change of when a medical device is used during a medical procedure.

**[2129]** Example 19: The non-transitory computer readable medium of any one of Examples 15-18, wherein the customized change comprises a change in a control algorithm of a medical device.

**[2130]** Example 20: The non-transitory computer readable medium of any one of Examples 15-19, wherein the customized change comprises a substitution of a first medical device for a second medical device during a medical procedure.

**[2131]** Various additional aspects of the subject matter described herein are set out in the following numbered examples:

**[2132]** Example 1: A cloud based security system for a medical data network, the security system comprising: at least one processor; at least one memory communicatively coupled to the processor; an input/output interface configured for accessing data from a plurality of medical hubs, each communicatively coupled to at least one surgical instrument; and a database residing in the at least one memory and configured to store the data; the at least one memory storing instructions executable by the at least one processor to: identify a first security threat by a first medical instrument communicatively coupled to a first medical hub located at a first medical facility; determine that a second security threat is present at a second medical hub located at a second medical facility, based on at least one common characteristic between the first medical instrument and a second medical instrument communicatively coupled to the second medical hub; and provide an alert to the second medical facility about the second security threat.

**[2133]** Example 2: The cloud based security system of Example 1, wherein identifying the first security threat comprises determining that an identification parameter of the first medical instrument is invalid.

**[2134]** Example 3: The cloud based security system of any of Examples 1-2, wherein identifying the first security threat comprises detecting that the first medical instrument is transmitting a virus.

**[2135]** Example 4: The cloud based security system of any of Examples 1-3, wherein identifying the first security threat comprises determining that the first medical instrument fails an authentication protocol.

**[2136]** Example 5: The cloud based security system of any of Examples 1-4, wherein the at least one processor is further programmed to lock out the first medical instrument from operating with the first medical hub and every other medical hub in the first medical facility.

**[2137]** Example 6: The cloud based security system of claim Examples 1-5, wherein the at least one processor is further configured to: analyze alert data associated with the first medical facility, in response to identifying the first security threat; determine an irregularity with the alert data associated with the first medical facility compared to alert data associated with other medical facilities; and determine a revised security procedure for the first medical facility in response to the determined irregularity.

**[2138]** Example 7: The cloud based security system of any of Examples 1-6, wherein the at least one common charac-

teristic comprises a common manufacturer between the first medical device and the second medical device.

**[2139]** Example 8: The cloud based security system of any of Examples 1-7, wherein the at least one common characteristic comprises a first identification parameter of the first medical device and a second identification parameter of the second medical device both within an invalid range.

**[2140]** Example 9: A method of a cloud based security system of a medical data network for improving security and authentication of the medical data network, the medical data network further comprising a plurality of medical hubs each communicatively coupled to the cloud based security system and at least one surgical instrument, the method comprising: identifying, by the cloud based security system, a first security threat by a first medical instrument communicatively coupled to a first medical hub located at a first medical facility; determining, by the cloud based security system, that a second security threat is present at a second medical hub located at a second medical facility, based on at least one common characteristic between the first medical instrument and a second medical instrument communicatively coupled to the second medical hub; and providing, by the cloud based security system, an alert to the second medical facility about the second security threat.

**[2141]** Example 10: The method of Example 9, wherein identifying the first security threat comprises determining that an identification parameter of the first medical instrument is invalid.

**[2142]** Example 11: The method of any of Examples 9-10, wherein identifying the first security threat comprises detecting that the first medical instrument is transmitting a virus.

**[2143]** Example 12: The method of any of Examples 9-11, wherein identifying the first security threat comprises determining that the first medical instrument fails an authentication protocol.

**[2144]** Example 13: The method of any of Examples 9-12, further comprising locking out the first medical instrument from operating with the first medical hub and every other medical hub in the first medical facility.

**[2145]** Example 14: The method of any of Examples 9-13, further comprising: analyzing alert data associated with the first medical facility, in response to identifying the first security threat; determining an irregularity with the alert data associated with the first medical facility compared to alert data associated with other medical facilities; and determining a revised security procedure for the first medical facility in response to the determined irregularity.

**[2146]** Example 15: The method of any of Examples 9-14, wherein the at least one common characteristic comprises a common manufacturer between the first medical device and the second medical device.

**[2147]** Example 16: The method of any of Examples 9-15, wherein the at least one common characteristic comprises a first identification parameter of the first medical device and a second identification parameter of the second medical device both within an invalid range.

**[2148]** Example 17: A non-transitory computer readable medium comprising instructions that, when executed by a processor of a cloud based security system of a medical data network, cause the processor to perform operations comprising: identifying a first security threat by a first medical instrument communicatively coupled to a first medical hub located at a first medical facility; determining that a second security threat is present at a second medical hub located at

a second medical facility, based on at least one common characteristic between the first medical instrument and a second medical instrument communicatively coupled to the second medical hub; and providing an alert to the second medical facility about the second security threat.

**[2149]** Example 18: The non-transitory computer readable medium of Example 17, wherein identifying the first security threat comprises determining that an identification parameter of the first medical instrument is invalid, detecting that the first medical instrument is transmitting a virus, or determining that the first medical instrument fails an authentication protocol.

**[2150]** Example 19: The non-transitory computer readable medium of any of Examples 17-18, wherein the operations further comprise locking out the first medical instrument from operating with the first medical hub and every other medical hub in the first medical facility.

**[2151]** Example 20: The non-transitory computer readable medium of any of Examples 17-19, wherein the operations further comprise: analyzing alert data associated with the first medical facility, in response to identifying the first security threat; determining an irregularity with the alert data associated with the first medical facility compared to alert data associated with other medical facilities; and determining a revised security procedure for the first medical facility in response to the determined irregularity.

**[2152]** Various additional aspects of the subject matter described herein are set out in the following numbered examples.

**[2153]** Example 1: A cloud based analytics medical system comprising: at least one processor; at least one memory communicatively coupled to the at least one processor; an input/output interface configured for accessing data from a plurality of surgical hubs, each of the plurality of surgical hubs communicatively coupled to at least one surgical instrument and the at least one processor; and a database residing in the at least one memory and configured to store the data; and wherein the at least one memory is configured to store instructions executable by the at least one processor to receive critical data from the plurality of surgical hubs, wherein the plurality of surgical hubs determine critical data based on screening criteria; determine a priority status of the critical data; route the critical data to a cloud storage location residing within the at least one memory; and determine a response to the critical data based on an operational characteristic indicated by the critical data, wherein a time component of the response is determined based on the priority status.

**[2154]** Example 2: The cloud based analytics medical system of Example 1, wherein the screening criteria comprises one or more of: severity, unexpectedness, suspiciousness, and security.

**[2155]** Example 3: The cloud based analytics medical system of any one of Examples 1-2, wherein the severity screening criteria comprises an extent of a perioperative device failure and a transition to non-standard post-operation treatment of a patient.

**[2156]** Example 4: The cloud based analytics medical system of any one of Examples 1-3, wherein the at least one memory is further configured to store instructions executable by the at least one processor to request the plurality of surgical hubs obtain additional data pertaining to the critical data.

[2157] Example 5: The cloud based analytics medical system of Example 4, wherein the at least one memory is further configured to store instructions executable by the at least one processor to request additional data based on a plurality of trigger conditions.

[2158] Example 6: The cloud based analytics medical system of Example 5, wherein the plurality of trigger conditions comprise one or more of: exceeding a predetermined unexpectedness threshold, unauthorized modification of the critical data, unsecure communication of data, placement of the at least one surgical instrument on a watch list.

[2159] Example 7: The cloud based analytics medical system of any one of Examples 1-6, wherein the critical data comprises aggregated data from the plurality of surgical hubs.

[2160] Example 8: The cloud based analytics medical system of any one of Examples 1-7, wherein the at least one processor transmits the critical data to the database.

[2161] Example 9: A non-transitory computer readable medium storing computer readable instructions executable by the at least one processor of a cloud-based analytics system to: receive critical data from a plurality of surgical hubs, wherein the plurality of surgical hubs determine critical data based on screening criteria and each of the plurality of surgical hubs are communicatively coupled to at least one surgical instrument and the at least one processor; determine a priority status of the critical data; route the critical data to a cloud storage location residing within at least one memory coupled to the at least one processor; and determine a response to the critical data based on an operational characteristic indicated by the critical data, wherein a time component of the response is determined based on the priority status.

[2162] Example 10: The non-transitory computer readable medium of Example 9, wherein the priority status is determined by the at least one processor based on one or more of: the critical data corresponds to the at least one surgical instrument placed on a watch list, the critical data corresponds to an automated response; the critical data corresponds to a notification response, the critical data corresponds to an urgent response.

[2163] Example 11: The non-transitory computer readable medium of Example 10, wherein the at least one surgical instrument is placed on the watch list based on one or more of: counterfeit products, deviation in surgical instrument performance, and unauthorized usage.

[2164] Example 12: The non-transitory computer readable medium of any one of Examples 10-11, wherein the automated response comprises a corrective and preventive action response.

[2165] Example 13: The non-transitory computer readable medium of any one of Examples 1-9, wherein the at least one processor stores the critical data in a hold list in the at least one memory and validates the accuracy of the critical data.

[2166] Example 14: A cloud based analytics medical system comprising: at least one processor; at least one memory communicatively coupled to the at least one processor; an input/output interface configured for accessing data from a plurality of surgical hubs, each of the plurality of surgical hubs communicatively coupled to at least one surgical instrument and the at least one processor; and a database residing in the at least one memory and configured to store the data; and wherein the at least one memory is configured to store instructions executable by the at least one processor

to: receive critical data from the plurality of surgical hubs, wherein the plurality of surgical hubs determine critical data based on screening criteria; determine a priority status of the critical data; route the critical data to a cloud storage location residing within the at least one memory; request the plurality of surgical hubs obtain additional data pertaining to the critical data based on a plurality of trigger conditions; determine the cause of an irregularity corresponding to the critical data and additional data; and determine a response to the irregularity, wherein a time component of the response is determined based on the priority status.

[2167] Example 15: The cloud based analytics medical system of Example 14, wherein the at least one processor responds to the irregularity by transmitting a signal to the at least one surgical instrument corresponding to the irregularity, wherein the signal causes an operational lockout of the at least one surgical instrument.

[2168] Example 16: The cloud based analytics medical system of any one of Examples 14-15, wherein the at least one processor requests the plurality of surgical hubs obtain the additional data for a predetermined amount of time.

[2169] Example 17: The cloud based analytics medical system of Example 16, wherein the at least one processor requests the plurality of surgical hubs obtain the additional data for the predetermined amount of time based on an occurrence of a predetermined medical event.

[2170] Example 18: The cloud based analytics medical system of any one of Examples 14-17, wherein the at least one processor responds to the irregularity by monitoring patient outcomes corresponding to irregularity for a predetermined amount of time.

[2171] Example 19: The cloud based analytics medical system of any one of Examples 14-18, wherein the at least one processor responds to the irregularity by transmitting a signal to the plurality of surgical hubs corresponding to the irregularity to indicate a corrective action.

[2172] Example 20: The cloud based analytics medical system of any one of Examples 14-19, wherein the at least one processor transmits the critical data to the database for aggregation of the critical data, wherein the critical data is classified as corresponding to a positive patient outcome or a negative patient outcome.

[2173] Various additional aspects of the subject matter described herein are set out in the following numbered examples:

[2174] Example 1: A surgical system, comprising: a surgical hub couplable with a plurality of inventory items of an institution, wherein the plurality of inventory items include medical devices, and wherein the surgical hub comprises: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to communicate with the plurality of inventory items; and a cloud-based analytics system communicatively coupled to the surgical hub, wherein the cloud-based analytics system comprises: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to: receive, via the surgical hub, data associated with the plurality of inventory items, wherein the received data comprises a unique identifier for each inventory item; determine whether each inventory item is available for use based on its respective unique identifier and system-defined constraints, wherein the system-defined constraints comprise at least one use restriction; generate a cloud interface for the institution, wherein the institution's cloud interface

comprises a plurality of user-interface elements, wherein at least one user-interface element enables selection of one or more than one surgical procedure to be performed, and wherein after selection of a surgical procedure, via the at least one user-interface element, the availability of each inventory item associated with the selected surgical procedure is dynamically generated on the institution's cloud interface; and transmit an alert for each inventory item determined as not available based on the system-defined constraints, wherein the alert is displayable on at least one of the institution's cloud interface or the inventory item.

**[2175]** Example 2: The surgical system of Example 1, wherein the system-defined constraints further comprise a list of unauthorized devices, and wherein the instructions are further executable by the processor of the cloud-based analytics system to: prevent each unauthorized device from being utilized in the surgical system to perform surgical procedures.

**[2176]** Example 3: The surgical system of any one of Examples 1-2, wherein the instructions are further executable by the processor of the cloud-based analytics system to: allow an unauthorized device to perform surgical procedures if at least one of the unauthorized device is subject to a usage fee, the unauthorized device is subject to limited functionality, or the unauthorized device is subject to secondary system-defined constraints.

**[2177]** Example 4: The surgical system of any one of Examples 1-3, wherein the instructions are further executable by the processor of the surgical hub to communicate wirelessly with the plurality of inventory items.

**[2178]** Example 5: The surgical system of any one of Examples 1-4, wherein the plurality of inventory items further comprises a surgical instrument to perform the selected surgical procedure, wherein the surgical instrument comprises a plurality of modular components, and wherein the instructions are further executable by the processor of the cloud-based analytics system to: determine whether each modular component of the surgical instrument is available for use based on its respective unique identifier and the system-defined constraints.

**[2179]** Example 6: The surgical system of any one of Examples 1-5, wherein the instructions are further executable by the processor of the cloud-based analytics system to: determine that a unique identifier, associated with a first modular component of the plurality of modular components, indicates the first modular component as at least one of counterfeit or defective; and transmit an alert displayable on a user interface of the first modular component.

**[2180]** Example 7: The surgical system of any one of Examples 1-6, wherein the cloud-based analytics system further comprises a database, and wherein the instructions are further executable by the processor of the cloud-based analytics system to: update a list of unauthorized devices stored on the database with the unique identifier of the first modular component.

**[2181]** Example 8: The surgical system of any one of Examples 1-7, wherein the instructions are further executable by the processor of the cloud-based analytics system to: determine at least one alternative modular component available, based on system-defined constraints, to perform the selected surgical procedure; and transmit an alert displayable on at least one of the institution's cloud interface or the user interface of the first modular component.

**[2182]** Example 9: The surgical system of any one of Examples 1-8, wherein a system-defined constraint comprises an expiration date associated with each modular component of the surgical instrument, and wherein the instructions are further executable by the processor of the cloud-based analytics system to: determine that a first modular component of the surgical instrument has exceeded an expiration date; transmit an alert displayable on a user interface of the first modular component, wherein the alert comprises a warning that the expiration date has been exceeded; and receive an input, via the user interface of the first modular component, to bypass the exceeded expiration date.

**[2183]** Example 10: The surgical system of Example 9, wherein the exceeded expiration date is associated with a control program stored on the first modular component.

**[2184]** Example 11: The surgical system of Example 5, wherein the at least one use restriction comprises a usable life metric associated with each modular component of the surgical instrument, and wherein the instructions are further executable by the processor of the cloud-based analytics system to: access a current usage parameter associated with each modular component of the surgical instrument; determine that a first modular component of the surgical instrument has exceeded its associated usable life metric; and transmit an alert displayable on a user interface of the first modular component.

**[2185]** Example 12: The surgical system of any one of Examples 1-11, further comprising: at least one modular component couplable with the surgical hub, wherein each modular component comprises: a processor; and a memory coupled to the processor, the memory storing instructions executable by the processor to communicate its identifier and at least one of a usage parameter or a usable life metric to the surgical hub.

**[2186]** Example 13: The surgical system of any one of Examples 1-12, wherein the instructions are further executable by the processor of each modular component to relay at least one of an identifier, a usage parameter, or a usable life metric received from another modular component to the surgical hub.

**[2187]** Example 14: The surgical system of any one of Examples 1-13, wherein each modular component further comprises a user interface, and wherein the instructions are further executable by the processor of each modular component to: display, via its user interface, an alert transmitted by the cloud-based analytics system, wherein the alert comprises a link associated with a violated system-defined constraint; receive, via its user interface, a selection of the link; receive, via its user interface, a selection to waive a flexible system-defined constraint; and transmit the selection to waive the flexible system-defined constraint to the cloud-based analytics system.

**[2188]** Example 15: A surgical system, comprising: a surgical hub couplable with a plurality of inventory items of an institution, wherein the plurality of inventory items include medical devices, and wherein the surgical hub comprises a control circuit configured to communicate with the plurality of inventory items; and a cloud-based analytics system communicatively coupled to the surgical hub, wherein the cloud-based analytics system comprises a control circuit configured to: receive, via the surgical hub, data associated with the plurality of inventory items, wherein the received data comprises a unique identifier for each inven-

tory item; determine whether each inventory item is available for use based on its respective unique identifier and system-defined constraints, wherein the system-defined constraints comprise at least one use restriction; generate a cloud interface for the institution, wherein the institution's cloud interface comprises a plurality of user-interface elements, wherein at least one user-interface element enables selection of one or more than one surgical procedure to be performed, and wherein after selection of a surgical procedure, via the at least one user-interface element, the availability of each inventory item associated with the selected surgical procedure is dynamically generated on the institution's cloud interface; and transmit an alert for each inventory item determined as not available based on the system-defined constraints, wherein the alert is displayable on at least one of the institution's cloud interface or the inventory item.

**[2189]** Example 16: The surgical system of Example 15, wherein the system-defined constraints further comprise a list of unauthorized devices, and wherein the control circuit of the cloud-based analytics system is further configured to: prevent each unauthorized device from being utilized in the surgical system to perform surgical procedures; or allow an unauthorized device to perform surgical procedures if at least one of the unauthorized device is subject to a usage fee, the unauthorized device is subject to limited functionality, or the unauthorized device is subject to secondary system-defined constraints.

**[2190]** Example 17: The surgical system of any one of Examples 15-16, wherein the plurality of inventory items further comprises a surgical instrument to perform the selected surgical procedure, wherein the surgical instrument comprises a plurality of modular components, and wherein the control circuit of the cloud-based analytics system is further configured to: determine whether each modular component of the surgical instrument is available for use based on its respective unique identifier and the system-defined constraints.

**[2191]** Example 18: The surgical system of any one of Examples 15-17, further comprising: at least one modular component couplable with the surgical hub, wherein each modular component comprises a control circuit configured to communicate its identifier and at least one of a usage parameter or a usable life metric to the surgical hub.

**[2192]** Example 19: The surgical system of any one of Examples 15-18, wherein each modular component further comprises a user interface, and wherein the control circuit of each modular component is further configured to: display, via its user interface, an alert transmitted by the cloud-based analytics system, wherein the alert comprises a link associated with a violated system-defined constraint; receive, via its user interface, a selection of the link; receive, via its user interface, a selection to waive a flexible system-defined constraint; and transmit the selection to waive the flexible system-defined constraint to the cloud-based analytics system.

**[2193]** Example 20: A non-transitory computer readable medium storing computer readable instructions which, when executed, causes a cloud-based analytics system to: receive, via a surgical hub, data associated with a plurality of inventory items of an institution, wherein the plurality of inventory items include medical devices, wherein the received data comprises a unique identifier for each inventory item, and wherein each unique identifier is received by

the surgical hub in a communication with each inventory item; determine whether each inventory item is available for use based on its respective unique identifier and system-defined constraints, wherein the system-defined constraints comprise at least one use restriction; generate a cloud interface for the institution, wherein the institution's cloud interface comprises a plurality of user-interface elements, wherein at least one user-interface element enables selection of one or more than one surgical procedure to be performed, and wherein after selection of a surgical procedure, via the at least one user-interface element, the availability of each inventory item associated with the selected surgical procedure is dynamically generated on the institution's cloud interface; and transmit an alert for each inventory item determined as not available based on the system-defined constraints, wherein the alert is displayable on at least one of the institution's cloud interface or the inventory item.

**[2194]** While several forms have been illustrated and described, it is not the intention of the applicant to restrict or limit the scope of the appended claims to such detail. Numerous modifications, variations, changes, substitutions, combinations, and equivalents to those forms may be implemented and will occur to those skilled in the art without departing from the scope of the present disclosure. Moreover, the structure of each element associated with the described forms can be alternatively described as a means for providing the function performed by the element. Also, where materials are disclosed for certain components, other materials may be used. It is therefore to be understood that the foregoing description and the appended claims are intended to cover all such modifications, combinations, and variations as falling within the scope of the disclosed forms. The appended claims are intended to cover all such modifications, variations, changes, substitutions, modifications, and equivalents.

**[2195]** The foregoing detailed description has set forth various forms of the devices and/or processes via the use of block diagrams, flowcharts, and/or examples. Insofar as such block diagrams, flowcharts, and/or examples contain one or more functions and/or operations, it will be understood by those within the art that each function and/or operation within such block diagrams, flowcharts, and/or examples can be implemented, individually and/or collectively, by a wide range of hardware, software, firmware, or virtually any combination thereof. Those skilled in the art will recognize that some aspects of the forms disclosed herein, in whole or in part, can be equivalently implemented in integrated circuits, as one or more computer programs running on one or more computers (e.g., as one or more programs running on one or more computer systems), as one or more programs running on one or more processors (e.g., as one or more programs running on one or more microprocessors), as firmware, or as virtually any combination thereof, and that designing the circuitry and/or writing the code for the software and or firmware would be well within the skill of one of skilled in the art in light of this disclosure. In addition, those skilled in the art will appreciate that the mechanisms of the subject matter described herein are capable of being distributed as one or more program products in a variety of forms and that an illustrative form of the subject matter described herein applies regardless of the particular type of signal-bearing medium used to actually carry out the distribution.

**[2196]** Instructions used to program logic to perform various disclosed aspects can be stored within a memory in the system, such as DRAM, cache, flash memory, or other storage. Furthermore, the instructions can be distributed via a network or by way of other computer-readable media. Thus a machine-readable medium may include any mechanism for storing or transmitting information in a form readable by a machine (e.g., a computer), but is not limited to, floppy diskettes, optical disks, CD-ROMs, magneto-optical disks, ROM, RAM, EPROM, EEPROM, magnetic or optical cards, flash memory, or tangible, machine-readable storage used in the transmission of information over the Internet via electrical, optical, acoustical, or other forms of propagated signals (e.g., carrier waves, infrared signals, digital signals). Accordingly, the non-transitory computer-readable medium includes any type of tangible machine-readable medium suitable for storing or transmitting electronic instructions or information in a form readable by a machine (e.g., a computer).

**[2197]** As used in any aspect herein, the term “control circuit” may refer to, for example, hardwired circuitry, programmable circuitry (e.g., a computer processor comprising one or more individual instruction processing cores, processing unit, processor, microcontroller, microcontroller unit, controller, DSP, PLD, programmable logic array (PLA), or FPGA), state machine circuitry, firmware that stores instructions executed by programmable circuitry, and any combination thereof. The control circuit may, collectively or individually, be embodied as circuitry that forms part of a larger system, for example, an integrated circuit, an application-specific integrated circuit (ASIC), a system on-chip (SoC), desktop computers, laptop computers, tablet computers, servers, smart phones, etc. Accordingly, as used herein, “control circuit” includes, but is not limited to, electrical circuitry having at least one discrete electrical circuit, electrical circuitry having at least one integrated circuit, electrical circuitry having at least one application-specific integrated circuit, electrical circuitry forming a general-purpose computing device configured by a computer program (e.g., a general-purpose computer configured by a computer program which at least partially carries out processes and/or devices described herein, or a microprocessor configured by a computer program which at least partially carries out processes and/or devices described herein), electrical circuitry forming a memory device (e.g., forms of random access memory), and/or electrical circuitry forming a communications device (e.g., a modem, communications switch, or optical-electrical equipment). Those having skill in the art will recognize that the subject matter described herein may be implemented in an analog or digital fashion or some combination thereof.

**[2198]** As used in any aspect herein, the term “logic” may refer to an app, software, firmware, and/or circuitry configured to perform any of the aforementioned operations. Software may be embodied as a software package, code, instructions, instruction sets, and/or data recorded on non-transitory computer-readable storage medium. Firmware may be embodied as code, instructions, instruction sets, and/or data that are hard-coded (e.g., non-volatile) in memory devices.

**[2199]** As used in any aspect herein, the terms “component,” “system,” “module,” and the like can refer to a computer-related entity, either hardware, a combination of hardware and software, software, or software in execution.

**[2200]** As used in any aspect herein, an “algorithm” refers to a self-consistent sequence of steps leading to a desired result, where a “step” refers to a manipulation of physical quantities and/or logic states which may, though need not necessarily, take the form of electrical or magnetic signals capable of being stored, transferred, combined, compared, and otherwise manipulated. It is common usage to refer to these signals as bits, values, elements, symbols, characters, terms, numbers, or the like. These and similar terms may be associated with the appropriate physical quantities and are merely convenient labels applied to these quantities and/or states.

**[2201]** A network may include a packet-switched network. The communication devices may be capable of communicating with each other using a selected packet-switched network communications protocol. One example communications protocol may include an Ethernet communications protocol which may be capable permitting communication using a Transmission Control Protocol/IP. The Ethernet protocol may comply or be compatible with the Ethernet standard published by the Institute of Electrical and Electronics Engineers (IEEE) titled “IEEE 802.3 Standard,” published in December 2008 and/or later versions of this standard. Alternatively or additionally, the communication devices may be capable of communicating with each other using an X.25 communications protocol. The X.25 communications protocol may comply or be compatible with a standard promulgated by the International Telecommunication Union-Telecommunication Standardization Sector (ITU-T). Alternatively or additionally, the communication devices may be capable of communicating with each other using a frame relay communications protocol. The frame relay communications protocol may comply or be compatible with a standard promulgated by Consultative Committee for International Telegraph and Telephone (CCITT) and/or the American National Standards Institute (ANSI). Alternatively or additionally, the transceivers may be capable of communicating with each other using an Asynchronous Transfer Mode (ATM) communications protocol. The ATM communications protocol may comply or be compatible with an ATM standard published by the ATM Forum, titled “ATM-MPLS Network Interworking 2.0,” published August 2001, and/or later versions of this standard. Of course, different and/or after-developed connection-oriented network communication protocols are equally contemplated herein.

**[2202]** Unless specifically stated otherwise as apparent from the foregoing disclosure, it is appreciated that, throughout the foregoing disclosure, discussions using terms such as “processing,” “computing,” “calculating,” “determining,” “displaying,” or the like, refer to the action and processes of a computer system, or similar electronic computing device, that manipulates and transforms data represented as physical (electronic) quantities within the computer system’s registers and memories into other data similarly represented as physical quantities within the computer system memories or registers or other such information storage, transmission, or display devices.

**[2203]** One or more components may be referred to herein as “configured to,” “configurable to,” “operable/operative to,” “adapted/adaptable,” “able to,” “conformable/conformed to,” etc. Those skilled in the art will recognize that “configured to” can generally encompass active-state com-

ponents, inactive-state components, and/or standby-state components, unless context requires otherwise.

**[2204]** The terms “proximal” and “distal” are used herein with reference to a clinician manipulating the handle portion of the surgical instrument. The term “proximal” refers to the portion closest to the clinician, and the term “distal” refers to the portion located away from the clinician. It will be further appreciated that, for convenience and clarity, spatial terms such as “vertical,” “horizontal,” “up,” and “down” may be used herein with respect to the drawings. However, surgical instruments are used in many orientations and positions, and these terms are not intended to be limiting and/or absolute.

**[2205]** Those skilled in the art will recognize that, in general, terms used herein, and especially in the appended claims (e.g., bodies of the appended claims), are generally intended as “open” terms (e.g., the term “including” should be interpreted as “including, but not limited to”; the term “having” should be interpreted as “having at least”; the term “includes” should be interpreted as “includes, but is not limited to”). It will be further understood by those within the art that if a specific number of an introduced claim recitation is intended, such an intent will be explicitly recited in the claim, and in the absence of such recitation, no such intent is present. For example, as an aid to understanding, the following appended claims may contain usage of the introductory phrases “at least one” and “one or more” to introduce claim recitations. However, the use of such phrases should not be construed to imply that the introduction of a claim recitation by the indefinite articles “a” or “an” limits any particular claim containing such introduced claim recitation to claims containing only one such recitation, even when the same claim includes the introductory phrases “one or more” or “at least one” and indefinite articles such as “a” or “an” (e.g., “a” and/or “an” should typically be interpreted to mean “at least one” or “one or more”); the same holds true for the use of definite articles used to introduce claim recitations.

**[2206]** In addition, even if a specific number of an introduced claim recitation is explicitly recited, those skilled in the art will recognize that such recitation should typically be interpreted to mean at least the recited number (e.g., the bare recitation of “two recitations,” without other modifiers, typically means at least two recitations or two or more recitations). Furthermore, in those instances where a convention analogous to “at least one of A, B, and C, etc.” is used, in general, such a construction is intended in the sense that one having skill in the art would understand the convention (e.g., “a system having at least one of A, B, and C” would include, but not be limited to, systems that have A alone, B alone, C alone, A and B together, B and C together, and/or A, B, and C together). In those instances where a convention analogous to “at least one of A, B, or C, etc.” is used, in general, such a construction is intended in the sense that one having skill in the art would understand the convention (e.g., “a system having at least one of A, B, or C” would include, but not be limited to, systems that have A alone, B alone, C alone, A and B together, A and C together, B and C together, and/or A, B, and C together). It will be further understood by those within the art that typically a disjunctive word and/or phrase presenting two or more alternative terms, whether in the description, claims, or drawings, should be understood to contemplate the possibilities of including one of the terms,

either of the terms, or both terms, unless context dictates otherwise. For example, the phrase “A or B” will be typically understood to include the possibilities of “A” or “B” or “A and B.”

**[2207]** With respect to the appended claims, those skilled in the art will appreciate that recited operations therein may generally be performed in any order. Also, although various operational flow diagrams are presented in a sequence(s), it should be understood that the various operations may be performed in other orders than those which are illustrated or may be performed concurrently. Examples of such alternate orderings may include overlapping, interleaved, interrupted, reordered, incremental, preparatory, supplemental, simultaneous, reverse, or other variant orderings, unless context dictates otherwise. Furthermore, terms like “responsive to,” “related to,” or other past-tense adjectives are generally not intended to exclude such variants, unless context dictates otherwise.

**[2208]** It is worthy to note that any reference to “one aspect,” “an aspect,” “an exemplification,” “one exemplification,” and the like means that a particular feature, structure, or characteristic described in connection with the aspect is included in at least one aspect. Thus, appearances of the phrases “in one aspect,” “in an aspect,” “in an exemplification,” and “in one exemplification” in various places throughout the specification are not necessarily all referring to the same aspect. Furthermore, the particular features, structures, or characteristics may be combined in any suitable manner in one or more aspects.

**[2209]** Any patent application, patent, non-patent publication, or other disclosure material referred to in this specification and/or listed in any Application Data Sheet is incorporated by reference herein, to the extent that the incorporated materials are not inconsistent herewith. As such, and to the extent necessary, the disclosure as explicitly set forth herein supersedes any conflicting material incorporated herein by reference. Any material, or portion thereof, that is said to be incorporated by reference herein but which conflicts with existing definitions, statements, or other disclosure material set forth herein will only be incorporated to the extent that no conflict arises between that incorporated material and the existing disclosure material.

**[2210]** In summary, numerous benefits have been described which result from employing the concepts described herein. The foregoing description of the one or more forms has been presented for purposes of illustration and description. It is not intended to be exhaustive or limiting to the precise form disclosed. Modifications or variations are possible in light of the above teachings. The one or more forms were chosen and described in order to illustrate principles and practical application to thereby enable one of ordinary skill in the art to utilize the various forms and with various modifications as are suited to the particular use contemplated. It is intended that the claims submitted herewith define the overall scope.

1. A surgical system configured to improve surgical outcomes for a procedure performed on a patient in a first operating room of a medical network, the surgical system comprising:

- a first surgical hub positioned in the first operating room;
- a second surgical hub positioned in a second operating room of the medical network; and



- an analytics subsystem communicably coupled to the first and second surgical hub, wherein the analytics subsystem comprises a control circuit configured to:
- receive a consult request from the first surgical hub;
  - transmit the consult request to the second surgical hub;
  - receive an acceptance of the consult request from the second surgical hub;
  - establish an active communications link between the first surgical hub and the second surgical hub based, at least in part, on the acceptance;
  - receive a signal comprising consult information from the second surgical hub;
  - correlate the consult information to a surgical outcome of the procedure; and
  - generate a control program update for the first surgical hub based, at least in part, on the correlation of the consult information to the surgical outcome, wherein the control program update is configured to improve a future outcome of a future procedure.
2. The system of claim 1, further comprising a modular device communicably coupled to the first surgical hub, wherein the modular device is configured to generate perioperative data associated with the procedure, wherein the first surgical hub is configured to generate and transmit a second signal comprising the perioperative data, and wherein the control circuit of the analytics subsystem is further configured to:
- receive the second signal from the first surgical hub; and
  - determine the surgical outcome of the procedure based, at least in part, on the perioperative data.
3. The system of claim 2, further comprising a display communicably coupled to the second surgical hub, and wherein the active communications link comprises a livestream of the procedure presented on the display.
4. The system of claim 3, wherein the consult information comprises a recommended setting of the modular device, a recommended position of the modular device, or a recommended step of the procedure, or combinations thereof.
5. The system of claim 4, wherein the consult information comprises a preoperative recommendation, and wherein the control circuit of the data analytics subsystem is further configured to present the consult information on the display.
6. The system of claim 5, further comprising an imaging device configured to generate image data associated with the surgical procedure, and wherein the control circuit of the data analytics subsystem is further configured to present the image data on the display in conjunction with the consult information.
7. The system of claim 6, wherein the second surgical hub is positioned within a sterile field of the second operating room.
8. The system of claim 7, wherein the display comprises a personal interface configured to receive a user input from an consulting clinician using the second surgical hub, such that the consulting operator can provide the consult information without leaving the sterile field of the second operating room.
9. The system of claim 8, wherein the personal interface comprises a navigation control configured to adjust a view of the livestream of the procedure.
10. The system of claim 9, wherein the navigation control is touch-based, such that the consulting operator can use a finger to change the view of the livestream of the procedure.
11. The system of claim 2, wherein the modular device comprises an electro-surgical instrument, wherein the consult information comprises a recommended energy level for the electro-surgical instrument based, at least in part, on the perioperative data, and wherein the control program update comprises a default energy level for the future procedure, wherein the default energy level for the future procedure is the same as the recommended energy level.
12. The system of claim 2, further comprising an aggregated medical database communicably coupled to the data analytics subsystem, and wherein the control circuit of the data analytics subsystem is further configured to:
- correlate the consult information to historical outcome data stored in the aggregated medical database, and
  - wherein the generation of the control program update for the first surgical hub is further based on the correlation of the consult information to the historical outcome data.
13. An analytics system configured to improve surgical outcomes for a procedure performed on a patient in a first operating room of a medical network, wherein the analytics system is configured to be communicably coupled to a first surgical hub positioned in the first operating room and a second surgical hub positioned in a second operating room, wherein the analytics system comprises a control circuit configured to:
- receive a consult request from the first surgical hub;
  - transmit the consult request to the second surgical hub;
  - receive an acceptance of the consult request from the second surgical hub;
  - establish an active communications link between the first surgical hub and the second surgical hub based, at least in part, on the acceptance;
  - receive a signal comprising consult information from the second surgical hub;
  - correlate the consult information to a surgical outcome of the procedure; and
  - generate a control program update for the first surgical hub based, at least in part, on the correlation of the consult information to the surgical outcome, wherein the control program update is intended to improve a future outcome of a future procedure.
14. The analytics system of claim 13, wherein the first hub is communicably coupled to a modular device configured to generate perioperative data associated with the procedure, and wherein the control circuit is further configured to:
- receive the perioperative data from the first surgical hub; and
  - determine the surgical outcome of the procedure based, at least in part, on the perioperative data.
15. The analytics system of claim 14, wherein the second surgical hub comprises a display, and wherein the active communications link comprises a livestream of the procedure presented on the display.
16. The analytics system of claim 15, wherein the consult information comprises a recommended setting of the modular device, a recommended position of the modular device, or a recommended step of the procedure, or combinations thereof.
17. The system of claim 16, wherein the consult information comprises a preoperative recommendation, and wherein the control circuit is further configured to present the consult information on the display.

**18.** The system of claim **17**, wherein the control circuit is further configured to present image data associated with the surgical procedure on the display in conjunction with the consult information.

**19.** A method of improving surgical outcomes for a procedure performed on a patient in a first operating room of a medical network, the method comprising:

receiving a consult request from the first surgical hub in use during the procedure;

transmitting the consult request to the second surgical hub positioned in a second operating room of the medical network;

receiving an acceptance of the consult request from the second surgical hub;

establishing an active communications link between the first surgical hub and the second surgical hub based, at least in part, on the acceptance;

receiving a signal comprising consult information from the second surgical hub;

correlating the consult information to a surgical outcome of the procedure; and

generating a control program update for the first surgical hub based, at least in part, on the correlation of the consult information to the surgical outcome, wherein the control program update is intended to improve a future outcome of a future procedure.

**20.** The method of claim **19**, further comprising:

receiving a second signal from the first surgical hub, wherein the second signal comprises perioperative data generated by a modular device communicably coupled to the first surgical hub; and

determining the surgical outcome of the procedure based, at least in part, on the perioperative data.

**21.** The method of claim **20**, further comprising:

retrieving historical outcome data from an aggregated medical database; and

correlating the consult information to historical outcome data retrieved from the aggregated medical database, and wherein the generation of the control program update for the first surgical hub is further based on the correlation of the consult information to the historical outcome data.

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