

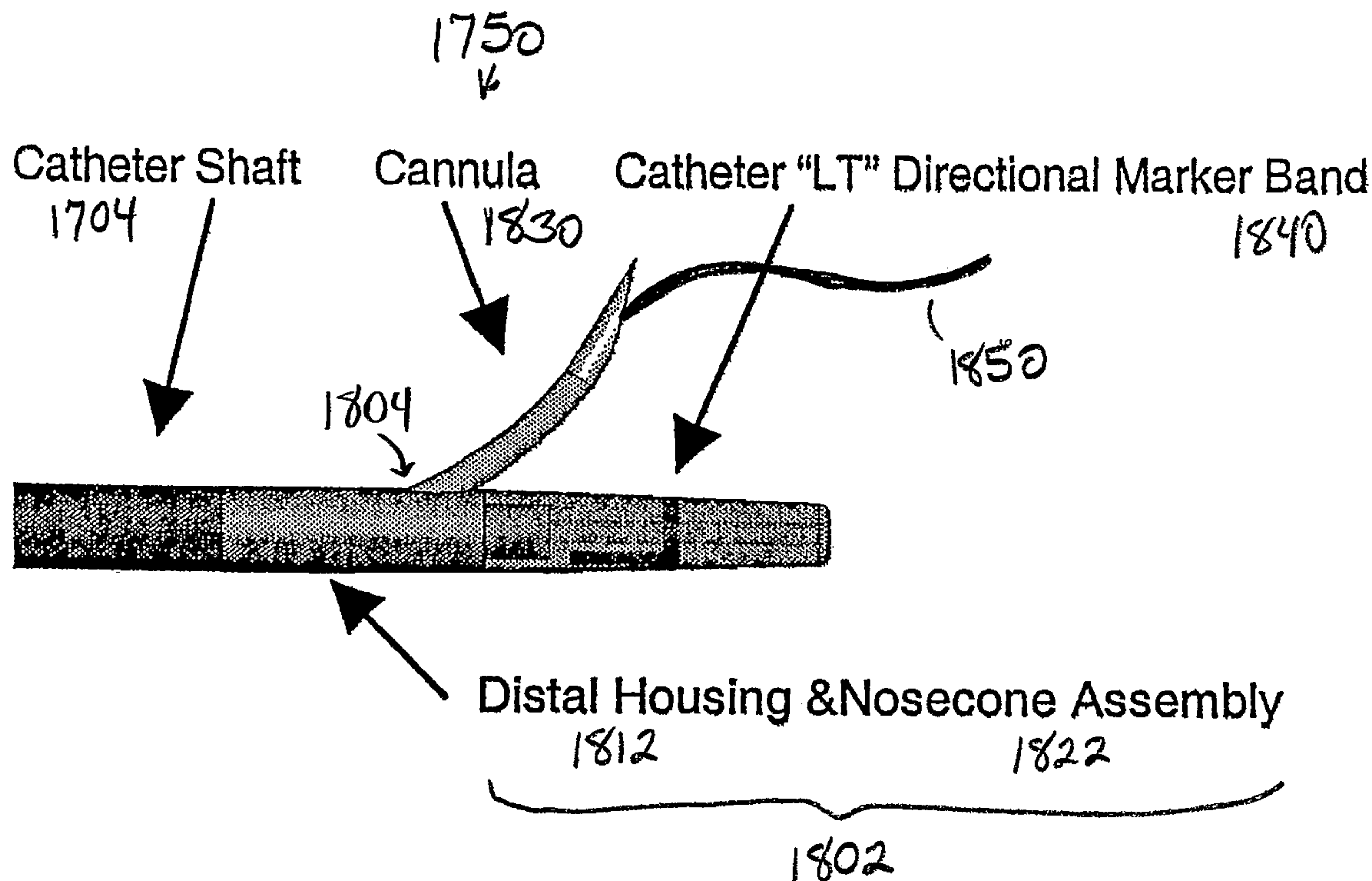


(86) Date de dépôt PCT/PCT Filing Date: 2006/03/30
 (87) Date publication PCT/PCT Publication Date: 2006/10/05
 (85) Entrée phase nationale/National Entry: 2007/09/27
 (86) N° demande PCT/PCT Application No.: US 2006/011547
 (87) N° publication PCT/PCT Publication No.: 2006/105244
 (30) Priorité/Priority: 2005/03/30 (US60/666,896)

(51) Cl.Int./Int.Cl. *B22D 19/00* (2006.01)
 (71) Demandeur/Applicant:
LUMEND, INC., US
 (72) Inventeurs/Inventors:
SELMON, MATTHEW R., US;
SPARKS, KURT D., US;
BETELIA, RAY, US;
CLARK, BENJAMIN J., US;
KAISER, JASON, US;
DECKMAN, ROBERT K., US;
THAI, ERIK, US
 (74) Agent: SIM & MCBURNEY

(54) Titre : SYSTEMES DE CATHETER DESTINES A TRAVERSER LES OCCLUSIONS TOTALES DANS UNE VASCULATURE

(54) Title: CATHETER SYSTEMS FOR CROSSING TOTAL OCCLUSIONS IN VASCULATURE



(57) **Abrégé/Abstract:**

Medical devices and methods are described that include catheter systems for use in vasculature. The catheter systems include a re-entry catheter for use with numerous guide wires to direct the guide wire from the extraluminal or subintimal space back into a true lumen after the guide wire has entered the subintimal space. An example of the re-entry catheter is a single lumen catheter configured to facilitate placement and positioning of guide wires and catheters within vasculature. An embodiment places and positions guide wires and catheters within peripheral vasculature. More specifically, the re-entry catheter provides for re-entry of a guide wire back into the true lumen of peripheral vasculature from a subintimal space.

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau(43) International Publication Date
5 October 2006 (05.10.2006)

PCT

(10) International Publication Number
WO 2006/105244 A2

(51) International Patent Classification:

B22D 19/00 (2006.01)

(21) International Application Number:

PCT/US2006/011547

(22) International Filing Date: 30 March 2006 (30.03.2006)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:

60/666,896 30 March 2005 (30.03.2005) US

(71) Applicant (for all designated States except US): LUMEND, INC. [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US).

(72) Inventors; and

(75) Inventors/Applicants (for US only): SELMON, Matthew, R. [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US). SPARKS, Kurt, D. [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US). BETELIA, Ray [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US). CLARK, Benjamin, J. [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US). KAISER, Jason

[US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US). DECKMAN, Robert, K. [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US). THAI, Erik [US/US]; 400 Chesapeake Drive, Redwood City, California 94063 (US).

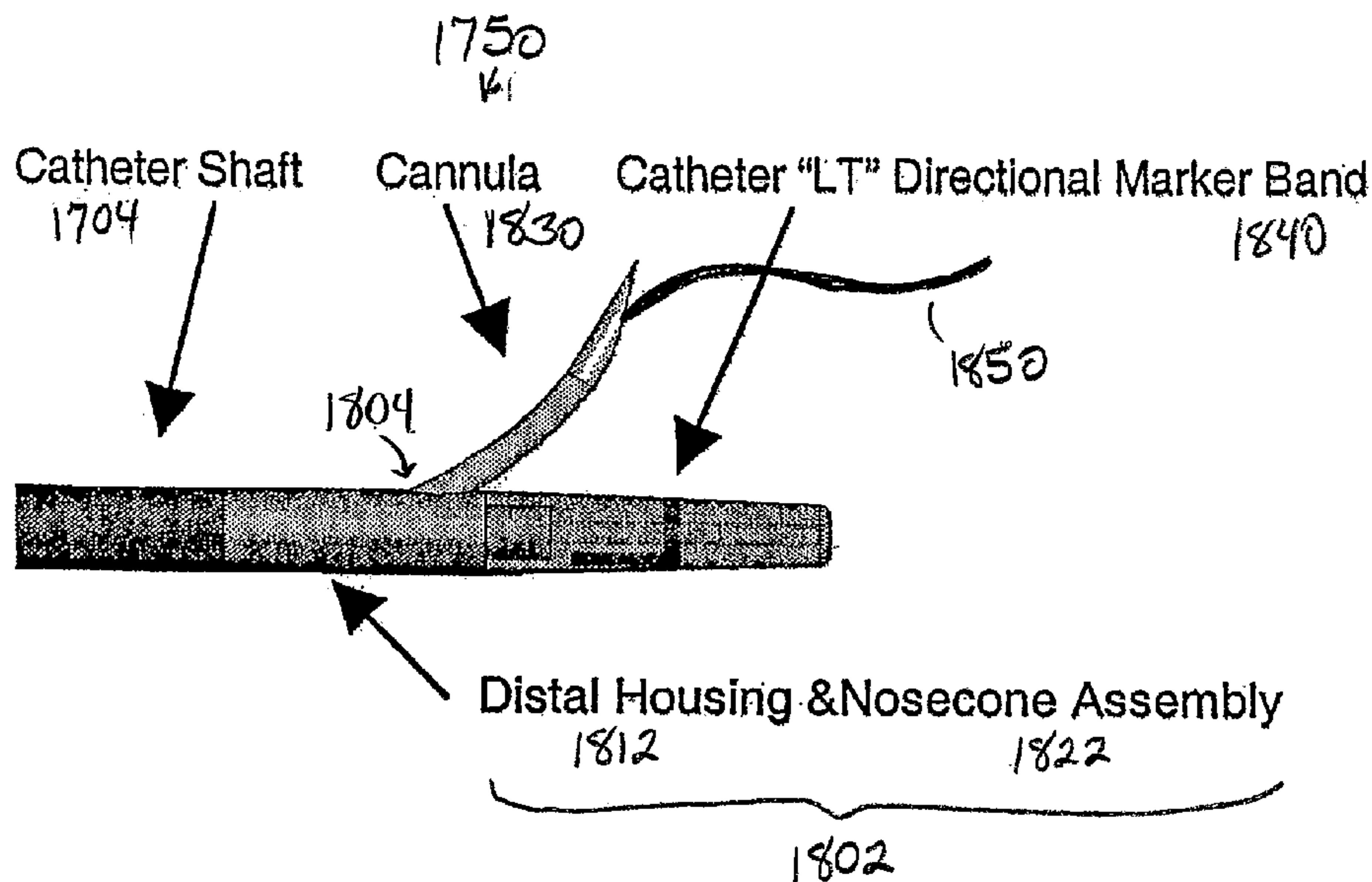
(74) Agent: GREGORY, Richard, L., Jr.; Courtney, Barbara B., and practitioners, Courtney Staniford & Gregory & LLP, P.O. Box 9686, San Jose, C 95157 (US).

(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI,

[Continued on next page]

(54) Title: CATHETER SYSTEMS FOR CROSSING TOTAL OCCLUSIONS IN VASCULATURE



(57) Abstract: Medical devices and methods are described that include catheter systems for use in vasculature. The catheter systems include a re-entry catheter for use with numerous guide wires to direct the guide wire from the extraluminal or subintimal space back into a true lumen after the guide wire has entered the subintimal space. An example of the re-entry catheter is a single lumen catheter configured to facilitate placement and positioning of guide wires and catheters within vasculature. An embodiment places and positions guide wires and catheters within peripheral vasculature. More specifically, the re-entry catheter provides for re-entry of a guide wire back into the true lumen of peripheral vasculature from a subintimal space.

WO 2006/105244 A2

WO 2006/105244 A2



FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT,
RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA,
GN, GQ, GW, ML, MR, NE, SN, TD, TG).

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

Published:

— *without international search report and to be republished upon receipt of that report*

Catheter Systems For Crossing Total Occlusions In Vasculature

Inventors:

Matthew R. Selmon

Kurt Sparks

Ray Betelia

Ben Clark

Jason Kaiser

Rob Deckman

Erik Thai

RELATED APPLICATIONS

This application claims the benefit of United States Patent Application Number 60/666,896, filed March 30, 2005. This application is a continuation-in-part application of United States Patent Application Number 10/823,416, filed April 12, 2004, which is a continuation-in-part application of United States Patent Application Number 10/308,568, filed December 3, 2002, which is a continuation of United States Patent Application Number 09/765,777, filed January 19, 2001, now United States Patent Number 6,511,458, which is a continuation of United States Patent Application Number 09/440,308, filed November 17, 1999, now United States Patent Number 6,235,000, and which is a division of United States Patent Application Number 09/006,563, filed January 13, 1998, now United States Patent Number 6,231,546.

TECHNICAL FIELD

The disclosure herein relates generally to medical devices and methods. In particular, this disclosure relates to systems, methods and procedures for crossing chronic total occlusions in vasculature.

BACKGROUND

Cardiovascular disease is a leading cause of mortality worldwide. Cardiovascular disease can take many forms, and a variety of specific interventional and pharmaceutical treatments have been devised over the years with varying levels of success.

A particularly troublesome form of cardiovascular disease results when a blood vessel becomes totally occluded with atheroma or plaque, referred to as a

chronic total occlusion (CTO). Until recently chronic total occlusions have typically been treated by performing a bypass procedure where an autologous or synthetic blood vessel is anastomotically attached to locations on the blood vessel upstream and downstream of the occlusion. While highly effective, such bypass procedures
5 are quite traumatic to the patient.

Recently, catheter-based intravascular procedures have been utilized to treat chronic total occlusions with increasing success. Catheter-based intravascular procedures include angioplasty, atherectomy, stenting, and the like, and are often preferred because they are much less traumatic to the patient. Before such catheter-
10 based treatments can be performed, however, it is usually necessary to cross the occlusion with a guide wire to provide access for the interventional catheter.

In some instances, crossing the occlusion with a guide wire can be accomplished simply by pushing the guide wire through the occlusion. After being advanced through the occlusion, the guide wire emerges in the blood vessel lumen
15 and provides the desired access path. In many cases, however, the guide wire inadvertently penetrates into the subintimal space between the intimal layer and the adventitial layer of the blood vessel as it attempts to cross the occlusion. Once in the subintimal space, it is very difficult and in many cases impossible for a physician/user to direct the guide wire back into the blood vessel lumen. In such
20 cases, it will usually be impossible to perform the catheter-based intervention and other, more traumatic, procedures may have to be employed. Catheters for use in treating chronic total occlusions are described in United States Patent Numbers 4,405,314, 4,947,864, 5,183,470, 5,190,528, 5,287,861, 5,409,019, 5,413,581, 5,429,144, 5,443,497, and 5,464,395 as well as in International Publication Numbers
25 WO 97/13463 and WO 97/13471. For these reasons, there is a need to provide devices and methods that facilitate the crossing of chronic total occlusions with guide wires.

INCORPORATION BY REFERENCE

30 Each patent, patent application, and/or publication mentioned in this specification is herein incorporated by reference in its entirety to the same extent as if each individual patent, patent application, and/or publication was specifically and individually indicated to be incorporated by reference.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows a section of artery and the tissue layers that form the artery.

Figure 2A is a section of a diseased artery that shows detail of the normal tissue of the arterial wall along with a total occlusion.

5 **Figure 2B** is a section of a diseased artery that shows an arterial wall with a modified tissue structure that can result from the presence of the total occlusion.

Figure 3A, 3B, 3C, 3D, and 3E shows a procedure for crossing a total occlusion using a catheter system, under an embodiment.

Figure 4 is a distal region of a deflecting catheter, under an embodiment.

10 **Figure 5** is a distal region of a deflecting catheter, under an alternative embodiment.

Figure 6 shows a distal region of a deflecting catheter, under another alternative embodiment.

15 **Figure 6A** shows a distal region of a deflecting catheter that assumes a straight configuration in the retracted state, under an alternative embodiment.

Figure 6B shows a distal region of a deflecting catheter that assumes a curved configuration in the retracted state, under an alternative embodiment.

Figure 7 is a deflecting catheter system, under an embodiment.

20 **Figure 8** is a cross-sectional view of a distal region of the deflecting catheter system that includes a cannula in a retracted configuration, under an embodiment.

Figure 9 is a cross-sectional view of a distal region of the deflecting catheter system that includes a cannula in an advanced configuration, under an embodiment.

Figure 10 is a cross-sectional view of a proximal region of the deflecting catheter system that includes a proximal hub, under an embodiment.

25 **Figure 10A and Figure 10B** are cross-sectional views of a proximal actuation handle with a locking mechanism that prevents inadvertent deployment of the cannula, under an embodiment.

Figure 11A and Figure 11B show rotational keying in cross-sectional views of a proximal region of the deflecting catheter system, under an embodiment.

30 **Figure 12** shows rotational keying in cross-sectional views of a distal region of the deflecting catheter system, under an embodiment.

Figure 13 is a catheter system that includes a catheter shaft having a distal end port and a proximal phased array ultrasound device, under an embodiment.

Figure 14A shows an ultrasound visualization system deployed beyond a distal end of the catheter shaft to image surrounding tissue, under an embodiment.

Figure 14B shows an ultrasound visualization system deployed to image surrounding tissue from a window within the nosecone of the catheter system, under an embodiment.

Figure 15 shows an ultrasound visualization system deployed within a cannula of the catheter shaft to image surrounding tissue, under an embodiment.

Figure 16 shows a composite distal end termination of a braided catheter shaft that includes a nosecone, under an embodiment.

Figure 17 shows a re-entry catheter in an extended configuration, under an embodiment.

Figure 18 shows the distal portion of the re-entry catheter in the extended configuration, under an embodiment.

Figure 19 shows the re-entry catheter in a retracted configuration, under an embodiment.

Figure 20A is a perspective view of the distal housing of the catheter nosecone, under an embodiment.

Figures 20B and 20C are side cross-sectional views of the distal housing, under an embodiment.

Figure 21A is a first side view of the LT directional marker band, under an embodiment.

Figure 21B is a second side view of the LT directional marker band, under an embodiment.

Figure 21C is an end view of the LT directional marker band, under an embodiment.

Figure 22 is a flow diagram for crossing occlusions using the re-entry catheter, under an embodiment.

Figure 23 shows how the LT directional marker band is used to locate and tune the position of the lateral port of the re-entry catheter, under an embodiment.

DETAILED DESCRIPTION

Devices and methods are provided below that include catheters, guides, and/or other apparatus for use in crossing total occlusions in vasculature. The total occlusions are also referred to as total vascular occlusions or chronic total occlusions

(CTOs). The devices and methods include medical devices for use by physicians/users with conventional and/or specialized guide wires to direct or redirect the guide wire from the subintimal space back into the blood vessel lumen after the guide wire has entered the subintimal space. These devices and methods are useful in the treatment of coronary artery disease in coronary arteries as well as other blood vessels and should be capable of being performed with or without imaging. The devices/methods are also useful for applications in other arteries and veins, such as the treatment of peripheral vascular diseases.

Figure 1 shows a section of artery A and the tissue layers that form the artery A. A normal (non-diseased) artery A comprises an arterial wall having a number of layers. The innermost layer is referred to herein as the intimal layer I which includes the endothelium, the subendothelial layer, and the internal elastic lamina IEL. A medial layer M is concentrically outward from the internal elastic lamina IEL, an external elastic lamina layer EEL is concentrically outward from the medial layer M, and an adventitial layer AL is the outermost layer. Beyond the adventitial layer AL is extravascular tissue. As used hereinafter, the region between the intimal layer I and the adventitial layer AL, generally including the medial layer M, is referred to as the subintimal space, but the subintimal space can include additional tissue types/layers as described below. This definition of subintimal space used herein is in addition to any meaning(s) provided by those skilled in the art.

Figure 2A is a section of a diseased artery A that shows detail of the normal tissue of the arterial wall along with a total occlusion TO. **Figure 2B** is a section of a diseased artery A that shows an arterial wall with a modified tissue structure that can result from the presence of the total occlusion TO. With reference to **Figure 2B**, a diseased artery A with a total occlusion TO comprises an arterial wall having a modified structure as compared to the tissue layers of a normal artery. The innermost layer is referred to as diffuse disease DD. The diffuse disease DD layer may range in thickness from approximately 50 microns to 500 microns, but is not limited to this thickness in heavily diseased arteries. A medial layer M may be located concentrically outward from the total occlusion TO and diffuse disease DD. In heavily diseased vessels the medial layer M may have eroded and may not be evident, as shown in **Figure 2B**. An external elastic lamina EEL may be located concentrically outward from either the medial layer M or total occlusion TO or

diffuse disease DD, and an adventitial layer AL is the outermost layer. As used hereinafter, the region within the diffuse disease DD and bounded by the adventitial layer AL is also referred to as the subintimal space, where this additional definition of subintimal space is in addition to any meaning(s) provided by those skilled in the art. The subintimal space is the region through which the wires, deflecting catheters, and other catheters described herein pass when crossing a total occlusion.

A total occlusion TO may comprise atheroma, plaque, thrombus, and/or other occluding materials normally associated with cardiovascular disease. By "total" occlusion, it is meant that the occluding material occludes substantially the entire lumen L of the artery or other blood vessel so that blood flow through the vessel is substantially blocked or stopped. The catheter systems and methods described herein are generally used with patients in whom the totally occluded artery is not immediately life threatening since the tissue distal to the occlusion will often receive oxygenated blood from collateral arteries. Usually, however, the blood supply in regions distal to the occlusion will be insufficient and it will be desirable to treat the occlusion by an intravascular intervention, such as angioplasty, atherectomy, stenting, or the like, to restore blood flow through the affected vessel.

Total occlusions are crossed by positioning a guide wire, or blunt dissection catheter (as described in United States Patent numbers 5,968,064, 6,217,549, 6,398,798, 6,508,825, 6,599,304, and 6,638,247, for example) at the proximal end of the occlusion and advancing the device through the occlusion using conventional interventional methods. Crossing of the total occlusions is defined herein, in addition to any meanings provided by those skilled in the art, as establishing a longitudinal path from the proximal end of the occlusion to the distal end of the occlusion. The longitudinal path of the guide wire and/or blunt dissection catheter is to remain as central as possible within the occluded vessel and emerge in the true lumen of the vessel after having traversed the occlusion. In practice however, the guide wire or blunt dissection catheter often tracks an eccentric pathway through the occlusion and, after having been advanced beyond the distal end of the occlusion itself, is contained within a layer of vascular tissue that is distal to the terminal end of the occlusion. A dissection track of this type is generally referred to as a subintimal track, i.e. between the intimal layer and adventitial layer of the vessel, and is typically contained within the medial layer but is not so limited (**Figures 1, 2A, and 2B**).

However, more often the disease state that forms a total occlusion erodes the inner layers of the vessel wall at the site of the occlusion, as described above with reference to **Figure 2B**. Further, the erosion of the vessel wall and the disease state often does not abruptly end at either of the proximal or distal ends of the occlusion; instead, the disease state also lines the vessel wall tapering up to and away from the occlusion. Accordingly, the structure of the vessel section that is proximal and distal to the terminal ends of the occlusion is often diffusely diseased. In these diseased vessel sections proximal and distal to the total occlusion, the intimal layer (I), internal elastic lamina layer (IEL) and the medial layer (M) may not be present, and are often replaced with a layer of diffuse disease (DD) that comprises at least one of atheroma, plaque, thrombus, fatty and fibo-calcific deposits/tissue. All of this diseased tissue is typically contained within the external elastic lamina (EEL) and the outer-most boundary of the vessel, the adventitial layer (AL), but is not so limited.

Therefore, taking into consideration that the vessel segment in the region of the occlusion may be either of normal structure (**Figure 2A**), or of diseased structure (**Figure 2B**), the subintimal track which is distal to the occlusion may be described in two corresponding ways. Within a normal vessel, the subintimal track is described as being bounded by the adventitial layer (AL) and the intimal layer (I), i.e. within the vessel wall, and usually thought to be within the medial layer (M). Within a diseased vessel, the subintimal track is described as being within a layer of diffuse disease (DD) that is also externally bounded by the external elastic lamina (EEL) or adventitial layer (AL).

Note, also, that there is a difference between a dissection tract that is contained within the total occlusion, and a dissection tract that is propagated beyond the total occlusion. When the dissection tract is propagated beyond the total occlusion, both types of subintmal tracks described above will be defined as being extra-luminal, i.e. outside the bounds of the vessel true lumen that is distal to the occlusion. The term "extra-luminal" only has relevance when the dissection track has been advanced beyond the distal end of the total occlusion. Note that a dissection track that is contained within a total occlusion can have no reference to an extra-luminal location, since no physical lumen exists within the occlusion. Hence for further reference, the subintimal tract distal to the occlusion is defined as extra-luminal, in addition to any meaning(s) provided by those skilled in the art.

Also notable is the difference between intra-vascular and extra-vascular locations. Since all dissection tracts described herein are contained within the boundary of the blood vessel, e.g. within the adventitial layer (AL), these are considered "intra-vascular". Accordingly, all locations outside of the adventitial layer (AL) are termed "extra-vascular". Extra-vascular locations are not material to the discussions described herein.

Having now established the different structures found in diseased vessels, from the subintimal, extra-luminal locations previously described, a passage or pathway is formed from these subintimal locations to the true lumen of the vessel via methods described herein. In the methods of an embodiment, a guide wire is deflected using a deflecting catheter. Typically, the deflecting catheter is advanced over a proximal end of the guide wire and advanced into the track within the subintimal space. The guide wire and the deflecting catheter are then manipulated so that the guide wire is deflected laterally through the intimal layer or diffuse disease back into the blood vessel lumen at a point distal to the occlusion. Such deflecting catheters also support the guide wire as it is advanced into and/or through the track, i.e. the catheter can enhance the pushability of the guide wire when it is advanced forward through any resisting material.

Alternatively, the guide wire which is initially positioned within the track in the subintimal space may be withdrawn through the deflecting catheter and exchanged for a second wire or other device suitable for penetrating through the intimal layer or diffuse disease back into the blood vessel lumen. The guide wires and/or deflecting catheters and other catheters can be freely exchanged over or through one another in a conventional matter without departing from the methods described herein.

In an embodiment, the physician/user determines when the guide wire and/or deflecting catheter is positioned distal to the total occlusion so that the guide wire can be returned to the blood vessel lumen beyond or distal to any occlusions. Most simply, such position determination can be made by fluoroscopically imaging the blood vessel in a conventional matter.

Alternatively or additionally to such fluoroscopic imaging, intravascular imaging, e.g. intravascular ultrasonic imaging (IVUS), and a variety of optical imaging modalities, such as optical coherence tomography (OCT), are used. For example, an ultrasonic imaging guide wire may be used to initially access the

subintimal space and/or may be exchanged for the guide wire which is used to access the subintimal space. Alternatively, the imaging guide wire may be advanced within a lumen of the catheter, or within a cannula to a distal location of the catheter suitable for viewing the surrounding vascular tissue. An imaging guide wire present
5 in the subintimal space may readily detect the presence or absence of occluding material within the blood vessel lumen. When the transition from occluding material to normal arterial tissue is detected, it is known that the position of the guide wire has advanced beyond that of a distal region of the total occlusion.

Alternatively, an imaging system or select imaging components like those
10 described in United States Patent Numbers 5,000,185 and 4,951,677 can be carried on and/or within the catheter system and/or advanced within a lumen of the deflecting catheter to a distal position within the catheter, wherein the surrounding tissue is imaged to determine if the catheter has been advanced beyond a distal region of the total occlusion. The catheter system of an alternative embodiment
15 may house and translate the imaging system or components within the lumen of the cannula itself so that both the cannula and imaging system are independently advanced distally and retracted proximally.

After a passage is formed back from the sub-intimal track into the blood vessel lumen and a wire is in place across the total occlusion, the wire is available
20 for use as a guide wire in positioning interventional and diagnostic catheters across the total occlusion. Most commonly, interventional catheters are positioned across the total occlusion for treating the occlusion. Interventional catheters include, for example, angioplasty balloon catheters, rotational atherectomy catheters, directional atherectomy catheters, and stent-placement catheters, but are not so limited.

Wire deflecting in the catheter system of an embodiment comprises
25 deflecting a cannula from the subintimal space back into the blood vessel lumen and thereafter passing the wire through a path defined/formed by the cannula, typically via a lumen within the cannula. The cannula is advanced over the wire after the wire is disposed within the subintimal space. Deflecting of the cannula in an embodiment
30 comprises advancing a resilient (pre-formed) curved end of the cannula from a constraining lumen of the catheter into the blood vessel lumen, as described below.

Wire deflecting in alternative embodiments of the catheter system comprises advancing a deflecting catheter over a wire that was previously advanced into the subintimal space. The cannula is subsequently advanced through a lateral opening

of the deflecting catheter and penetrated through the intimal layer or diffuse disease to define a path for the guide wire back into the blood vessel lumen.

Wire deflecting in other alternative embodiments comprises advancing a deflecting catheter over a wire which was previously advanced into the subintimal
5 space. The cannula is subsequently advanced through a distal opening of the deflecting catheter and penetrated through the intimal layer or diffuse disease to define a path for the guide wire back into the blood vessel lumen. Steerable and other actively deployed cannulas may also be used.

Figure 3A, 3B, 3C, 3D, and 3E shows a crossing of a total occlusion using a
10 catheter system, under an embodiment. The catheter system includes a deflecting catheter 20 and at least one wire 10, or guide wire 10, but is not so limited. With reference to **Figure 2A** and **Figure 2B**, this procedure is performed in an upper portion of the artery, but is not so limited. Referring to **Figure 3A**, a wire 10 is advanced through the lumen L of the artery A until it encounters material of a total
15 occlusion TO, as described above. At that time, it is possible that the wire 10 will advance through the occlusion TO without deflecting into the blood vessel wall. Should that occur, subsequent repositioning of the guide wire according to the methods of the present invention may not be necessary.

More usually, however, the wire 10 will advance into the subintimal space
20 within either the medial layer M, or the diffuse disease DD, as shown in **Figure 2B** (as described above, the intimal and medial layers of advanced atherosclerotic occlusions may evolve into a heterogeneous layer of diffuse disease DD). The intimal layer I and adventitial layer AL together define a tissue plane through which the wire 10 can naturally pass as the wire 10 is pushed distally from its proximal
25 end. Alternatively, the wire 10 may be advanced through the diffuse disease DD and take a similar pathway through the occlusion TO. The wire 10 will continue to advance until the distal tip of the wire 10 passes beyond the distal end of the total occlusion TO, as shown in **Figure 3B**. The distal tip of wire 10 can axially advance well beyond the total occlusion until advancement is ceased by the physician/user.

Figure 3B shows the guide wire 10 advancing without support. In some
30 instances, however, the guide wire 10 may encounter significant resistance as it enters and/or passes through the space between the intimal layer I and the adventitial layer AL, or the diffuse disease DD. If resistance is encountered, the deflection catheter 20 may be used to support and enhance the pushability of the guide wire 10

by advancing the deflection catheter 20 to a location just proximal of the distal tip of the guide wire 10, as shown in **Figure 3C**. The guide wire 10 and catheter 20 may then be advanced sequentially, e.g. advancing the guide wire 10 a short distance followed by advancing the catheter 20 to a location just proximal of the distal tip of the guide wire 10, and so on.

Regardless of the procedure used, however, once the guide wire 10 is advanced to a point that positions the distal tip beyond the total occlusion TO, deflecting catheter 20 is advanced over the wire 10, by coaxial introduction over the proximal end of the wire 10, until it approaches the total occlusion TO, as shown in **Figure 3B**. The deflecting catheter 20 is then further advanced over the wire 10 until its distal tip also extends beyond the total occlusion TO, as shown in **Figure 3D**. The deflecting catheter 20 of an embodiment includes at least one mechanism for laterally deflecting the guide wire 10 so that the guide wire 10 can pass in a radially inward direction through the intimal layer I or the diffuse disease DD back into the blood vessel lumen L.

The deflection mechanism of an embodiment takes a variety of forms as described below. For example, referring to **Figure 3D**, the deflection mechanism of an embodiment includes a lateral port 22 in the deflecting catheter 20. The guide wire 10 can be retracted so that its distal tip lies proximal to the lateral port 22 and then advanced distally so that the wire 10 passes laterally outwardly through the lateral port 22 and back into the blood vessel lumen L, as shown in **Figure 3E**.

The physician/user of the catheter system of an embodiment can assure that the distal tip of the guide wire 10 and the deflecting port 22 (or other deflecting mechanism) of the deflecting catheter 20 are properly positioned beyond the total occlusion TO without being advanced excessively beyond the end of the total occlusion TO. The proper positioning of the deflecting catheter 20 can vary approximately in the range of 0 cm to 2 cm beyond the distal end of the total occlusion TO, but is not so limited. In one embodiment, for example, the deflecting catheter 20 is positioned approximately 0 to 0.5 cm beyond the end of the total occlusion TO.

As described above, such positioning can in some instances be performed using fluoroscopic imaging. For example, in some instances it may be sufficient to provide suitable radiopaque markers on components of the catheter system that include at least one of the guide wire, the cannula, the deflecting mechanism of the

catheter, and some combination of any of these components permitting visual positioning of a distal region of the component via fluoroscopy. Fluorescence imaging is described, for example, in United States Patent Numbers 4,718,417 and 5,106,387.

5 In addition to fluoroscopy, active imaging systems/methods and other imaging modalities that include, but are not limited to, optical coherence tomography (OCT) and Raman spectroscopy can also be used to provide imaging information in a catheter system. The OCT is described, for example, in United States Patent Numbers 5,321,501, 5,459,570, 5,383,467, and 5,439,000. Raman
10 spectroscopy is described, for example, in International Publication Number WO 92/18008.

The catheter system of an embodiment provides for rotational positioning of the deflecting catheter 20. The rotational positioning allows the direction of deflection of the cannula or guide wire to be selective by allowing the physician/user
15 to aim the deflecting mechanism from the subintimal space back toward the arterial or other blood vessel lumen L.

If the catheter is provided with ultrasonic imaging, such imaging can be used for rotationally positioning the distal tip of the catheter. The catheter of an embodiment is rotationally rigid so that rotation of the proximal end allows for
20 positioning of the distal end. Using the detected presence of the blood vessel lumen, the deflecting port 22, and/or other deflecting mechanisms, the guide wire and/or cannula can be rotationally positioned towards the vessel true lumen via ultrasonically identifiable features of these components.

In an alternative embodiment, a rotationally specific fluoroscopic marker can
25 be provided on the catheter 20, or directly on the cannula. The marker is configured so that the rotational direction of the catheter tip or cannula can be determined by observing the two-dimensional image of the marker using fluoroscopic imaging.

Devices described herein for use in crossing vascular occlusions include catheter systems, also referred to as wire deflection or wire deflecting systems. The
30 wire deflection systems of an embodiment generally comprise a wire deflecting catheter that includes a catheter body and a deflecting cannula. The catheter body includes a proximal end, a distal end, and at least one lumen extending through at least a distal portion of the catheter body. In one embodiment, the lumen communicates with at least one of a distal port and/or a lateral port in at least one of

a distal section, distal region, or end zone of the catheter. In various alternative embodiments, the lumen communicates with one or more distal ports and/or one or more lateral ports.

5 The cannula of an embodiment also includes a proximal end, a distal end, and at least one lumen extending through a distal portion of the cannula. The distal portion of the cannula may include a pre-formed resilient curve, as described below. The cannula is slidably disposed within the lumen of the catheter body but is not so limited. Upon full proximal retraction of the cannula within the catheter body lumen, the distal section of the catheter can be configured to assume at least one of a
10 straight configuration and a curved configuration. The cannula can be deployed through at least one of a lateral port and an end port as appropriate to a procedure to establish a pathway through the subintimal tissue/diffuse disease according to the methods described herein.

The choice of materials and fabrication methods for the catheter shaft and/or
15 the cannula determine the resultant shape of the distal section of the catheter when the cannula is fully retracted. Either the straight or curved configuration of the distal catheter shaft may apply to two embodiments of the catheter as follows. A first embodiment of the catheter includes both a distal and lateral port, and full advancement of the cannula selectively deploys the pre-shaped cannula from the
20 lateral port. A second embodiment of the catheter includes a single distal port, and full advancement of the cannula deploys the pre-shaped cannula from the distal port. Various alternative embodiments include different combinations of lateral ports, distal ports, and cannula deployment options.

The catheter system of an embodiment further comprises a wire configured
25 to pass through the cannula lumen. The wire may be a conventional guide wire, a wire having a sharpened distal tip extended particularly for penetrating the intimal layer of the blood vessel wall and/or diffuse disease of the blood vessel, and/or other wires known in the art. Alternatively, the wire may include passive and/or active visualization or imaging means.

30 Regarding passive visualization or imaging systems, the catheter body of an embodiment includes one or more fluoroscopically visible markers near the distal end. The markers are configured to permit visual determination of the rotational orientation of the distal end of the catheter body when viewed in a two-dimensional fluoroscopic image. The catheter body can be reinforced to enhance torsional

rigidity, and can further comprise a distal nose cone wherein the distal and/or lateral openings may be defined within the nose cone. The distal end of the cannula of an embodiment is pre-formed in a smooth curve which may extend over an arc approximately in the range of 15 to 135 degrees, but is not limited to this range. The pre-formed curve may have a radius approximately in the range of 0.5 millimeters (mm) to 15 mm, but is not limited to this range.

A number of specific embodiments of the catheter system are described below, wherein use of the catheter system is generally described above with reference to **Figures 3A-3E**. These specific embodiments are provided as examples only and do not limit the catheter systems provided herein.

Figure 4 is a distal region of a deflecting catheter 30, under an embodiment. The deflecting catheter 30 includes a distal end having at least one distal port 32, at least one lateral port 34, and a passive deflecting mechanism 36. The catheter 30 can be advanced over the proximal end of a wire like a guide wire so that the wire passes over the deflecting mechanism 36 and back into the main lumen of the catheter 30. The catheter 30 can then be advanced over the wire until the distal tip enters the subintimal space and approaches the distal end of the wire. By retracting the distal end of the wire within the lumen of catheter 30 so that its distal tip is proximal to the deflecting mechanism 36, subsequent distal advancement of the wire engages the proximal surface of the deflecting mechanism and causes the wire to be deflected laterally through the lateral port 34. The deflecting catheter 30 of an embodiment is dimensioned as follows, but is not so limited: catheter shaft inner diameter approximately in the range 0.012 inches (to accommodate a 0.010 inch guide wire) to 0.043 inches (to accommodate a 0.039 inch guide wire); and catheter shaft outer diameter approximately in the range 0.020 inches to 0.050 inches.

Figure 5 is a distal region of a deflecting catheter 40, under an alternative embodiment. Deflecting catheter 40 includes at least one distal port 42 and at least one lateral port 44. Instead of a passive deflecting mechanism, deflecting catheter 40 includes an active deflecting mechanism in the form of an axially translatable cannula 46. The cannula 46 is configured to include a resilient pre-formed distal tip that can be advanced through port 44 (shown in broken line). The cannula 46 has a lumen which provides a guide path for the wire. The deflecting catheter 40 of an embodiment is dimensioned as follows, but is not so limited: catheter shaft inner diameter approximately in the range 0.025 inches to 0.055 inches; catheter shaft

outer diameter approximately in the range 0.035 inches to 0.065 inches; cannula inner diameter approximately in the range 0.012 inches (to accommodate a 0.010 inch guide wire) to 0.043 inches (to accommodate a 0.039 inch guide wire); and cannula outer diameter approximately in the range 0.020 inches to 0.050 inches.

5 The cannula 46 of an embodiment is constructed from a composite of braided stainless steel wire that is laminated with polymers such as nylons, urethanes, polyimides or polycarbonates, but is not so limited and can be formed from other materials as appropriate to the medical applications. The distal end of the cannula 46 can be terminated in an angled cut of the material, forming a needle-like
10 tip, or terminated with a sharpened, fluoroscopic hollow metallic tip formed to include Platinum-Iridium, for example. Alternatively, the cannula 46 can be fabricated from a uniform material such as nitinol (nickel-titanium), and terminated in a needle type tip via sharpening into an appropriate shape.

To further increase torque control of the cannula 46, especially in high
15 tortuosity applications, stainless steel wire or other suitable filaments are braided onto the cannula 46 of an embodiment. The braided cannula 46 can also be laminated with appropriate polymers (nylons, polyurethanes) to produce a smooth exterior surface of the cannula shaft. Further, the polymer of an embodiment is coated with a hydrophilic agent to decrease frictional effects as the cannula 46 is
20 translated within the catheter shaft. A resilient curve may be set into the distal section of the cannula 46 via heat setting methods for these materials, such heat setting methods known in the art. Materials and methods of cannula construction also allow some degree of radiopacity, i.e. ability to produce an image under
fluoroscopy.

25 Upon full extension of the cannula 46 from the catheter shaft, the cannula 46 is unconstrained, allowing the as-manufactured curved shape of the distal cannula 46. Upon full retraction of the cannula 46 into the catheter shaft, two configurations are possible. A first configuration is one in which the distal end of the catheter assumes a curved shape. The degree of the curve can range from the as-
30 manufactured curve of the cannula to something approaching a straight configuration. A second configuration is one in which the catheter assumes a straight configuration.

Each of these configurations is possible through the selection of materials used to construct both the distal section of the catheter shaft, and the distal section of

the cannula. To achieve the first or curved configuration, the material used to construct the cannula is more robust (for example, nitinol), and the material used to construct the distal section of the catheter shaft is comparatively less robust (for example, a braided shaft laminated with low durometer urethane). This combination
5 allows the catheter shaft to conform more to the shape of the cannula as the cannula is retracted into the catheter shaft. These fabrications and fabrication materials are provided as examples only, as many others allow the distal catheter shaft to follow the shape of the cannula.

The second or straight configuration is achieved by fabricating a less robust
10 or “softer” cannula with wire braid laminated with a medium durometer nylon such as 55D Pebax, and fabricating the catheter shaft with wire braid laminated with polyamide. The softer design of the cannula shaft allows the cannula to conform to the straight configuration of the catheter shaft as the cannula is retracted into the catheter shaft. Alternatively, the distal end of the catheter shaft may be fabricated
15 with an integral section of stainless steel hypotube, or other non-flexible material. This section of hypotube performs in similar fashion to maintain the straight configuration of the distal catheter shaft as the cannula is retracted. Again, these fabrications and fabrication materials are provided as examples only, as many others allow the distal region of the cannula, upon retraction, to conform to the straight
20 configuration of the distal region of the catheter shaft.

Figure 6 shows a distal region of a deflecting catheter 50, under another alternative embodiment. The deflecting catheter 50 includes a lumen and at least one distal port 54. A cannula 52 having a pre-formed distal end may be advanced and retracted through the lumen and out of the distal port 54, but the embodiment is
25 not so limited. As described above with reference to the catheter system in **Figure 5**, the cannula 52 and catheter shaft of the deflecting catheter 50 can be fabricated of various materials that allow the cannula to assume an as-manufactured curved shape (broken line) when extended from the catheter. The deflecting catheter 50 of an embodiment is dimensioned as follows, but is not so limited: catheter shaft inner
30 diameter approximately in the range 0.025 inches to 0.055 inches; catheter shaft outer diameter approximately in the range 0.035 inches to 0.065 inches; cannula inner diameter approximately in the range 0.012 inches (to accommodate a 0.010 inch guide wire) to 0.043 inches (to accommodate a 0.039 inch guide wire); and cannula outer diameter approximately in the range 0.020 inches to 0.050 inches.

Figure 6A shows a distal region of a deflecting catheter 50A that assumes a straight configuration in the retracted state, under an alternative embodiment. The materials of the cannula 52A and catheter shaft of the deflecting catheter 50A are fabricated of materials that allow the distal end of the catheter system 50A, when the
5 cannula 52A is fully retracted into the catheter shaft, to assume a straight configuration.

Figure 6B shows a distal region of a deflecting catheter 50B that assumes a curved configuration in the retracted state, under an alternative embodiment. The materials of the cannula 52B and catheter shaft of the deflecting catheter 50B are
10 fabricated of materials that allow the distal end of the catheter system 50B, when the cannula 52B is fully retracted into the catheter shaft, to assume a curved configuration. Further, the shape of the distal end of the catheter system (either straight 50A or curved 50B, with cannula retracted) determines the type of rotational
15 keying (if any) between the cannula and catheter shaft, and the type of fluoroscopic marking used to assist a physician/user in directing the cannula deployment towards the vessel true lumen.

The three catheter systems 30, 40, and 50 presented with reference to **Figures 4, 5, and 6** are presented as examples only and do not limit the catheter system to these exact embodiments. A wide variety of other passive and active
20 deflecting mechanisms can be provided on deflecting catheters for use in the methods described herein.

The distal catheter terminations of the catheter system embodiments described herein can be fabricated as a continuation of the catheter shaft polymers, which have been formed or molded into the configurations shown. Alternatively,
25 the distal terminations include a separate nosecone component attached to the terminal end of the catheter shaft. Regarding the attachment of the nosecone, the catheter shaft can be attached to the nosecone by lamination of the shaft polymer onto features at the proximal end of the nosecone, but is not so limited.

Alternative embodiments of the catheter systems described herein, however,
30 include composite terminations that provide strong flexible connections of the nosecone to the catheter shaft. **Figure 16** shows a composite distal end termination 1600 of a braided catheter shaft 1602 that includes a nosecone 1604, under an embodiment. This composite termination 1600 includes an internal metallic ring 1610 (also referred to as the internal or inner ring 1610) and an external metallic ring

1612 (also referred to as the external or outer ring 1612) between which a braid wire 1614 of the laminated shaft 1602 distally terminates. The internal 1610 and external 1612 rings are each approximately in the range of 1 to 2 mm in length, and have a nominal thickness of approximately 0.002", but are not so limited. The internal
5 1610 and external 1612 rings are attached to the braid wire 1614 via at least one of soldering, gluing, and resistance welding, as examples, but other suitable attachment methods can be used. Following attachment of the internal 1610 and external 1612 rings the tubular braid wire 1614 is then laminated with appropriate polymers 1616 and 1618, producing a completed catheter shaft component 1602. This composite
10 termination 1600 of the catheter shaft produces a short, integral metallic "ring". A nosecone 1604 can be machined with mating features as appropriate and attached to the ring via at least one of welding, gluing and soldering.

The composite termination 1600 provides a very strong, flexible connection of the nosecone 1604 to the catheter shaft 1602. One challenge of terminating the
15 outer polymer layer 1618 to the internal 1610 and external 1612 rings is that if both the shaft polymer 1618 and ring termination are bluntly terminated to each other, the distinct boundary existing between the two materials may tend to delaminate during operation of the catheter system due to bending stresses. The termination of an embodiment alleviates this issue through the inclusion of an internal taper 1620 in
20 the external ring 1612 at the terminal end of the catheter shaft. This taper 1620 of the external ring 1612 allows a small continuous taper of polymer 1618 to be produced underneath the external ring 1612 to afford stress relief upon bending of the catheter shaft 1602 at this boundary location. This taper 1620 prevents delamination of the polymer 1618 from the internal 1610 and external 1612 rings.

25 **Figure 7** is a deflecting catheter system 100, under an embodiment. **Figure 8** is a cross-sectional view of a distal region 104 of the deflecting catheter system 100 that includes a cannula 114 in a retracted configuration, under an embodiment. **Figure 9** is a cross-sectional view of a distal region 104 of the deflecting catheter system 100 that includes a cannula 114 in an advanced configuration, under an
30 embodiment. **Figure 10** is a cross-sectional view of a proximal region 106 of the deflecting catheter system 100 that includes a proximal hub 112, under an embodiment.

Referring to **Figures 7, 8, 9, and 10**, the deflecting catheter 100 comprises a catheter body 102 having a distal end 104 and a proximal end 106. Catheter body

102 includes a single lumen 108, and a deflection housing 110 secured to the distal end 104 of the catheter body 102. An actuator hub 112 is secured to the proximal end 106 of the catheter body 102, and an axially translatable cannula 114 is disposed within lumen 108. A distal length 118 of the cannula 114 is pre-formed in a curved
5 shaped, but is not so limited. The cannula 114 has a sharpened tip 116, formed using at least one of metal, hard plastic, composite, and/or combinations of these materials. The cannula tip 116 of an embodiment is radiopaque, but is not so limited. Alternatively or additionally, at least one separate radiopaque marker, similar to marker 120, is included in the catheter system on the cannula at or near its
10 distal end to facilitate visualization under fluoroscopic imaging. A rotationally specific radiopaque marker 120 is mounted near the distal end of catheter body 102. The marker has a generally U-shaped configuration so that the rotational position of the distal end of the catheter body 102 is apparent when observing the marker using a two-dimensional fluoroscopic image, but the marker is not so limited.

15 The deflecting catheter 100, in operation, laterally deflects the distal tip of the cannula 114 through a lateral opening 122 in the deflector housing 110. The deflector housing 110 also includes a distal port 124 to permit introduction of the catheter 100 over the proximal end of a guide wire GW, as shown in **Figure 8** in broken line. The guide wire GW passes through the distal port 124 and into the
20 distal end of the cannula 114 and passes through a lumen of cannula 114 to the proximal end of the catheter 100. In one embodiment, the distal length 118 of cannula 114 is straightened and deflected by axially retracting and advancing the cannula 114 between the configurations shown in **Figure 8** and **Figure 9**, respectively. Consistent with the description above that references **Figure 5**, an
25 alternative embodiment of the catheter system includes a catheter shaft that maintains some degree of curve as the cannula is retracted.

30 With reference to **Figure 10**, the actuator hub 112 comprises a pair of coaxial, telescoping tubes 130 and 132. The outer telescoping tube 132 is connected to a proximal end of cannula 114 using, for example, an adhesive 134. A proximal fitting 136 is further attached to the proximal end of tube 132 so that the assembly of the cannula 114, tube 132, and fitting 136 move together as a unit through the hemostatic fitting 140 at the proximal end of the hub 112. The hub 112 further includes a rotational fitting 142 which permits the catheter body 102 to be rotated relative to the hub body. The cannula 114 and catheter body 102 are rotationally

coupled or keyed together to limit and/or prevent relative rotation. The cannula 114 and catheter body 102 of an embodiment are coupled using keying within the hub and/or near the distal end so that rotation of the catheter body 102 causes a like rotation of the cannula 114 as the catheter is rotationally positioned within a blood vessel. A side port 148 is provided on the hub 112 to permit perfusion and/or infusion through the lumen 108 or catheter 102.

Figure 10A and Figure 10B are cross-sectional views of a proximal actuation handle 1000 with a locking mechanism that prevents inadvertent deployment of a cannula, under an embodiment. The actuation handle 1000 of an embodiment includes a slide mechanism 1002 with an integral lock 1004 that travels in a linear slot of a handle body 1006. The slide 1002 attaches to the working element 1010 which, in this embodiment, is a cannula 1010. Proximal and distal movement of the slide mechanism advances 1030 and retracts 1040 the working element 1010. Upon full retraction of the slide 1002, the spring-loaded 1008 lock mechanism 1004 is activated, automatically tripping the distal end of the lock 1004 into the distal end of the slot in the handle body 1006 through which the slide translates. Subsequent distal advancement of the slide 1002 and working element 1010 is only possible upon depressing the proximal end of the lock 1004, which disengages the distal end of the lock 1004 from the slot in the handle body 1006. Use of this locking mechanism 1004 prevents the working element 1010 from inadvertent deployment while the catheter system is being tracked within the vasculature. Components like the hemostatic fittings and keying can also be included, as described above with reference to **Figure 10**.

Keying of components of the catheter system, as described above, can be accomplished with a variety of techniques at both the proximal and distal end of the catheter. Keying at the proximal end of the catheter 100 can be achieved in a variety of ways. **Figure 11A and Figure 11B** show rotational keying in cross-sectional views of a proximal region 104 of the deflecting catheter system 100, under an embodiment. As an example, telescoping tubes 130 and 132 of the catheter 1102 of an embodiment include asymmetric, mating peripheral geometries having oval cross-sections. Likewise, telescoping tubes 130 and 132 of the catheter 1112 of an alternative embodiment include asymmetric, mating peripheral geometries having triangular cross-sections. These geometries are shown as examples only, and do not limit the catheter systems described herein to these geometries.

Keying at the distal end of the catheter 100 can also be achieved in a number of ways. **Figure 12** shows rotational keying in cross-sectional views of a distal region of the deflecting catheter system 100, under an embodiment. For example, the catheter body 102 can include an asymmetric lumen 108. The cannula 114 uses a mating cross-section, e.g. a D-shaped cross-section in this example. The ability to limit relative rotation of the cannula 114 within the catheter body 102 assures that the curved distal length 118 of the cannula 114 is properly oriented (directed radially outwardly) when the cannula tip 116 emerges through the lateral opening 122.

In use, and with reference to **Figure 8 and Figure 9**, catheter 100 is advanced over guide wire GW while the cannula 114 is retracted. Once the catheter is properly positioned, the guide wire is retracted a few centimeters proximal to the cannula tip 116, and the cannula 114 may be distally advanced. Distal advancement is achieved by forwardly advancing the sleeve 132/hub 136 relative to the body of the hub 112 so that the cannula advances within the lumen 108 of catheter body 102. Prior to advancing the cannula, the lateral port 122 is properly positioned so that it is directed toward the blood vessel lumen. Positioning of the lateral port 122 includes, in one embodiment, rotation of the catheter body 102 using the rotational hub 142. The physician/user observes the marker 120 so that the lateral port 122 is directed in the proper direction, for example radially inward. Following advancement of the cannula into the blood vessel, the guide wire GW can be advanced into the lumen. The cannula 114 is subsequently withdrawn proximally, and the entire catheter assembly is then withdrawn from over the guide wire, leaving the guide wire in place for introduction of other interventional and/or diagnostic catheters.

When the distal end of the catheter system assumes a straight configuration upon retraction of the cannula into the catheter shaft, keying and fluoroscopic marker options are numerous. As described with reference to **Figure 7** above, the distal end of the catheter shaft can specify the direction of cannula deployment via fluoroscopic indicator(s) 120. To support this capability in the catheter system, the cannula and catheter shaft of an embodiment are keyed to each other, as described herein. With the cannula retracted, its "curve" will be straightened, and the fluoroscopic image of the cannula will not indicate the direction of deployment. Therefore, the straightened "curve" rotationally follows (is keyed to) the catheter shaft marker used to indicate the direction the cannula will take when deployed.

This rotational keying (alignment of cannula curve to catheter shaft marker 120) is set during the manufacturing of the catheter.

5 In an alternative embodiment, the features of the fluoroscopic marker 120 are directly incorporated into the catheter nosecone or distal termination of the catheter shaft. Keying of the catheter shaft and cannula is also incorporated as described above.

10 In another alternative embodiment, the cannula can include a similar marking system 120 as the shaft. Since the cannula marker directly indicates deployment direction of the cannula, keying is not used between catheter shaft and cannula, nor is directional marking on the catheter shaft, but the embodiment is not so limited.

15 Yet another alternative embodiment includes a combination of catheter shaft marker and cannula marker. In this embodiment, since the cannula includes directional (deployment) marking, the catheter marker is not directional, and it signifies the location of the catheter distal end. A simple fluoroscopic nosecone or ring suffices for this type of marking.

20 When the distal end of the catheter system assumes a curved configuration upon retraction of the cannula into the catheter shaft, there are also numerous keying and fluoroscopic marker options. In an embodiment, non-directional fluoroscopic markers as described herein are used on either the distal catheter shaft or the distal cannula. Keying is not included since the curve of the cannula in the retracted position automatically angles the distal portion of the catheter shaft in the direction of the cannula deployment.

25 In an alternative embodiment a directional marker 120 is included on/in the cannula to indicate deployment direction, and works in concert with the directional curve at the distal end of the catheter shaft. Keying is not included in this embodiment (Figure 6b).

Another alternative embodiment includes a directional marker on the catheter for use in concert with the directional curve at the distal end of the catheter shaft. Keying is included in this embodiment, as described above.

30 Yet another alternative embodiment includes one or more combinations of the marking schemes described herein. In general, whenever directional marking is used on the catheter shaft, keying is used to align the catheter shaft marker to the cannula curve deployment direction.

In addition to the numerous passive visualization systems described above, various embodiments of the catheter systems described herein can include active on-board visualization systems and methods. One example of an on-board visualization system is described as a rotational ultrasound system in United States Patent
5 Numbers 4,951,677 and 5,000,185, but the on-board visualization systems used in the catheter systems described herein are not so limited.

An embodiment of the catheter system, however, provides ultrasonic or other imaging at or near the total occlusion to assist the physician/user in positioning the catheter system. In one embodiment, guide wire includes at least one ultrasonic
10 imaging component or device to detect the presence and absence of the occluding material as the wire is advanced past the total occlusion. In an alternative embodiment, the deflecting catheter includes such ultrasonic imaging, e.g. in the form of a phased array located near the distal tip of the deflecting catheter. United States Patent Numbers 4,917,097 and 5,368,037 describe a phased array system for
15 use in a deflecting catheter, but the embodiment is not limited to these visualization systems.

Figure 13 is a catheter system 1300 that includes a catheter shaft 1302 having a distal end port 1304 and a proximal phased array ultrasound device 1310, under an embodiment. The phased array ultrasound device 1310 is included within
20 and/or on the nosecone 1306 of the catheter system 1300 at the distal end of the catheter shaft 1302, but can be included within/on other components of the catheter system 1300. As the catheter shaft 1302 is rotated, the phased array ultrasound device 1310 produces an image of the surrounding tissue, and the image is used to identify the vessel true lumen. Once the vessel true lumen is correctly identified, a
25 cannula (not shown) is advanced through the lumen 1308 of the catheter shaft 1302; the cannula is keyed to exit in the direction of the vessel true lumen, as identified by the ultrasound image. Alternatively, a rotational imaging catheter system or its functional components are advanced either within the lumen of the catheter system or within the cannula of the catheter system, as appropriate, to a distal site of the
30 catheter from which the surrounding tissue can be imaged.

The visualization system of an embodiment includes ultrasonic imaging guide wires, but is not so limited. For example, United States Patent Number 5,095,911 describes an ultrasonic imaging guide wire, but the embodiment is not limited to this particular imaging guide wire. As yet another alternative, an imaging

guide wire can be advanced to the region of the total occlusion in a direction opposite to that of the guide wire and catheter. In this way the imaging guide wire need not advance through the total occlusion, but could still detect advancement of the catheter and/or guide wire, particularly if ultrasonically opaque components are provided on the catheter and/or the guide wire. In still another alternative, an ultrasonic imaging catheter or guide wire is positioned in a vein adjacent to the arterial occlusion site, allowing imaging of the entire occluded region while the guide wire is advanced through the occluded region.

Generally, an ultrasound visualization system can be used under at least two methods. Under a first method, the rotational ultrasound catheter system, or its functional components are advanced first within a catheter lumen to a distal region of the catheter shaft suitable for imaging the surrounding tissue. As an example, the ultrasound components can be advanced beyond a distal end of the catheter. **Figure 14A** shows an ultrasound visualization system 1402 deployed beyond a distal end 1404 of the catheter shaft 1406 to image surrounding tissue, under an embodiment.

As another example, the ultrasound components can be to a distal region of the catheter shaft while remaining housed in the catheter. **Figure 14B** shows an ultrasound visualization system 1402 deployed to image surrounding tissue from a window 1410 within the nosecone 1412 of the catheter system, under an embodiment. The window can be formed from a polymer like polyethylene that has acoustically transparent properties at the operating frequency of the ultrasound system, but is not so limited.

Following advancement of the ultrasound system to a location appropriate for imaging, the catheter is aligned to the vessel true lumen under a number of methods. Under a first method of catheter alignment, features at the distal end of the catheter, such as those that are machined into a nosecone as the viewing window, are identified via ultrasound imaging, and used to align a lateral or distal port to the true lumen. The ultrasound system is subsequently extracted and a keyed cannula system is advanced within the catheter lumen to exit in the direction of the viewing window and true lumen.

Under a second method of catheter alignment, ultrasound systems/components similar to those described in the first method of catheter alignment are used to align a catheter port with the vessel true lumen. However, instead of relying on a keying mechanism to align the cannula with the vessel true

lumen (as described above), use is made of the ultrasonic components of the catheter as fluoroscopic alignment features. In this way, once the catheter port is aligned to the vessel true lumen using the ultrasonic components, the same features may provide fluoroscopic alignment, indicating direction of the vessel true lumen. A
5 cannula with a directional fluoroscopic marker is then advanced within the catheter and brought into fluoroscopic alignment with the catheter marker, providing deployment guidance towards the vessel true lumen.

A second method under which ultrasound visualization systems are used includes advancing the rotational ultrasound catheter system or its functional
10 components within the cannula. **Figure 15** shows an ultrasound visualization system 1502 deployed within a cannula 1504 of the catheter shaft 1506 to image surrounding tissue, under an embodiment. Under this method, the ultrasound imaging is used to first identify the vessel true lumen via either images taken with the ultrasound system extended out from the distal port or ultrasound images taken
15 through a lateral window at the distal end of the catheter shaft. Further, the ultrasound imaging system may be retracted slightly to identify features on the cannula shaft or tip which indicate cannula deployment direction. The cannula can then be rotated to bring the identified cannula features into alignment with the vessel true lumen, as appropriate. The cannula is then deployed to gain access to the vessel
20 true lumen.

The deflecting catheter system of an embodiment includes a re-entry catheter. One example of a re-entry catheter is the Outback® LTD Re-Entry Catheter available from LuMend® of Redwood City, California. The re-entry catheter is a single lumen catheter configured to facilitate placement and positioning
25 of guide wires and catheters within vasculature. An embodiment places and positions guide wires and catheters within peripheral vasculature, but is not limited to peripheral vasculature. More specifically, the re-entry catheter allows for re-entry of a guide wire back into the true lumen of a peripheral artery from a subintimal space.

30 The re-entry catheter of an embodiment generally includes but is not limited to a catheter shaft or body with a catheter nosecone on a distal end and a deployment handle with a control knob on a proximal end. The re-entry catheter includes and/or works in conjunction with a guide or guide system that includes a cannula and/or a guide wire. A concentric lumen extends from the handle, through the catheter shaft,

and exits through the catheter nosecone. The lumen is configured to accept and pass a cannula having a cannula tip and/or a guidewire. The term “working element” is used herein to refer to the cannula, the guide wire, or the cannula in combination with the guide wire. A concentric lumen extending through the cannula and cannula tip is configured to accept and pass the guide wire.

Proximal retraction of a deployment slide of the handle positions the cannula tip coaxially within the distal lateral port, allowing the catheter and cannula to be tracked over a guide wire (e.g. 0.014 inch guide wire). Once at a target or desired vascular site, the catheter is aligned via visualization (e.g. fluoroscopic) and positioned by rotation of the handle rotating luer and the catheter lateral exit port is aligned via the catheter directional marker band located on the nosecone. Once in position, the guide wire is retracted into the cannula, allowing the curved cannula tip to be advanced from the catheter lateral port as required to access the target vascular location. The cannula curve of an embodiment is visible (e.g. fluoroscopically) and can indicate the point of entry during the re-entry process. The guide wire is advanced to extend from the cannula tip and into the target vascular site. The cannula tip is subsequently retracted into the catheter lateral port, and the catheter is proximally retracted leaving the guide wire in place in the vasculature. Operation of the re-entry catheter is described in detail below.

As one example of a re-entry catheter, **Figure 17** shows a re-entry catheter 1700 in an extended configuration, under an embodiment. The re-entry catheter 1700 includes a deployment handle 1702 (handle), a rotating hemostasis valve (RHV), a catheter shaft 1704, a catheter nosecone 1706, and a cannula having a cannula tip 1708. One example of a deployment handle 1702 is described above with reference to Figure 10. The catheter shaft of an embodiment is approximately 120 centimeters long but is not limited to this length. The cannula (not shown) is positioned in an interior lumen of the catheter shaft and includes a cannula wire port 1710 on a proximal end for accepting a guide wire. The cannula is axially translatable. A concentric guide wire lumen (not shown) extends from the handle 1702, through the cannula and cannula tip 1708, and exits through the catheter nosecone 1706. The re-entry catheter 1700 also can include a deployment slide release button 1712, a flush port 1714, and other components as appropriate to the medical procedure in which it is used. The re-entry catheter 1700 has a working length approximately in a range of 60 centimeters to 160 centimeters, and is 8

French to 4 French sheath compatible, but is not so limited. Guide wires compatible with the re-entry catheter 1700 include 0.010 inch to 0.038 inch coaxial guide wires.

The cannula of an embodiment has an approximately 22 gauge outside diameter and is a pre-formed curved re-entry cannula formed from materials that include shape memory alloys like Nickel Titanium (Nitinol), but is not so limited. The cannula, which can be covered (e.g., plated, adhered, etc.) with a radio-dense material (e.g., gold, platinum, tantalum, and/or other materials or metals appropriate to use in the human body and under the procedures employed), can be keyed to the catheter shaft 1704 of an embodiment but is not so limited. An example of keying of the cannula to the catheter shaft 1704, when used, is described above with reference to Figure 11 and Figure 12 but is not so limited. The cannula has a sharpened tip, formed using at least one of metal, hard plastic, composite, and/or combinations of these materials. The cannula tip of an embodiment can include radiopaque material, but is not so limited. Alternatively or additionally, at least one separate radiopaque marker can be included in the catheter system on the cannula at or near its distal end to facilitate visualization under fluoroscopic imaging.

In the extended configuration, the cannula and cannula tip 1708 or guide tip is extended through a port in a distal region of the re-entry catheter 1700. **Figure 18** shows the distal portion 1750 of the re-entry catheter 1700, under an embodiment. The distal portion 1750 includes a catheter nosecone 1802 coupled to a distal end of the catheter shaft 1704. The catheter nosecone 1802 includes a distal housing 1812 and nosecone assembly 1822. The catheter nosecone also includes an LT directional marker band 1840 as described in detail below. The cannula 1830, when deployed from the re-entry catheter 1700, extends through a lateral port 1804 or lateral exit port of the catheter nosecone 1802 as the distal end of the cannula 1830 exits the catheter nosecone 1802. The cannula 1830 of an embodiment is configured to deploy a guide wire 1850.

Figure 19 shows the re-entry catheter 1700 in a retracted configuration, under an embodiment. In the retracted configuration the cannula 1830 is retracted into the catheter shaft 1704 so that the cannula tip (not shown) does not protrude through the exit port of the catheter nosecone 1706.

Figure 20A is a perspective view of the distal housing 1812 of the catheter nosecone 1802, under an embodiment. The distal housing 1812 together with the nosecone assembly (not shown) forms the catheter nosecone. The distal housing

1812 includes a proximal region 2002 that couples or connects to a distal end or region of catheter shaft (not shown). A distal region 2004 of the distal housing also includes a lateral port 1804 as described above.

Figures 20B and 20C are side cross-sectional views of the distal housing 1812, under an embodiment. All dimensions shown are in inches. The distal housing 1812 includes a lumen 2010 in communication with a lateral port 1804 and a distal port 2012. A proximal region 2010P of the distal housing lumen 2010 couples to a distal region of the catheter shaft lumen (not shown) and is configured to accept and pass the cannula and/or a guide wire. A middle region 2010M of the distal housing lumen 2010 couples the lumen proximal region 2010P to the lateral port 1804 and is configured to accept and pass the cannula and/or guidewire. The lumen middle region 2010m is also configured to include a curved region that is a passive deflecting mechanism. The passive deflecting mechanism, in operation, laterally deflects the distal tip of the cannula and/or guide wire (not shown) through the lateral port 1804 of the distal housing 1812.

The lumen middle region 2010M also couples to a distal region 2010D of the distal housing lumen 2010. The lumen distal region 2010D is relatively smaller in cross-sectional area than the lumen proximal 2010P and middle regions 2010M; the relatively smaller cross-sectional area allows the distal region 2010D to accommodate passage of the guide wire through the distal port 2010 while preventing passage of the larger cannula. The distal port 2012 permits introduction of the re-entry catheter over the proximal end of a guide wire (not shown), where the guide wire passes through the nosecone assembly, distal port of the distal housing, and through a lumen of cannula to the proximal end of the catheter.

The re-entry catheter of an embodiment includes a marker for use in locating and/or positioning the re-entry catheter in a patient's vasculature. The marker, also referred to herein as the LT directional marker band, is located in the nosecone assembly (element 1822 with reference to Figure 18) of the catheter nosecone of an embodiment, but can be located in one or more other portions of the re-entry catheter. The LT directional marker band includes a first portion 2110 that is in a circular configuration. The first portion 2110 includes an orifice 2111 for accommodating a lumen (not shown) of the catheter nosecone. The first portion 2110 is connected to a second portion 2112 that extends outward from the first portion 2110 and is oriented approximately orthogonally to a plane that includes the

first portion 2110. The LT directional marker band is positioned in the nosecone assembly distal to the lateral port so that the first portion 2110 of the marker is distal to the second portion 2112 but is not so limited.

Figure 21A is a first side view 2101 of the LT directional marker band, under an embodiment. The first side view 2101 presents the LT directional marker band as an “L” shape for use in locating the cannula tip towards a target tissue site, as described below. **Figure 21B** is a second side view 2102 of the LT directional marker band, under an embodiment. The second side view 2102 is a view of the LT directional marker band when rotated ninety (90) degrees from the position of the first side view 2101. The second side view 2102 presents the LT directional marker band as a “T” shape for use in tuning the cannula tip location relative to the target tissue site, as described below. **Figure 21C** is an end view 2103 of the LT directional marker band, under an embodiment. All dimensions shown are in inches.

The re-entry catheter, as described above, is configured to facilitate placement and positioning of guide wires and catheters within vasculature. More specifically, the re-entry catheter facilitates crossing chronic total occlusions in vasculature. **Figure 22** is a flow diagram 2200 for crossing occlusions using the re-entry catheter, under an embodiment. A physician/user or other clinician (referred to herein as a user) positions 2202 the re-entry catheter in a patient’s vasculature using a guide wire, and orientates 2204 an exit port of the catheter towards a desired vascular target site. The user retracts 2206 the guide wire tip into the re-entry catheter. The cannula tip is extended 2208 from the catheter lateral port, and positioned at the vascular target site. The user advances the guide wire through the cannula tip to position 2210 it at the vascular target site. The user retracts 2212 the catheter over the guide wire, leaving the guide wire in place for subsequent therapeutic procedures.

When preparing to use the re-entry catheter of an embodiment to place and/or position a guide wire within vasculature, a physician or other clinical user fully retracts the cannula tip via proximal retraction of the handle deployment slide until it hard stops. Prior to insertion into a patient’s body, the user ensures the cannula tip is fully retracted into the catheter lateral port and the handle deployment slide is locked in the most proximal position.

The user back-loads a guide wire (e.g. 0.014 inch guide wire) into the catheter through the distal end port of the catheter nosecone, and introduces the

catheter and guide wire into the vasculature using appropriate percutaneous techniques. If a guide wire has already been placed in the vasculature, the guide wire is back-loaded into the catheter through the distal end port of the catheter nosecone. Before back-loading the guide wire, the user ensures the cannula tip is fully retracted back into the catheter shaft. Advancement, manipulation, and withdrawal of the catheter are performed under high-quality fluoroscopic guidance, but other types of guidance can be used as appropriate to a configuration of the re-entry catheter. While tracking the catheter over a guide wire, the user should ensure the cannula tip is fully retracted inside the catheter lateral port and the handle deployment slide is locked in the most proximal position.

Upon introducing the catheter into the patient's vasculature, the user tracks the catheter over the guide wire to a desired vascular site. The catheter is torqued as needed during delivery via the handle RHV. If strong resistance is felt during catheter manipulation/delivery, the user should determine the cause of the resistance before proceeding further with the procedure. Consider, for example, using a 3-4 millimeter balloon to dilate points of resistance, as needed, along the catheter delivery track.

When near the desired vascular site, the re-entry catheter is positioned prior to deployment of a cannula and/or guide wire. As described above, the re-entry catheter distal housing includes an LT directional marker band for use in locating, tuning catheter position, and deploying working elements. The LT directional marker band is used generally to locate and tune a position of the lateral port through which the cannula and/or guide wire is deployed relative to a target re-entry site in the vasculature. **Figure 23** shows how the LT directional marker band is used to locate and tune the position of the lateral port of the re-entry catheter, under an embodiment. The "L" shape 2302 of the LT directional marker band is used to locate the lateral port, and hence the cannula tip when it is deployed, towards the target re-entry site. The "T" shape 2304 of the LT directional marker band is used to tune or fine tune the lateral port position, and hence the location/direction of the deploying cannula tip, relative to the target re-entry site. A clinician using the LT directional marker band therefore locates 2312 or identifies the facing direction of the lateral port by rotating the re-entry catheter so the LT directional marker appears as the "L" shape 2302; the direction of the lower horizontal portion of the "L" shape 2302 indicates the deployment direction of the cannula tip. The clinician fine tunes

the cannula tip deployment position by rotating 2313 the catheter in a direction that results in the LT directional marker band appearing 2314 as the “T” shape 2304. The directional capability of the marker thus allows for placement of a guide wire into a true lumen of a peripheral artery for example, without ultrasound guidance.

5 The LT direction marker band is used during a procedure as follows. When in an approximate area of the desired vascular site the user orientates the re-entry catheter (e.g. rotate) so the distal housing of the re-entry catheter is located adjacent to the target re-entry site (e.g. true lumen) when visualized using fluoroscopy. The lateral exit port of the catheter is orientated towards the desired vascular target site
10 via rotation of the RHV. Using fluoroscopic guidance, the user orientates the distal housing to position the “L” marker leg on the catheter LT directional marker band towards the target re-entry site (e.g. true lumen) using an initial fluoroscopic view.

 Once the initial location and orientation is confirmed, an orthogonal fluoroscopic view is made (e.g. view from a position rotated ninety (90) degrees
15 relative to the previous fluoroscopic view) to confirm that the distal housing of the re-entry catheter is positioned “in line” with the target re-entry site as indicated by the visible “L” marker leg. The user then orientates the lateral exit port of the catheter towards the desired vascular target site via rotation of the RHV. The user performs this tuning of the initial orientation, under fluoroscopic guidance, by
20 rotating the RHV until the catheter LT directional marker band on the nosecone appears as a “T”. If additional orientation adjustments are necessary, this can be achieved through rotation of the RHV. A confirming view (e.g. orthogonal view) should be considered after each new adjustment of the catheter towards the re-entry target as appropriate to the procedure.

25 Upon completion of the locating and tuning of the distal housing position to properly position the re-entry catheter, as described above, the user releases any stored torque in the catheter shaft. The user ensures the catheter LT directional marker band slot is oriented toward the desired vascular location (target site) prior to actuation of the handle deployment slide, and retracts the guide wire tip into the
30 catheter approximately five (5) centimeters, using fluoroscopic guidance to confirm guide wire position. The handle deployment slide release button is depressed and the slide is incrementally advanced, as appropriate, to extend or deploy the cannula tip from the catheter lateral port and position it at the vascular target site.

The guide wire is advanced through the deployed cannula tip to position it as desired at the vascular target site. If after advancing the guide wire distally, it is desired to retract the guide wire and resistance is experienced, first retract the cannula (guide) fully into the catheter, and proceed with retraction of the guide wire.

5 Following advancement and placement of the guide wire, the cannula tip is retracted into the catheter by fully retracting the handle deployment slide until it hard stops, and the handle deployment slide button is released to lock the deployment slide in the retracted position. The user should ensure the cannula tip is fully retracted into the catheter lateral port, and the handle deployment slide is locked, prior to

10 withdrawing the catheter over the guide wire. The catheter is then retracted over the guide wire, leaving the guide wire in place for one or more subsequent therapeutic procedures.

The re-entry catheter of an embodiment can include an anchoring device or system for anchoring a distal region of the re-entry catheter. The catheter anchoring

15 systems include use of a biodegradable gel/glue, a balloon, a wire mesh/net, anchoring wires, and/or an extended catheter tip. Each of these anchoring systems is described in more detail below.

The gel anchor uses a biodegradable gel or glue to fill spaces or gaps in the subintimal layer around the catheter and host artery. The catheter is tracked over the

20 guide wire into the subintimal space and positioned as appropriate to the re-entry point. The biodegradable gel is delivered into the subintimal layer through the catheter from the proximal end. The gel, once delivered, steadies the catheter during the re-entry procedure with the cannula and/or guide wire. Once the catheter has been stabilized, the re-entry procedure continues with extension of the cannula

25 and/or guide wire.

The balloon anchor uses gap-filling balloons formed from various materials to fill the expanded subintimal space around the catheter and the host artery. The catheter is tracked over the guide wire into the subintimal space and positioned as appropriate to the re-entry point. The cannula is deployed from the lateral port in

30 order to re-enter the true lumen. If re-entry is unsuccessful due to instability of the catheter distal end, then the balloon is inflated an amount appropriate to the configuration of the subintimal space. The properly inflated balloon reduces an amount of subintimal space around the distal region of the catheter and thus stabilizes the distal region of the catheter. Once the catheter has been stabilized, the

re-entry procedure continues with extension of the cannula and/or guide wire. The balloon is deflated following successful re-entry into the true lumen.

The wire mesh or net anchor uses a mesh or net to fill the loose space or larger dissection plane of the subintimal layer. The catheter is tracked over the
5 guide wire into the subintimal space and positioned as appropriate to the re-entry point. The cannula is deployed from the lateral port in order to re-enter the true lumen. If re-entry is unsuccessful due to instability of the catheter distal end, then the mesh is introduced into the subintimal layer via the proximal end of the catheter and a catheter lumen. The introduction of the mesh reduces an amount of subintimal
10 space around the distal region of the catheter and thus stabilizes the distal region of the catheter. Once the catheter has been stabilized, the re-entry procedure continues with extension of the cannula and/or guide wire. The mesh is retracted or pulled back into the catheter following successful re-entry into the true lumen.

The anchoring wire uses wires, coils, and/or prongs made of an appropriate
15 material (e.g. metal, polymer, etc.) to push against the tissue and/or muscle in the subintimal space. The pressure against the tissue stabilizes the catheter during deployment of the cannula and/or guide wire. Once the catheter has been stabilized, the re-entry procedure continues with extension of the cannula and/or guide wire. The anchoring wires are retracted or pulled back into the catheter following
20 successful re-entry into the true lumen.

The anchoring that uses the extended tip of the catheter provides an extended tip that submerges between the target lumen and tissue beyond the target re-entry site. Once the catheter has tracked over the wire to the target re-entry site and is appropriately positioned, the portion of the extended tip that extends beyond the
25 lateral port of the nosecone is trapped between the lumen and vasculature tissue. Consequently, the extended tip uses the anatomy of the vasculature to stabilize or hold the distal region of the catheter. The re-entry procedure then continues with extension of the cannula and/or guide wire.

The catheter system of an embodiment includes a catheter system for use in
30 vasculature. The catheter system of an embodiment comprises a catheter body including a body lumen. The catheter system of an embodiment includes a nosecone coupled to a distal end of the catheter body. The nosecone of an embodiment includes a nosecone lumen, a lateral port, and a distal port.

A proximal section of the nosecone lumen of an embodiment is in communication with the body lumen.

A middle section of the nosecone lumen of an embodiment is configured to include a passive deflection region in communication with the lateral port and the proximal section.

A distal section of the nosecone lumen of an embodiment is in communication with the distal port and the proximal section.

The catheter system of an embodiment includes a marker in the nosecone configured to present as a plurality of symbols that indicate relative orientation of the lateral port to a target site in the vasculature.

A cross-sectional area of the distal section of the nosecone lumen of an embodiment is relatively smaller than the cross-sectional area of the proximal section and the middle section.

The middle section of the nosecone lumen of an embodiment is configured for passage of a cannula to the lateral port.

The distal section of the nosecone lumen of an embodiment is configured for passage of a guide wire to the distal port.

The marker of an embodiment is located distal to the lateral port.

The plurality of symbols of the marker of an embodiment includes a first symbol and a second symbol.

The catheter system of an embodiment includes a working element having a distal end configured to deploy through the lateral port for delivery from a first vascular location within a subintimal space to a second vascular location within a true lumen of the vasculature when the working element is advanced distally through the lateral port.

The working element of an embodiment is keyed to the catheter body.

The working element of an embodiment includes a cannula having at least one lumen.

The working element of an embodiment includes a cannula and at least one guide wire slidably disposed within the cannula.

The distal end of the working element of an embodiment includes a preformed resilient tip.

The distal end of the working element of an embodiment assumes a first configuration when the working element is retracted into the catheter body and a second configuration when the working element is extended through the lateral port.

The catheter system of an embodiment includes a method for re-entering a true lumen of vasculature from a subintimal space. The method of an embodiment includes advancing a catheter over a wire into the subintimal space and retracting the wire. The method of an embodiment includes locating and positioning a lateral port of the catheter approximately adjacent a target re-entry site of the true lumen using information of a plurality of symbols presented by a marker in a distal region of the catheter during visualization. The method of an embodiment includes advancing the wire through the lateral port in to the true lumen, wherein the true lumen is re-entered by the wire.

The locating and positioning of an embodiment includes positioning the lateral port approximately adjacent the true lumen by rotating the catheter to a position that presents a first symbol.

The locating and positioning of an embodiment includes tuning the positioning of the lateral port by rotating the catheter to a position that presents a second symbol. A position of an imager during visualization resulting in presentation of the second symbol is approximately orthogonal to the position of the imager during visualization resulting in presentation of the first symbol.

The method of an embodiment includes advancing a cannula through the lateral port toward the target re-entry site.

The advancing of the cannula of an embodiment comprises deflecting the cannula from the catheter.

The advancing of the cannula of an embodiment comprises advancing the wire through the cannula.

The visualization of an embodiment includes fluoroscopy.

The subintimal space is located within diffuse disease of the vasculature.

The subintimal space is located between an adventitial layer and an intimal layer of the vasculature.

Unless the context clearly requires otherwise, throughout the description and the claims, the words “comprise,” “comprising,” and the like are to be construed in an inclusive sense as opposed to an exclusive or exhaustive sense; that is to say, in a sense of “including, but not limited to.” Words using the singular or plural number

also include the plural or singular number respectively. Additionally, the terms “herein,” “hereunder,” “above,” “below,” and terms of similar import, when used in this application, refer to this application as a whole and not to any particular portion of this application. When the word “or” is used in reference to a list of two or more
5 items, that word covers all of the following interpretations of the word: any of the items in the list, all of the items in the list and any combination of the items in the list.

The above description of illustrated embodiments of the catheter system is not intended to be exhaustive or to limit the catheter system to the precise form
10 disclosed. While specific embodiments of, and examples for, the catheter system are described herein for illustrative purposes, various equivalent modifications are possible within the scope of the catheter system, as those skilled in the relevant art will recognize. The teachings of the catheter system provided herein can be applied to other medical devices and systems, not only for the catheter systems described
15 above.

The elements and acts of the various embodiments described above can be combined to provide further embodiments of the catheter system. These and other changes can be made to the catheter system in view of the above detailed description. Furthermore, aspects of the catheter system can be modified, if
20 necessary, to employ the systems, functions and concepts of the various patents and applications described above to provide yet further embodiments of the system.

In general, in the following claims, the terms used should not be construed to limit the catheter system to the specific embodiments disclosed in the specification and the claims, but should be construed to include all catheter systems and medical
25 devices that operate under the claims to cross vascular occlusions. Accordingly, the catheter system is not limited by the disclosure, but instead the scope of the catheter system is to be determined entirely by the claims.

While certain aspects of the catheter system are presented below in certain claim forms, the inventors contemplate the various aspects of the catheter system in
30 any number of claim forms. Accordingly, the inventors reserve the right to add additional claims after filing the application to pursue such additional claim forms for other aspects of the catheter system.

CLAIMS

What is claimed is:

- 1 1. A catheter system for use in vasculature, comprising:
2 a catheter body including a body lumen;
3 a nosecone coupled to a distal end of the catheter body, the nosecone
4 including a nosecone lumen, a lateral port, and a distal port, wherein a proximal
5 section of the nosecone lumen is in communication with the body lumen, a middle
6 section of the nosecone lumen is configured to include a passive deflection region in
7 communication with the lateral port and the proximal section, and a distal section of
8 the nosecone lumen is in communication with the distal port and the proximal
9 section; and
10 a marker in the nosecone configured to present as a plurality of symbols that
11 indicate relative orientation of the lateral port to a target site in the vasculature.
- 1 2. The catheter system of claim 1, wherein a cross-sectional area of the distal
2 section is relatively smaller than the cross-sectional area of the proximal section and
3 the middle section.
- 1 3. The catheter system of claim 1, wherein the middle section is configured for
2 passage of a cannula to the lateral port.
- 1 4. The catheter system of claim 1, wherein the distal section is configured for
2 passage of a guide wire to the distal port.
- 1 5. The catheter system of claim 1, wherein the marker is located distal to the
2 lateral port, wherein the plurality of symbols include a first symbol and a second
3 symbol.
- 1 6. The catheter system of claim 1, further comprising a working element having
2 a distal end configured to deploy through the lateral port for delivery from a first
3 vascular location within a subintimal space to a second vascular location within a
4 true lumen of the vasculature when the working element is advanced distally
5 through the lateral port.

1 7. The catheter system of claim 6, wherein the working element is keyed to the
2 catheter body.

1 8. The catheter system of claim 6, wherein the working element includes a
2 cannula having at least one lumen.

1 9. The catheter system of claim 6, wherein the working element includes a
2 cannula and at least one guide wire slidably disposed within the cannula.

1 10. The catheter system of claim 6, wherein the distal end of the working
2 element includes a preformed resilient tip.

1 11. The catheter system of claim 6, wherein the distal end of the working
2 element assumes a first configuration when the working element is retracted into the
3 catheter body and a second configuration when the working element is extended
4 through the lateral port.

1 12. A method for re-entering a true lumen of vasculature from a subintimal
2 space, comprising:
3 advancing a catheter over a wire into the subintimal space and retracting the
4 wire;
5 locating and positioning a lateral port of the catheter approximately adjacent
6 a target re-entry site of the true lumen using information of a plurality of symbols
7 presented by a marker in a distal region of the catheter during visualization; and
8 advancing the wire through the lateral port in to the true lumen, wherein the
9 true lumen is re-entered by the wire.

1 13. The method of claim 12, wherein locating and positioning includes
2 positioning the lateral port approximately adjacent the true lumen by rotating the
3 catheter to a position that presents a first symbol.

1 14. The method of claim 13, wherein locating and positioning includes tuning
2 the positioning of the lateral port by rotating the catheter to a position that presents a
3 second symbol.

1 15. The method of claim 14, wherein a position of an imager during visualization
2 resulting in presentation of the second symbol is approximately orthogonal to the
3 position of the imager during visualization resulting in presentation of the first
4 symbol.

1 16. The method of claim 12, further comprising advancing a cannula through the
2 lateral port toward the target re-entry site.

1 17. The method of claim 16, wherein advancing comprises deflecting the
2 cannula from the catheter.

1 18. The method of claim 16, wherein advancing comprises advancing the wire
2 through the cannula.

1 19. The method of claim 12, wherein the visualization includes fluoroscopy.

1 20. The method of claim 12, wherein the subintimal space is located within
2 diffuse disease of the vasculature.

1 21. The method of claim 12, wherein the subintimal space is located between an
2 adventitial layer and an intimal layer of the vasculature.

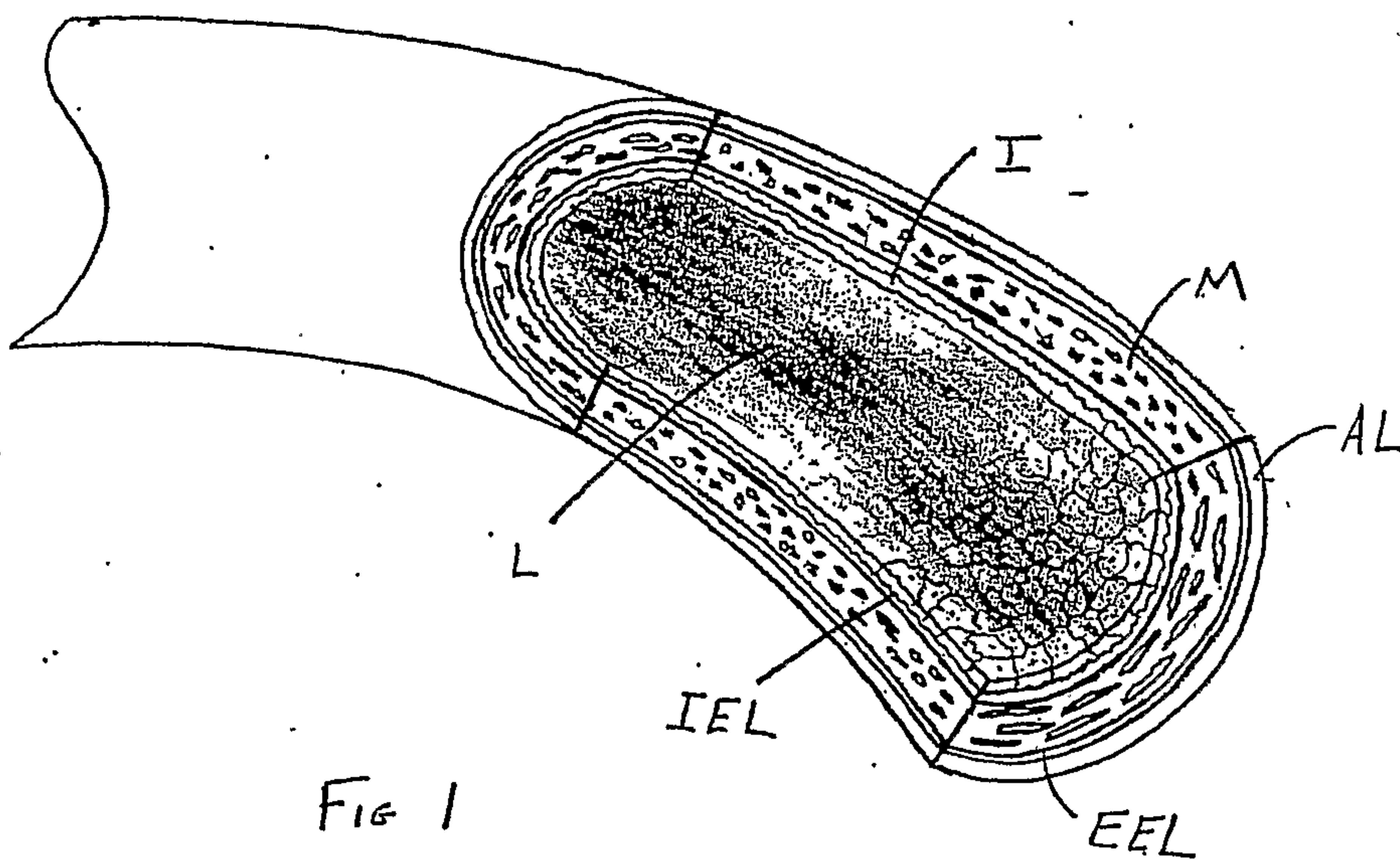


FIG 1

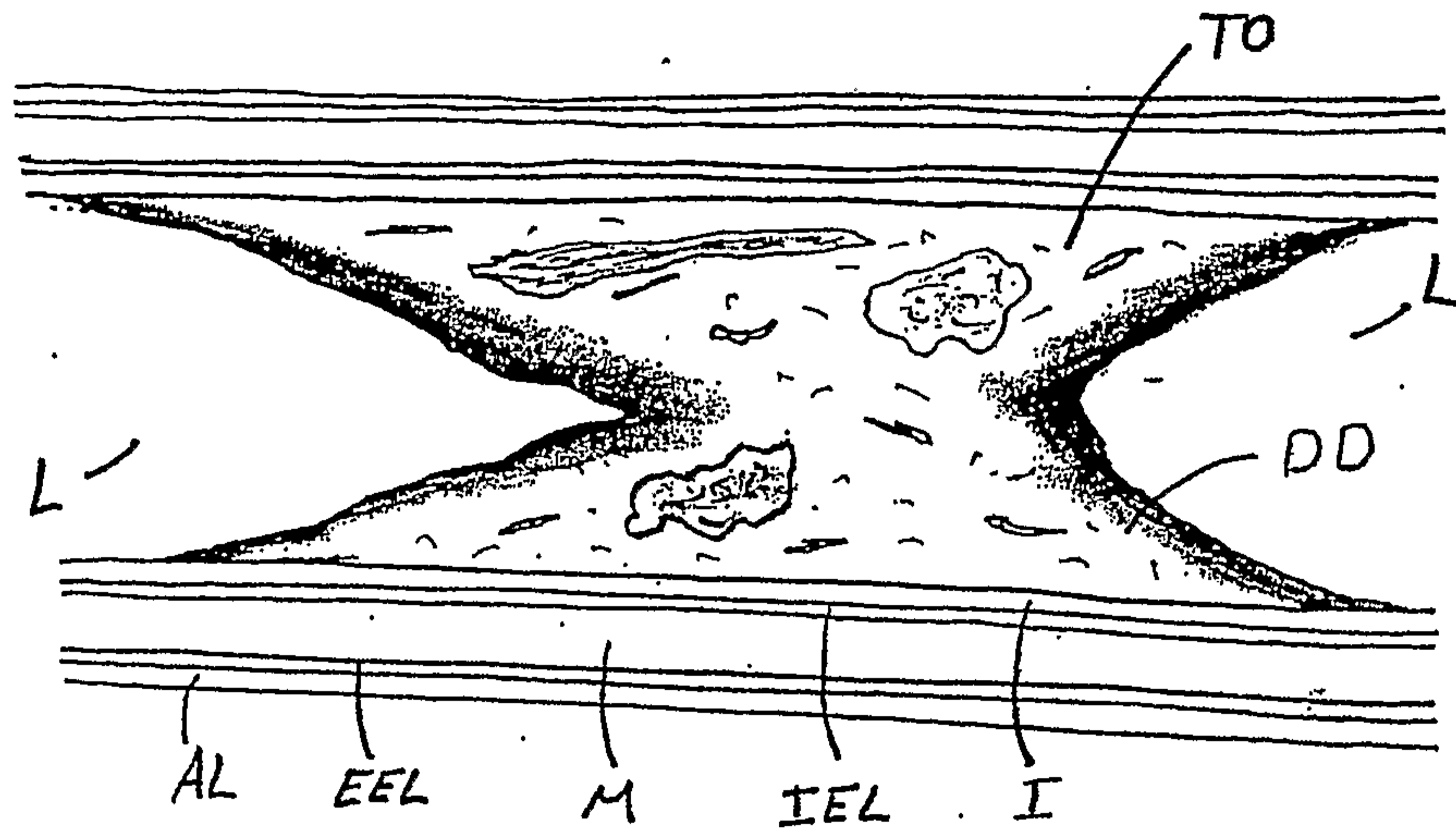


FIG 2A

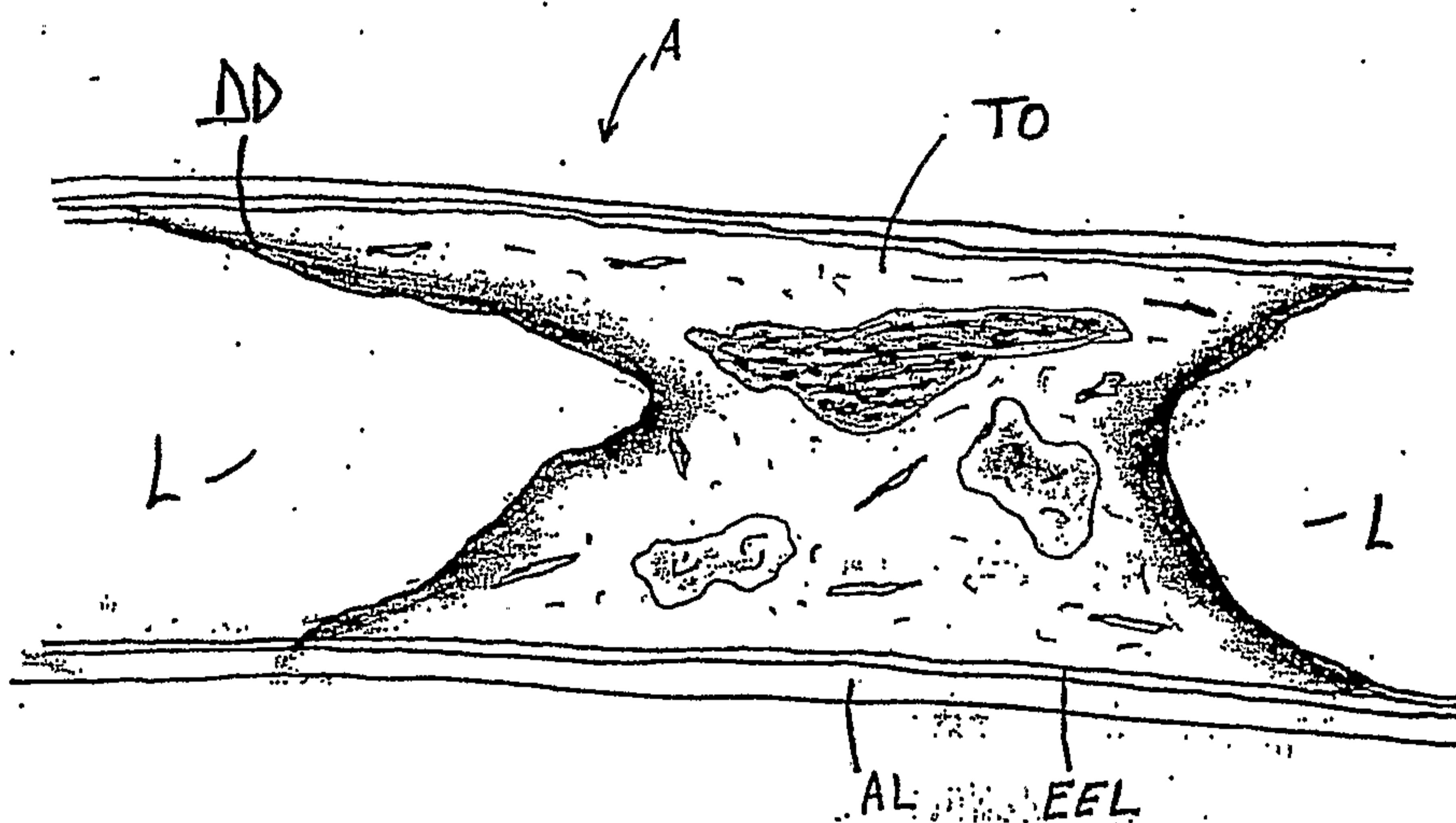


FIG 2B

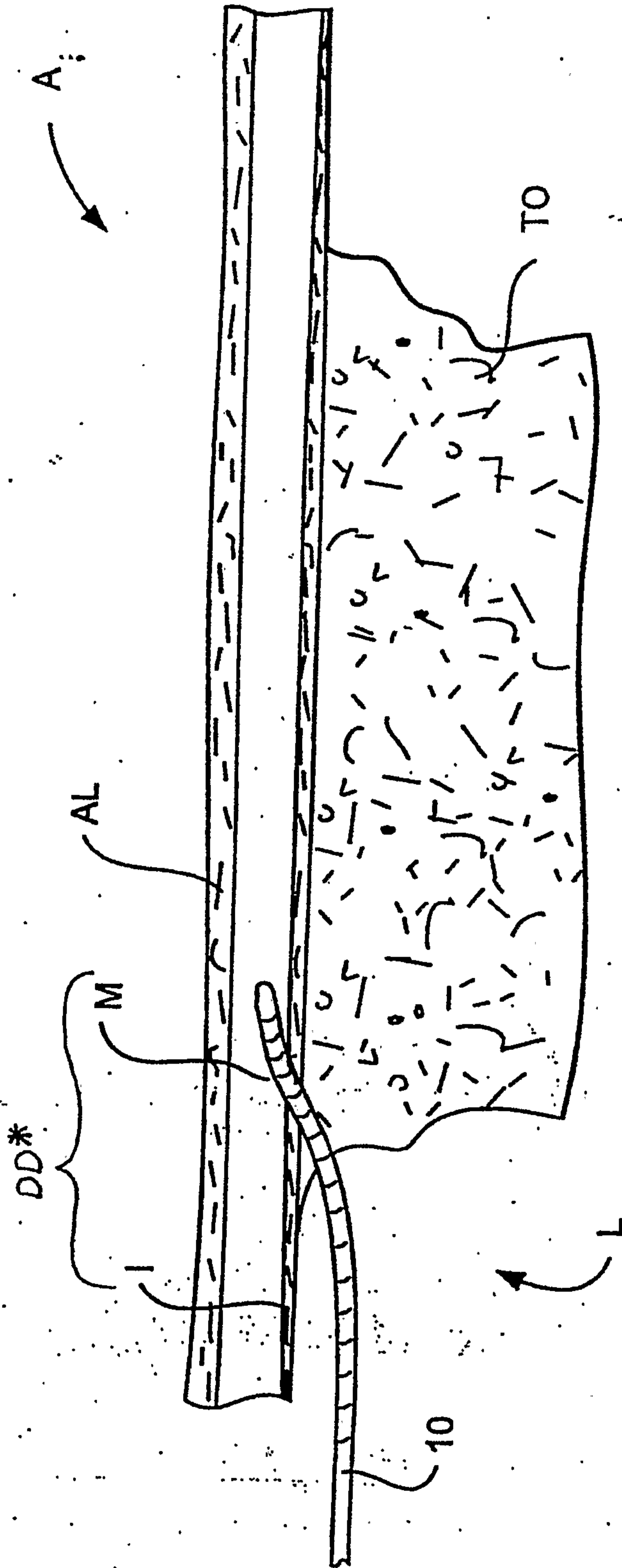


FIG. 3A

* THE INTIMA (I) AND MEDIA (M) LAYERS OF ADVANCED ATHEROSCLEROTIC OCCLUSIONS MAY EVOLVE INTO A HETEROGENEOUS LAYER OF DIFFUSE DISEASE (DD).

*

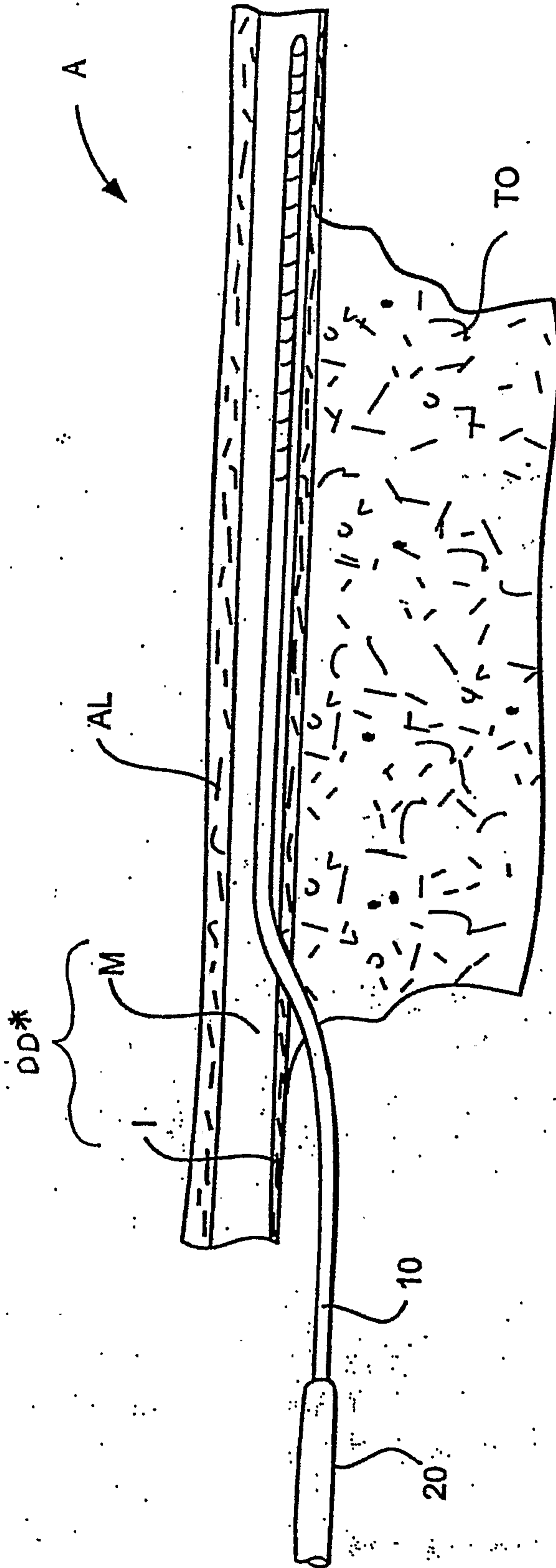


FIG. 3B

* THE INTIMA (I) AND MEDIA (M) LAYERS OF ADVANCED ATHEROSCLEROTIC OCCASIONS MAY EVOLVE INTO A HETEROGENEOUS LAYER OF DIFFUSE DISEASE (DD).

*

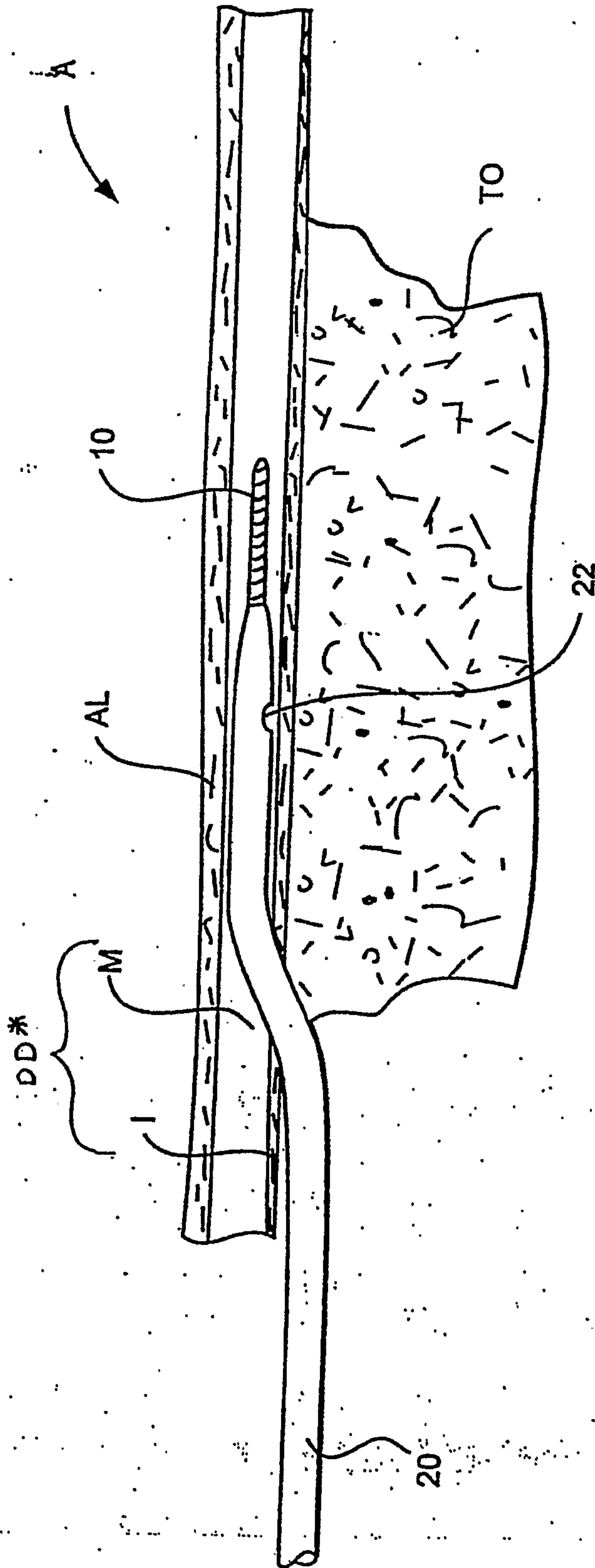


FIG. 3c

* THE INTIMA (I) AND MEDIA (M) LAYERS OF ADVANCED ATHEROSCLEROTIC OCCLUSIONS MAY EVOLVE INTO A HETEROGENEOUS LAYER OF DIFFUSE DISEASE (DD).

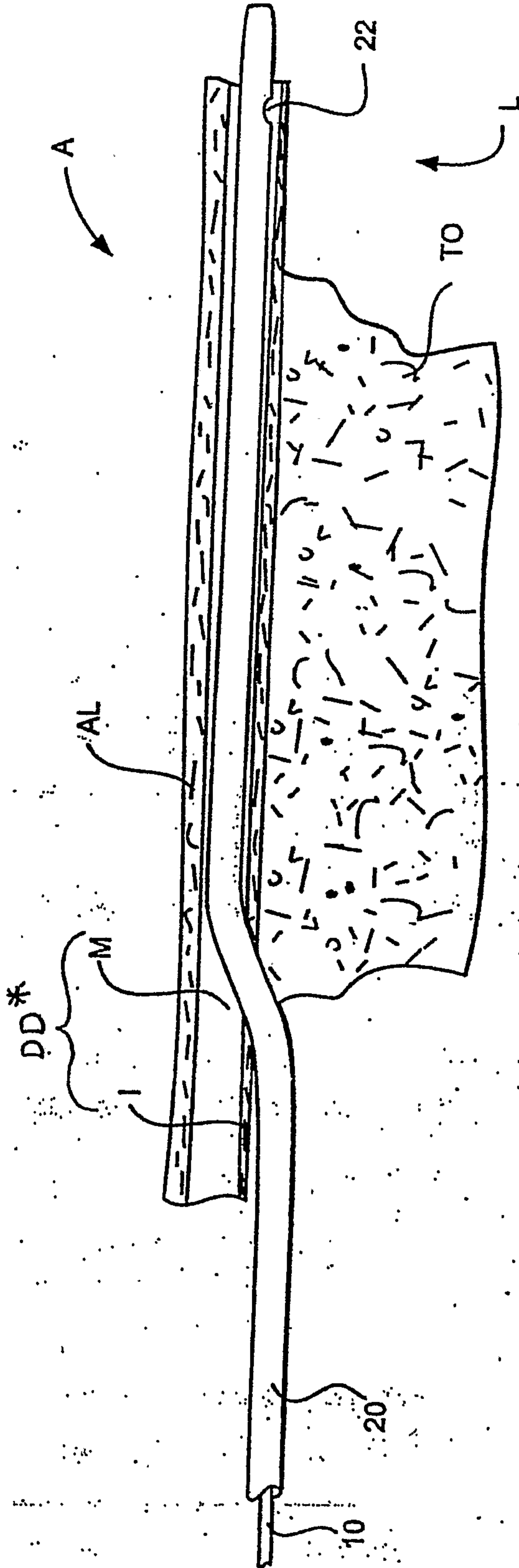


FIG. 3D

THE INTIMA (I) AND MEDIA (M) LAYERS OF ADVANCED ATHEROSCLEROTIC OCCLUSIONS MAY EVOLVE INTO A HETEROGENEOUS LAYER OF DIFFUSE DISEASE (DD).

*

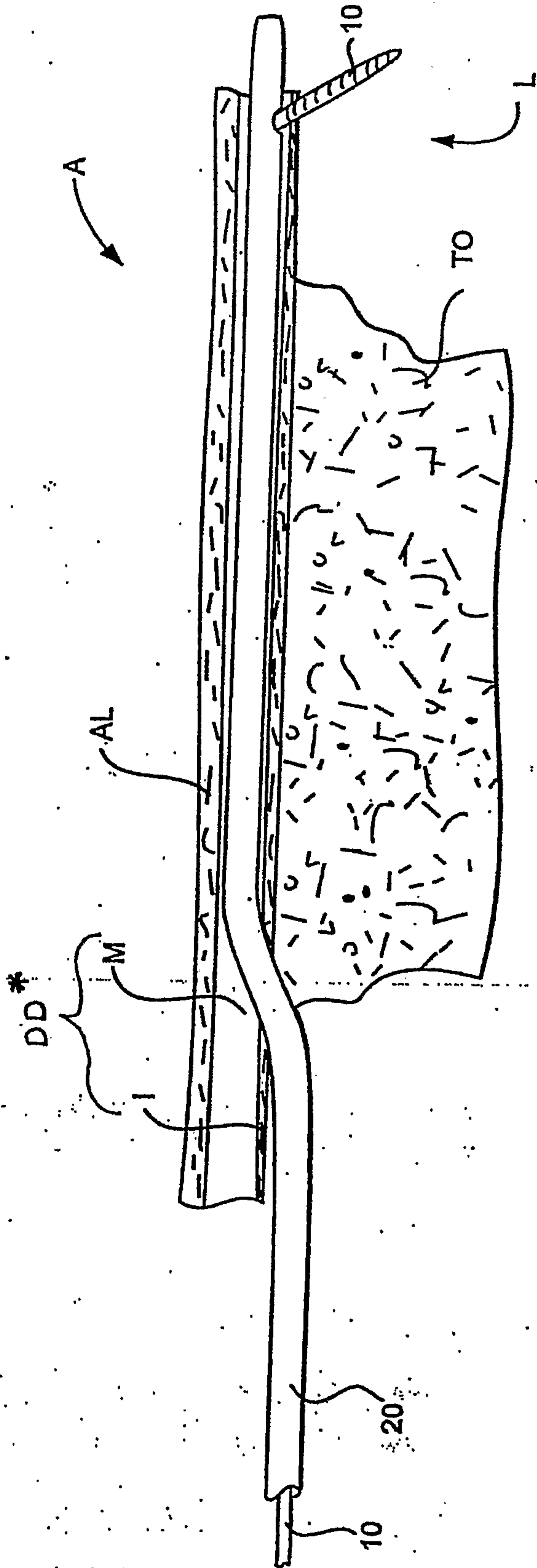


FIG. 3E

* THE INTIMA (I) AND MEDIA (M) LAYERS OF ADVANCED ATHEROSCLEROTIC OCCLUSIONS MAY EVOLVE INTO A HETEROGENEOUS LAYER OF DIFFUSE DISEASE (DD).

*

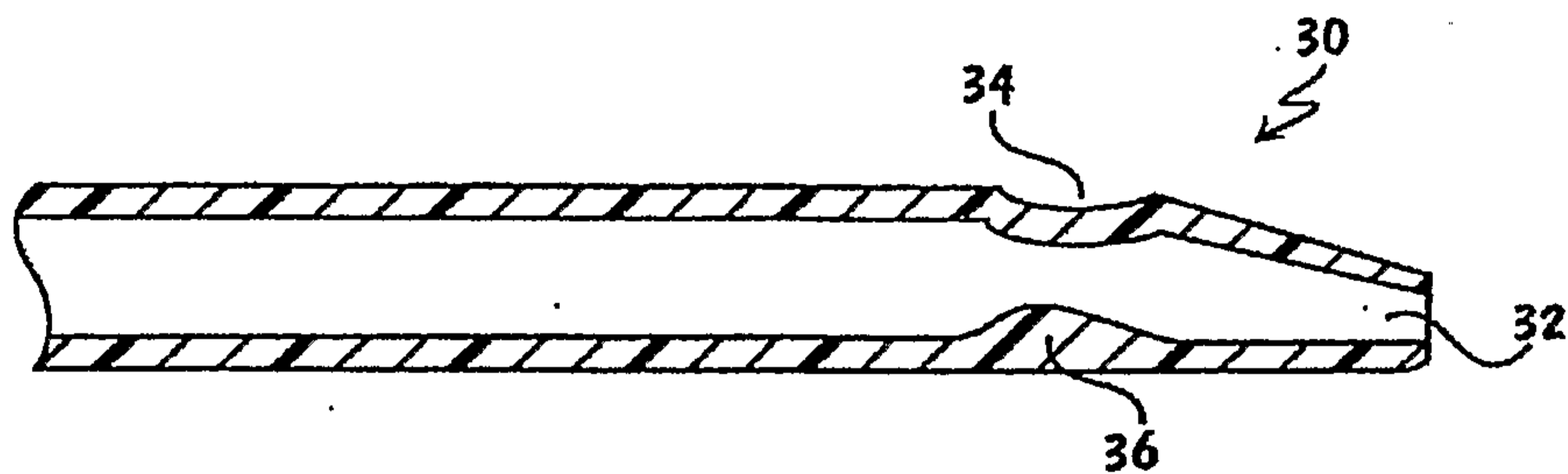


FIG. 4

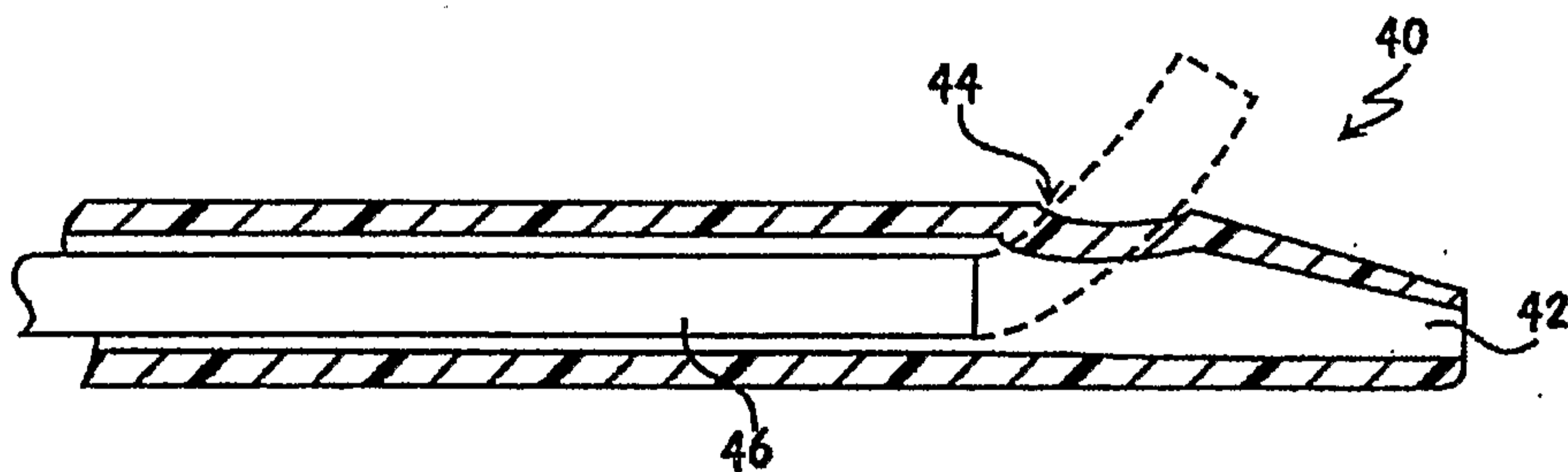


FIG. 5

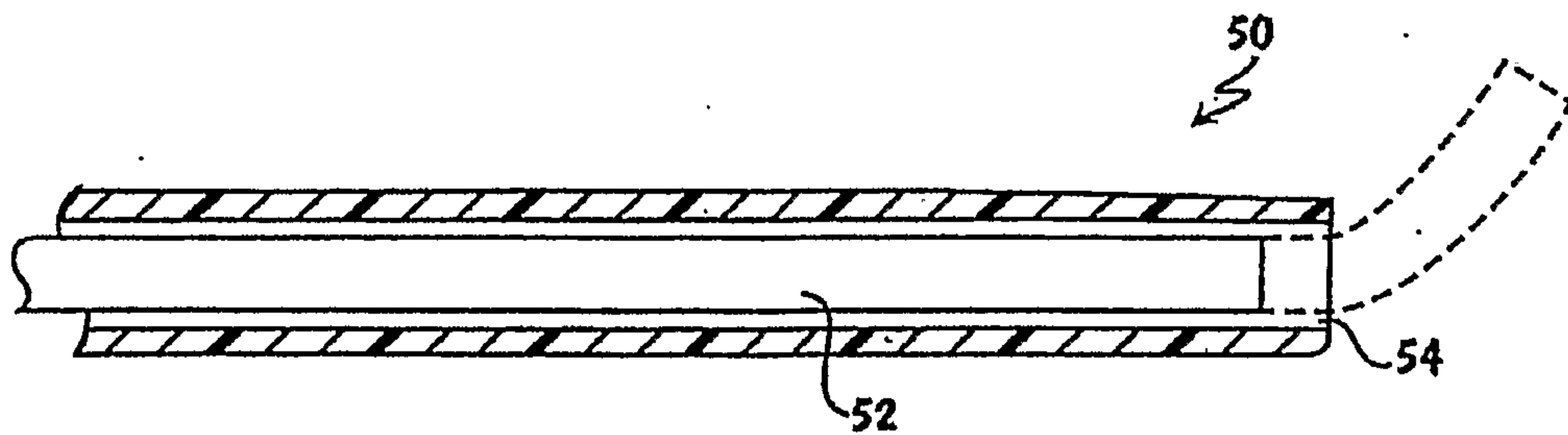


FIG. 6

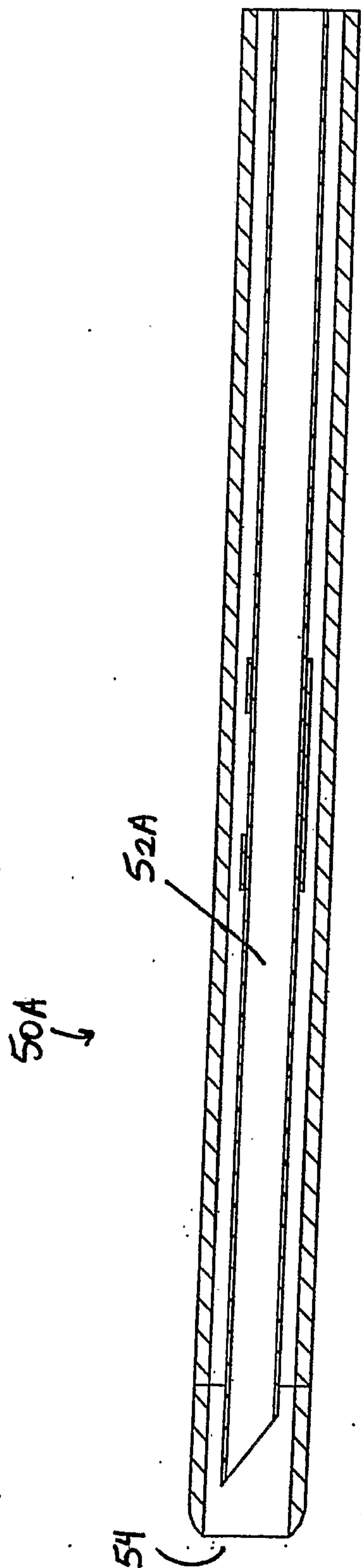


Figure 6a

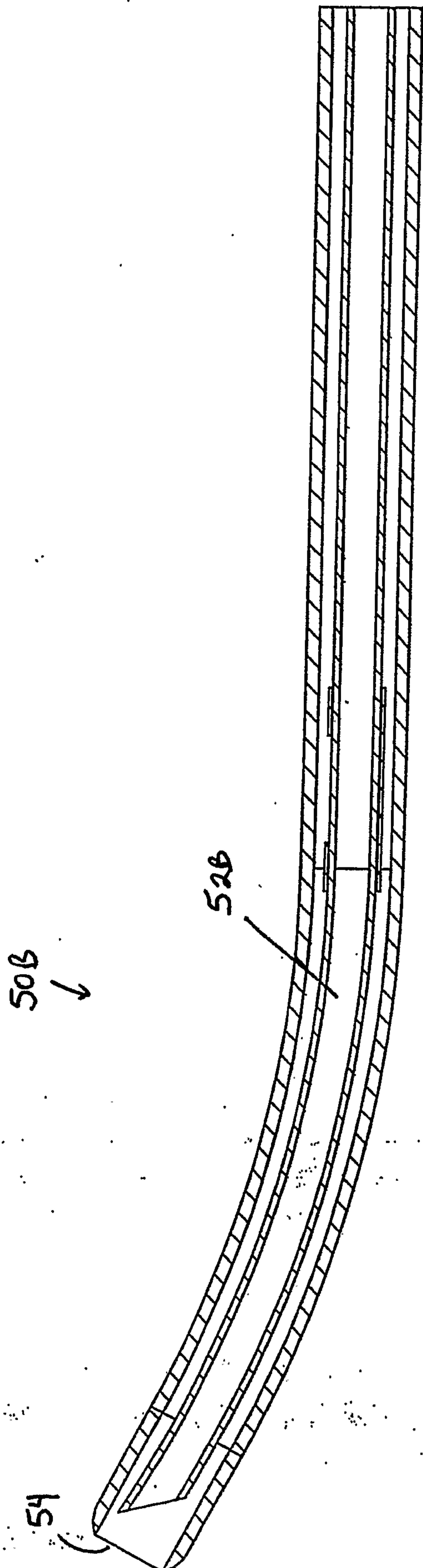


Figure 6b

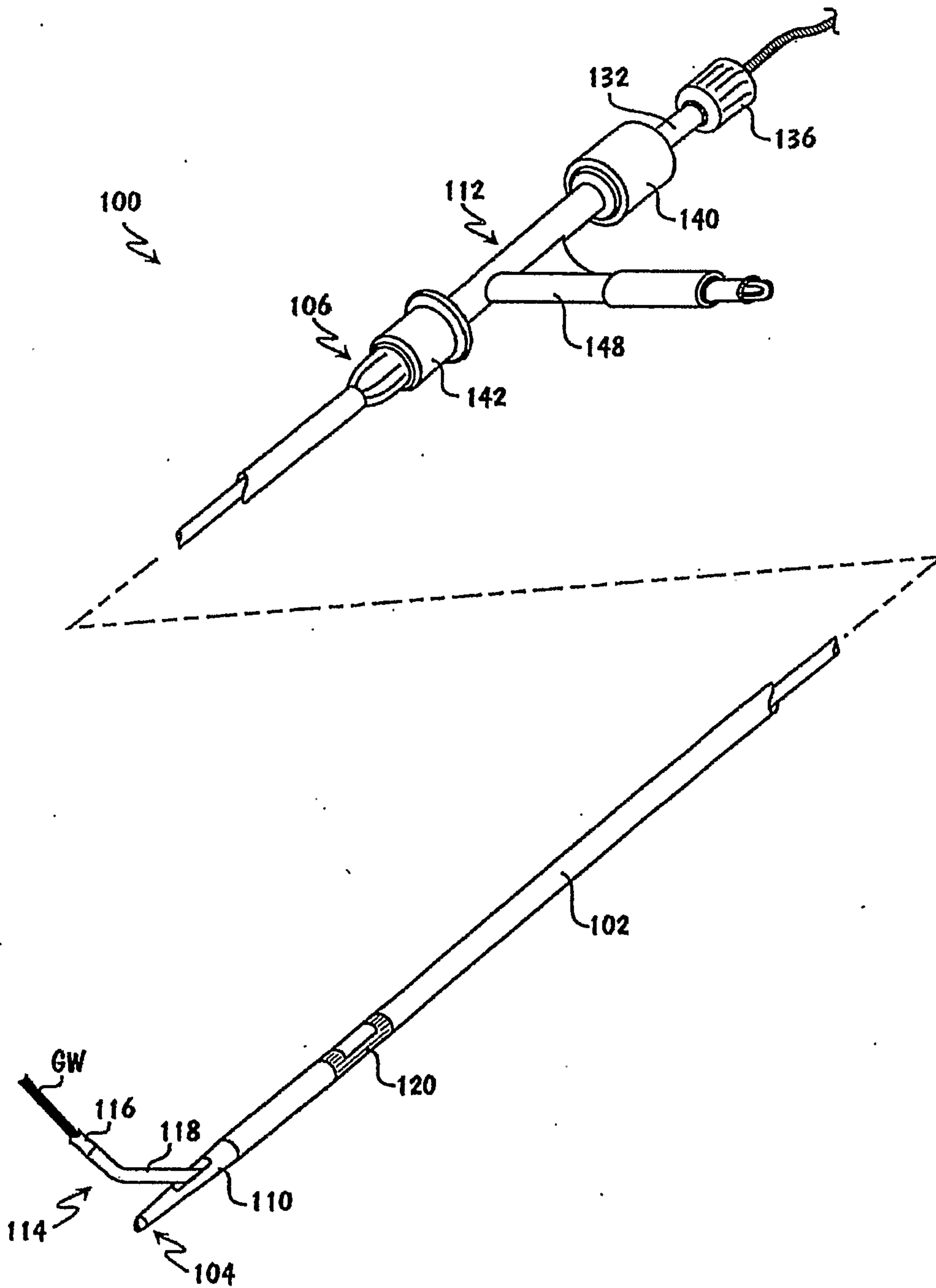


FIG. 7

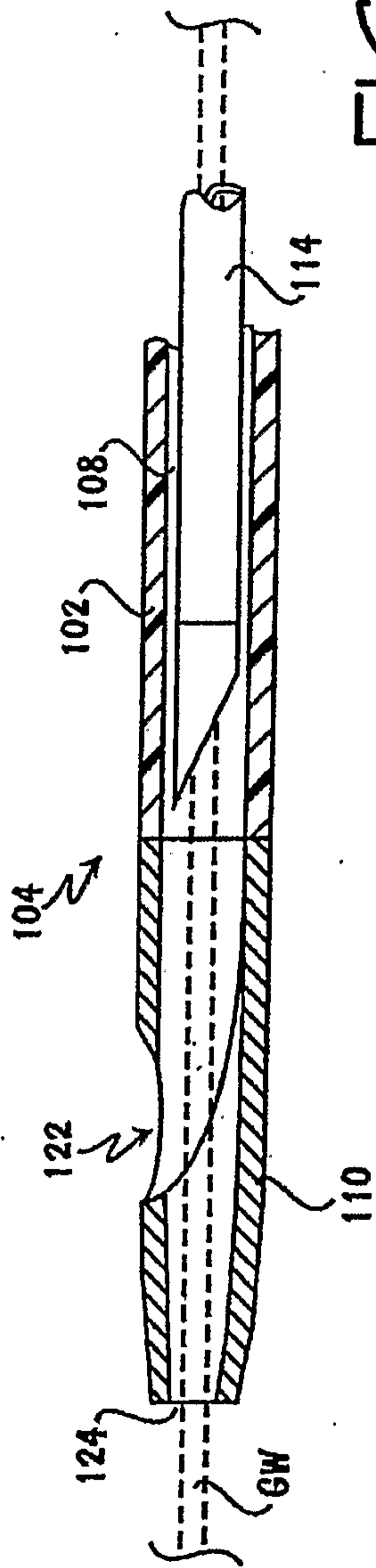


FIG. 8

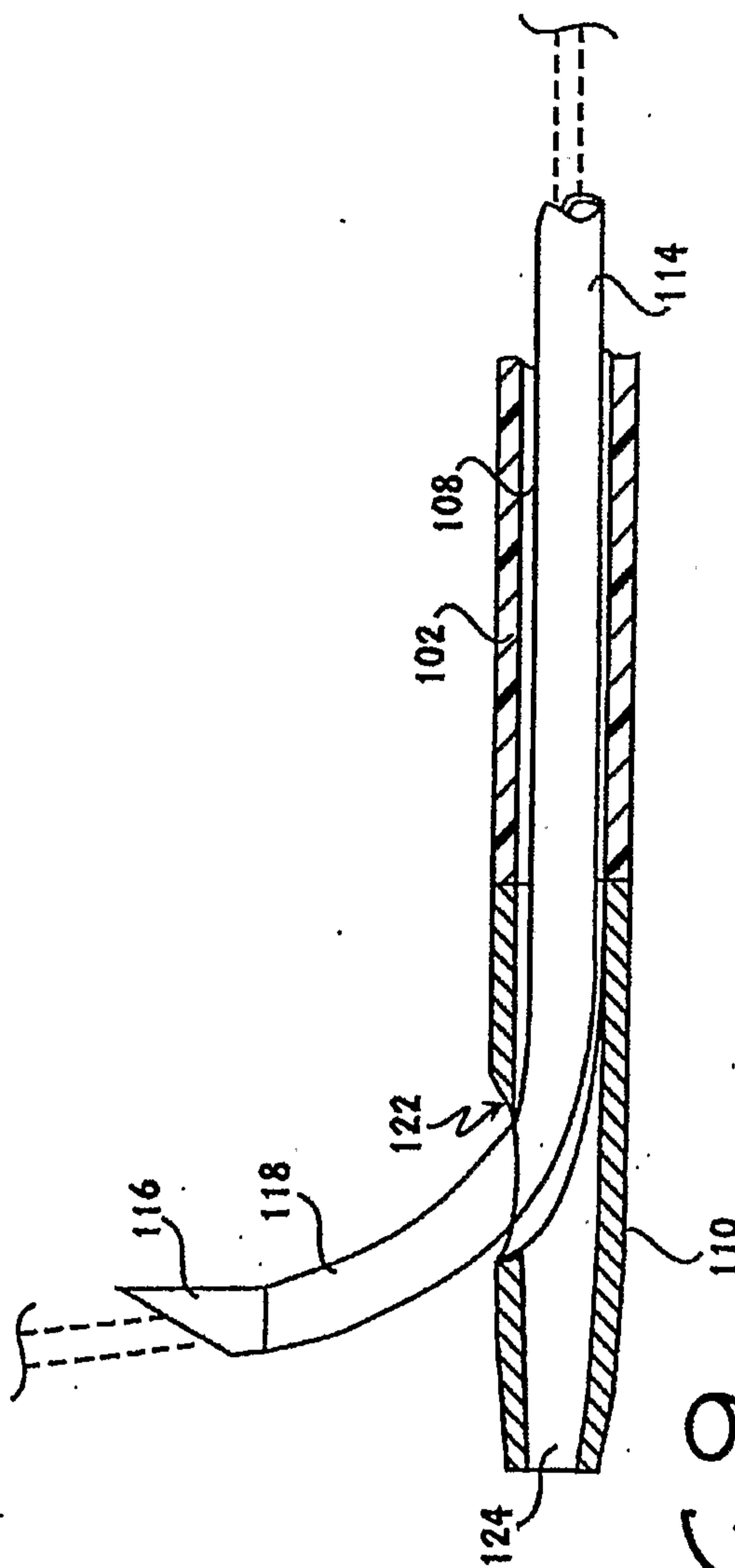


FIG. 9

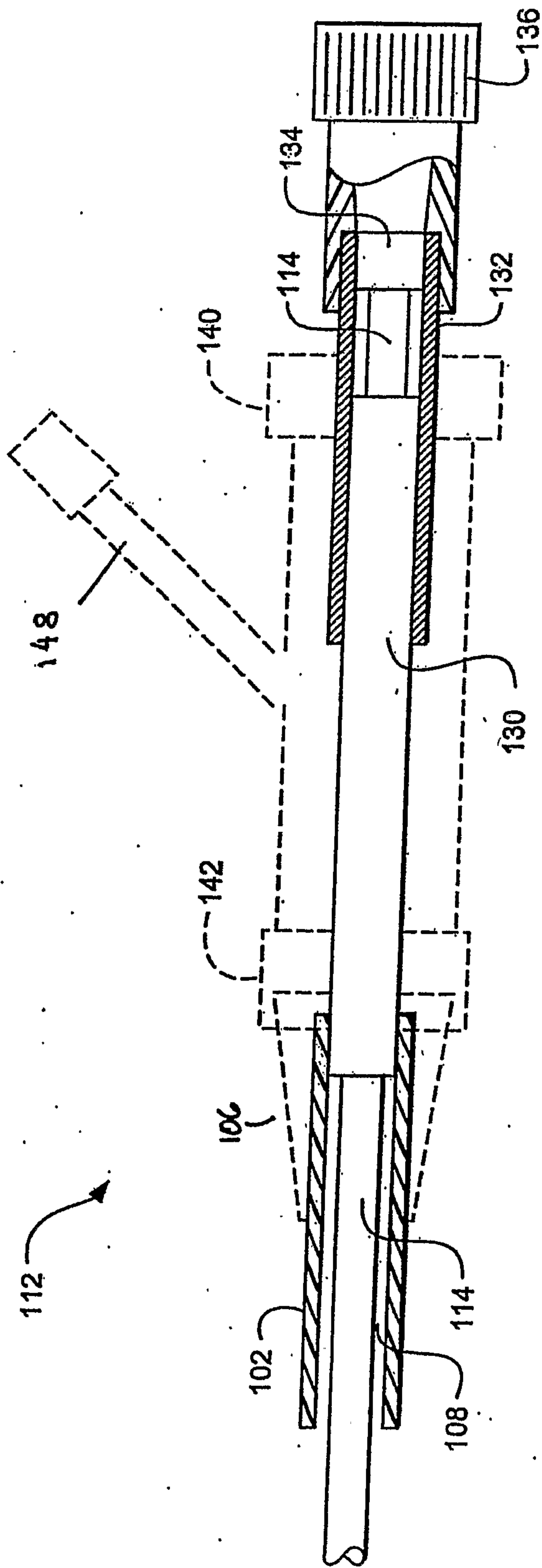


FIG. 10

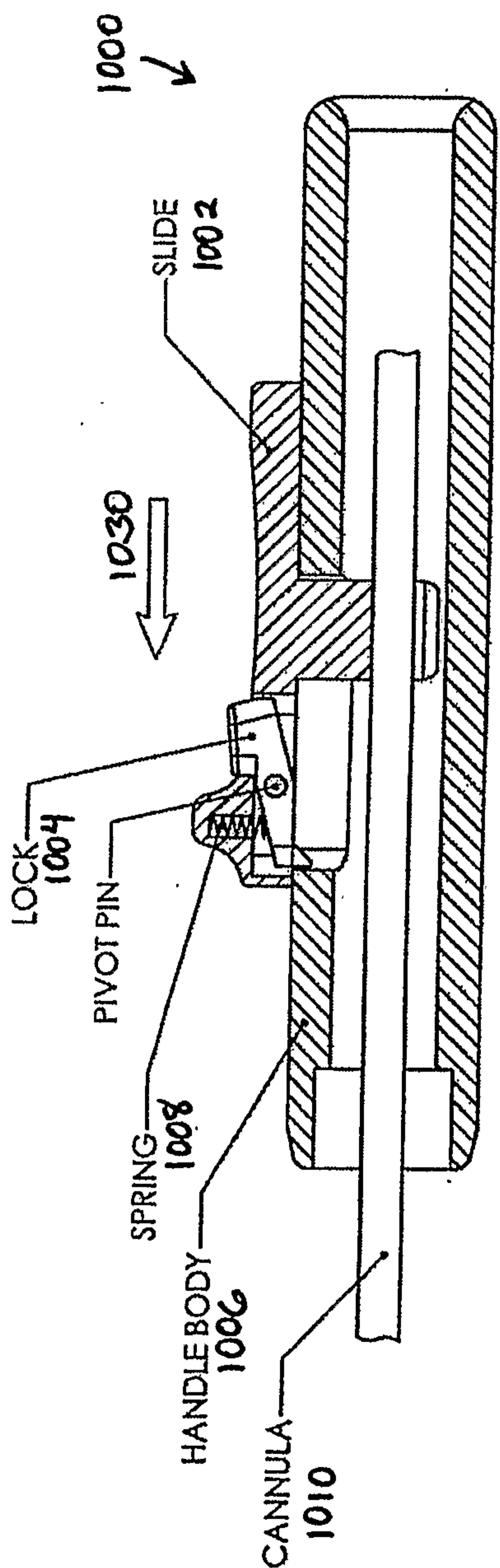


FIGURE 10A

CANNULA RETRACTED - LOCKED CONFIGURATION

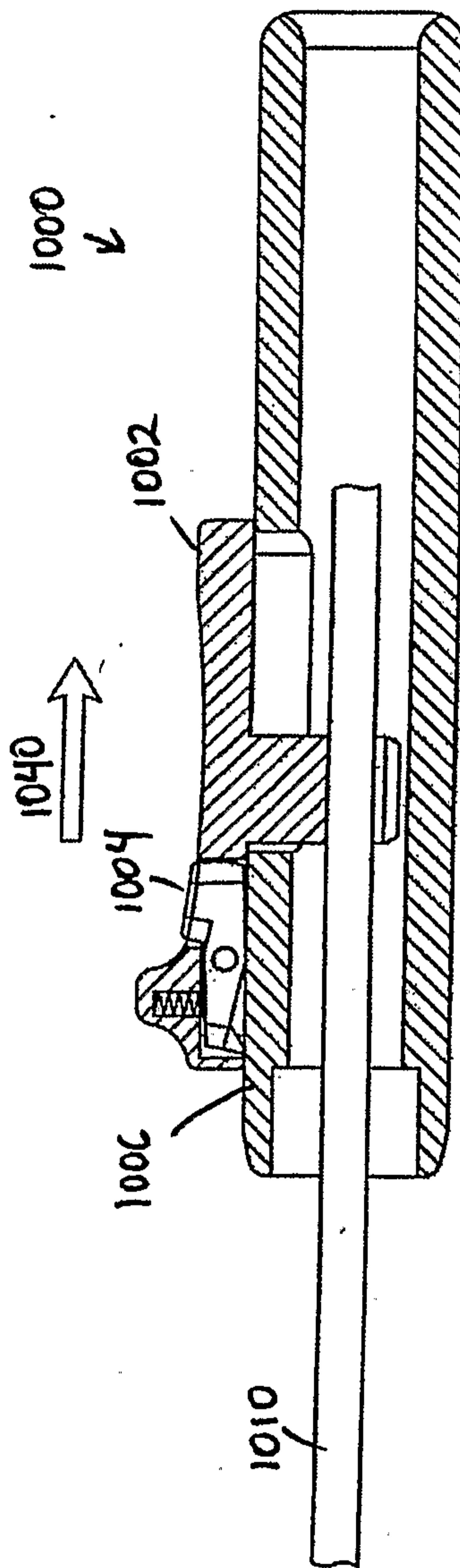


FIGURE 10B

CANNULA EXTENDED - UNLOCKED CONFIGURATION

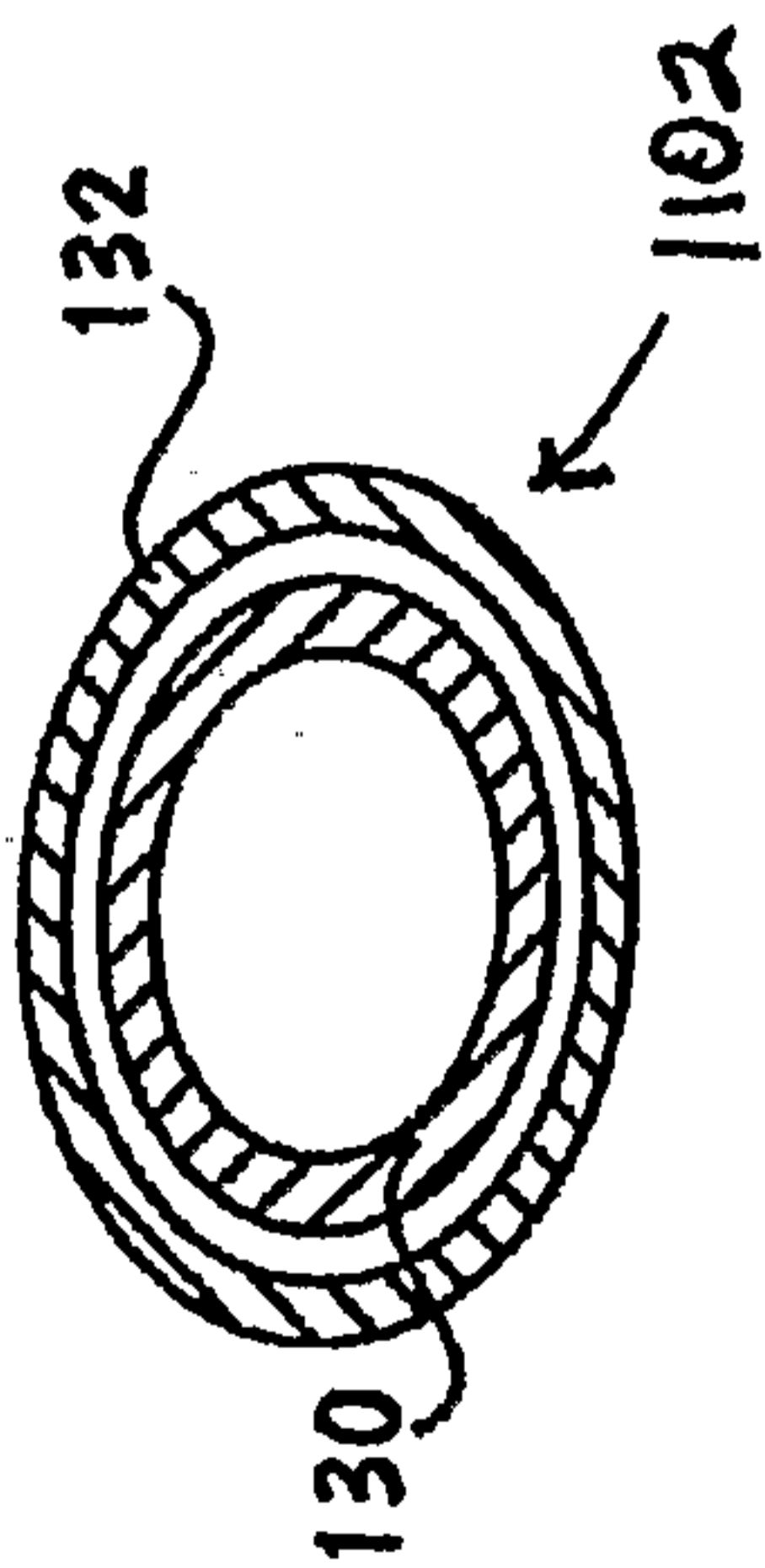


FIG. 11A

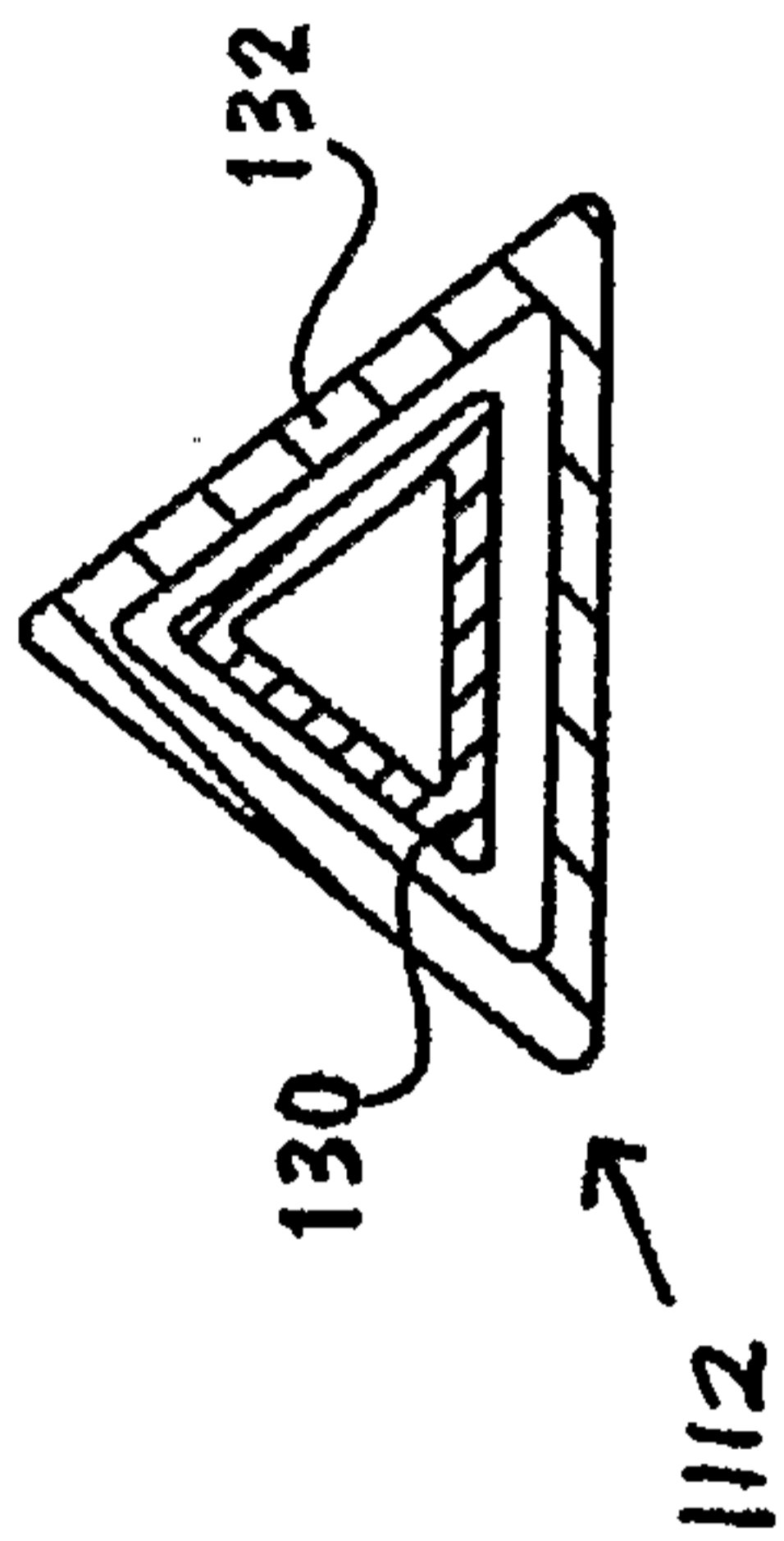


FIG. 11B

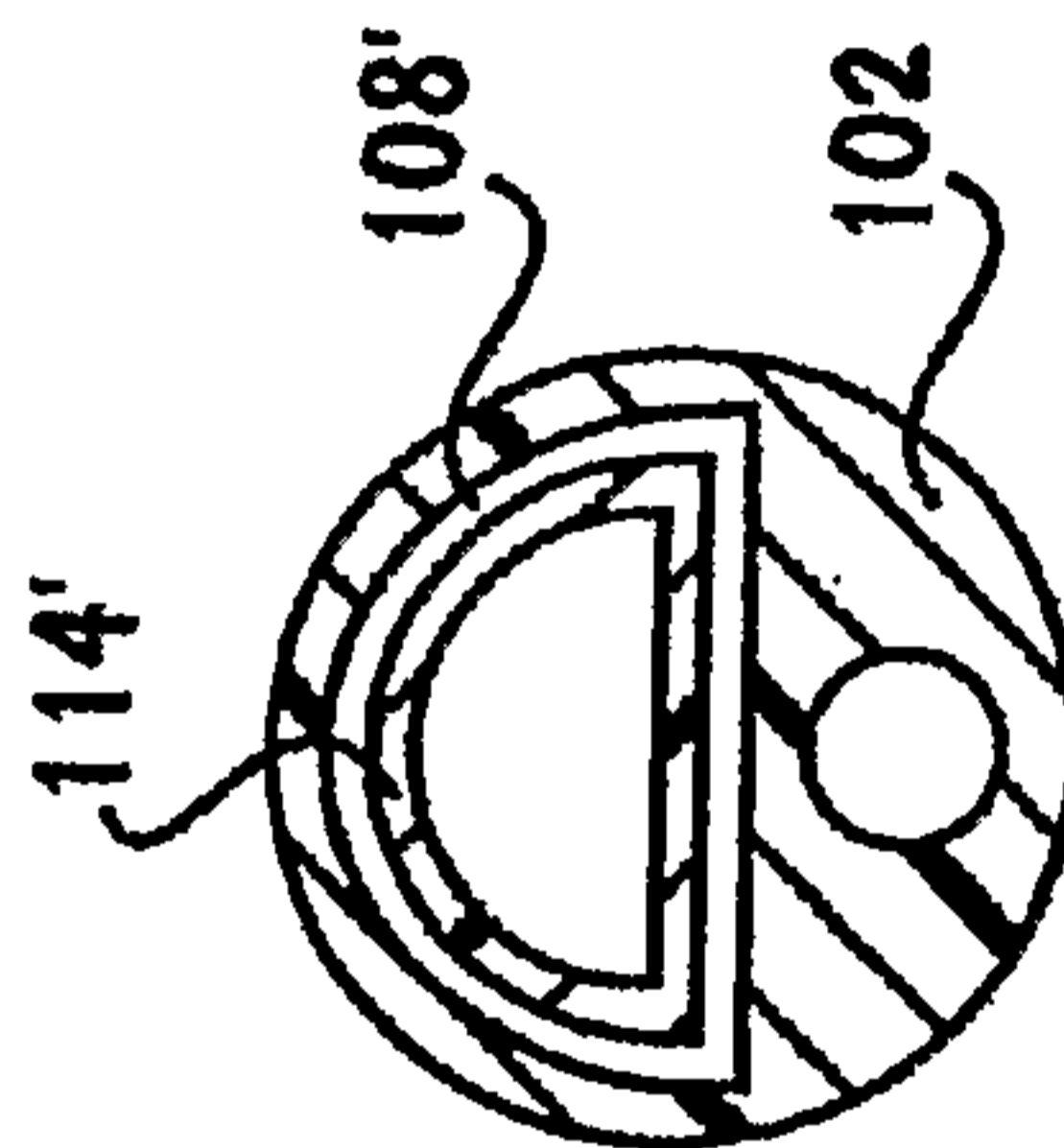


FIG. 12

1300
↓

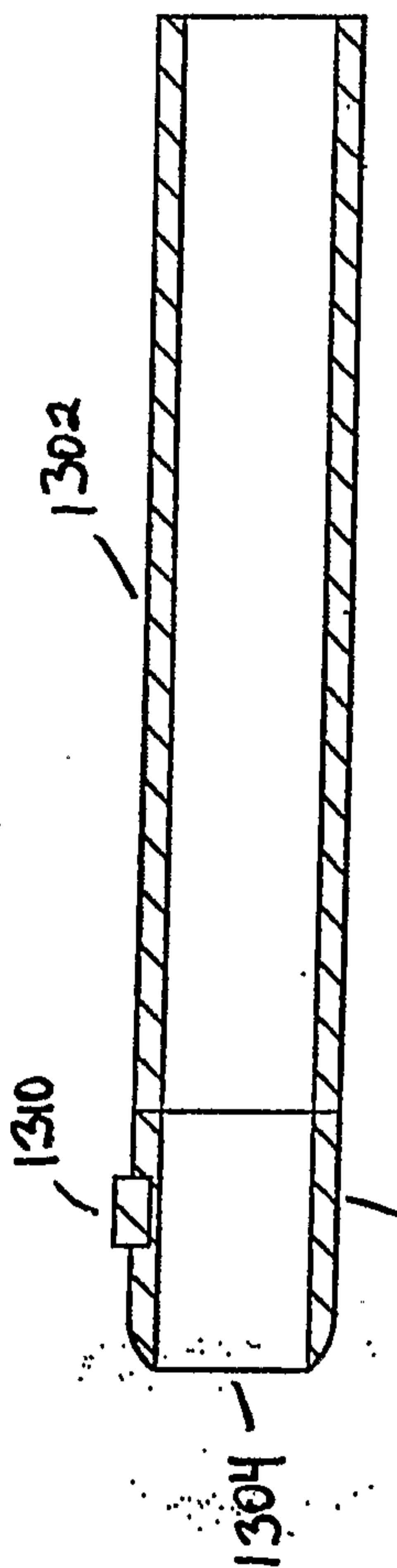


Figure 13

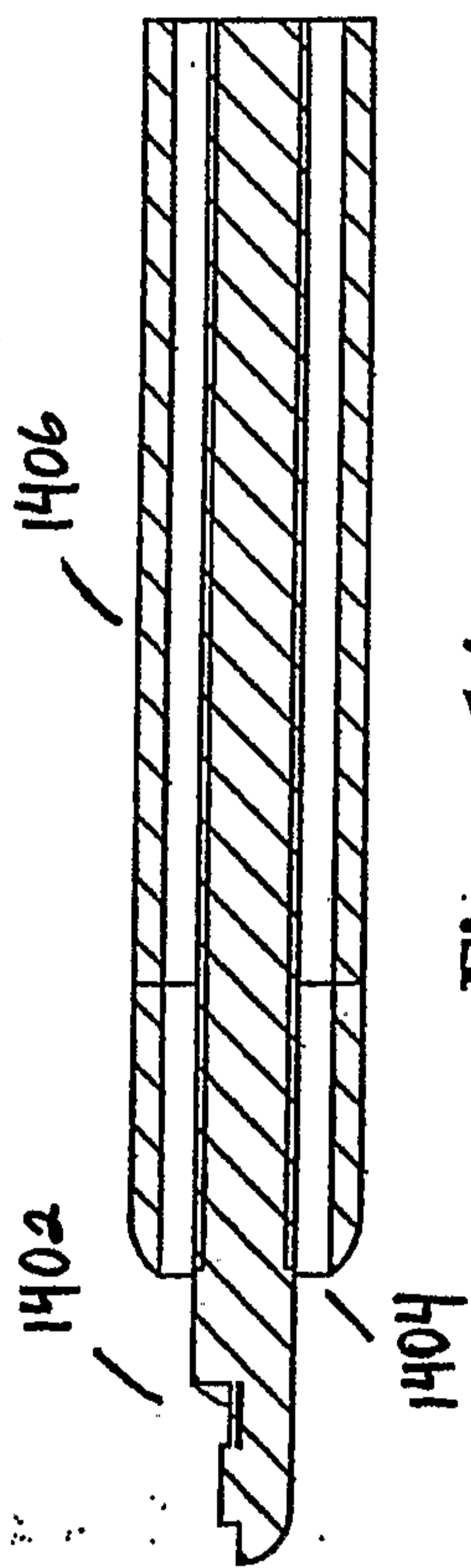


Figure 14a

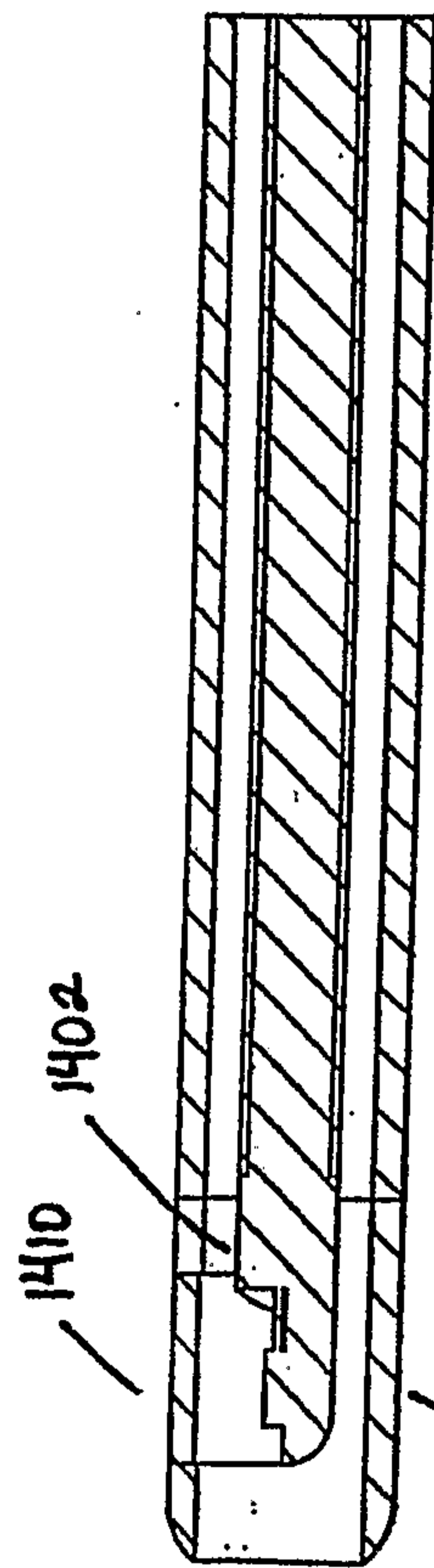


Figure 14b

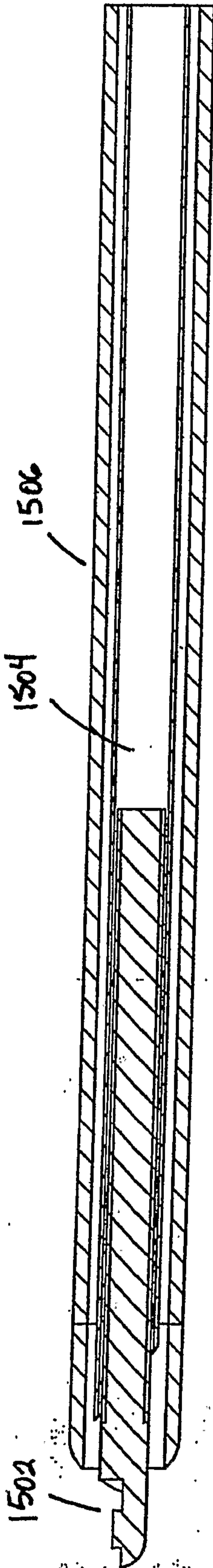


Figure 15

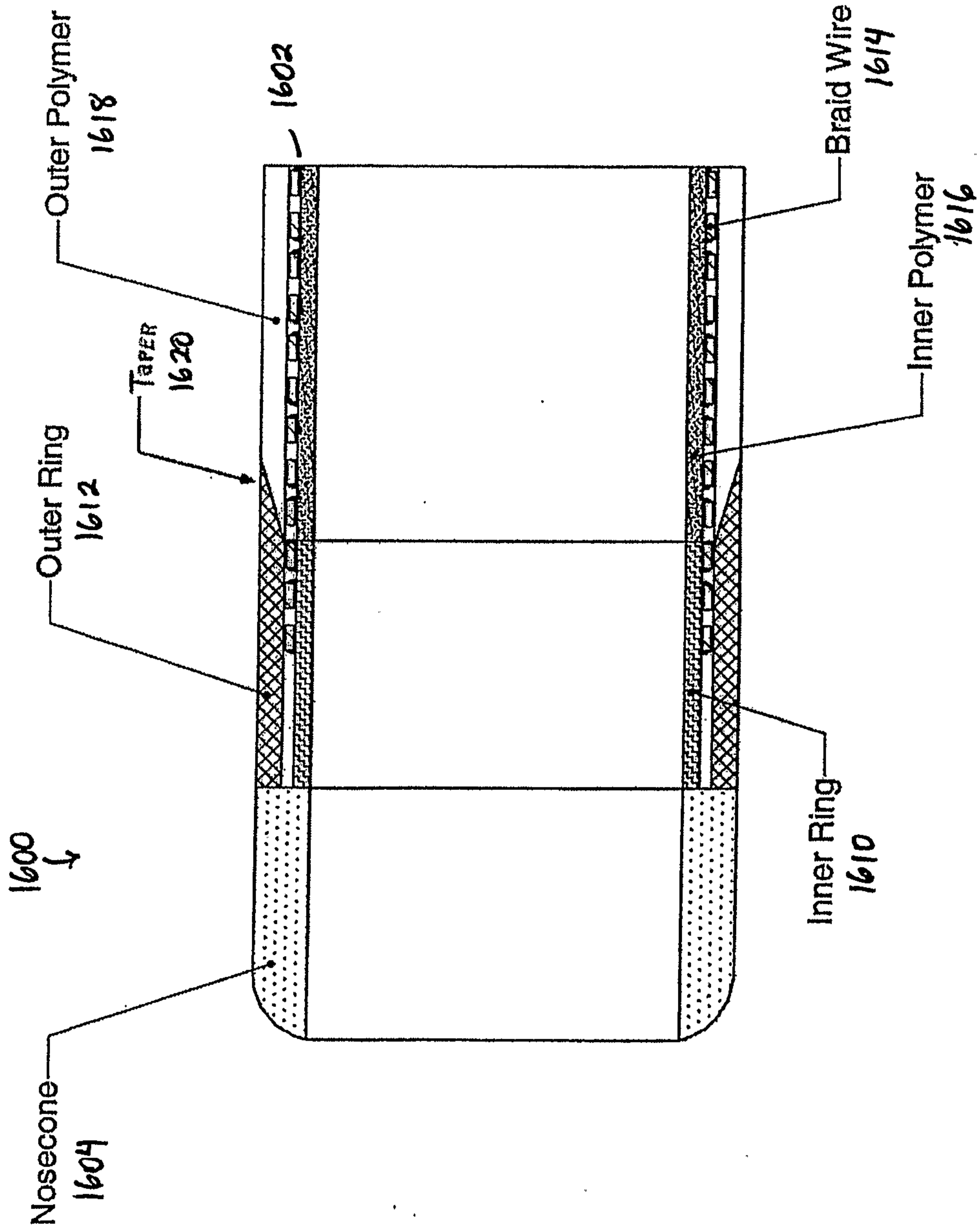


Figure 16

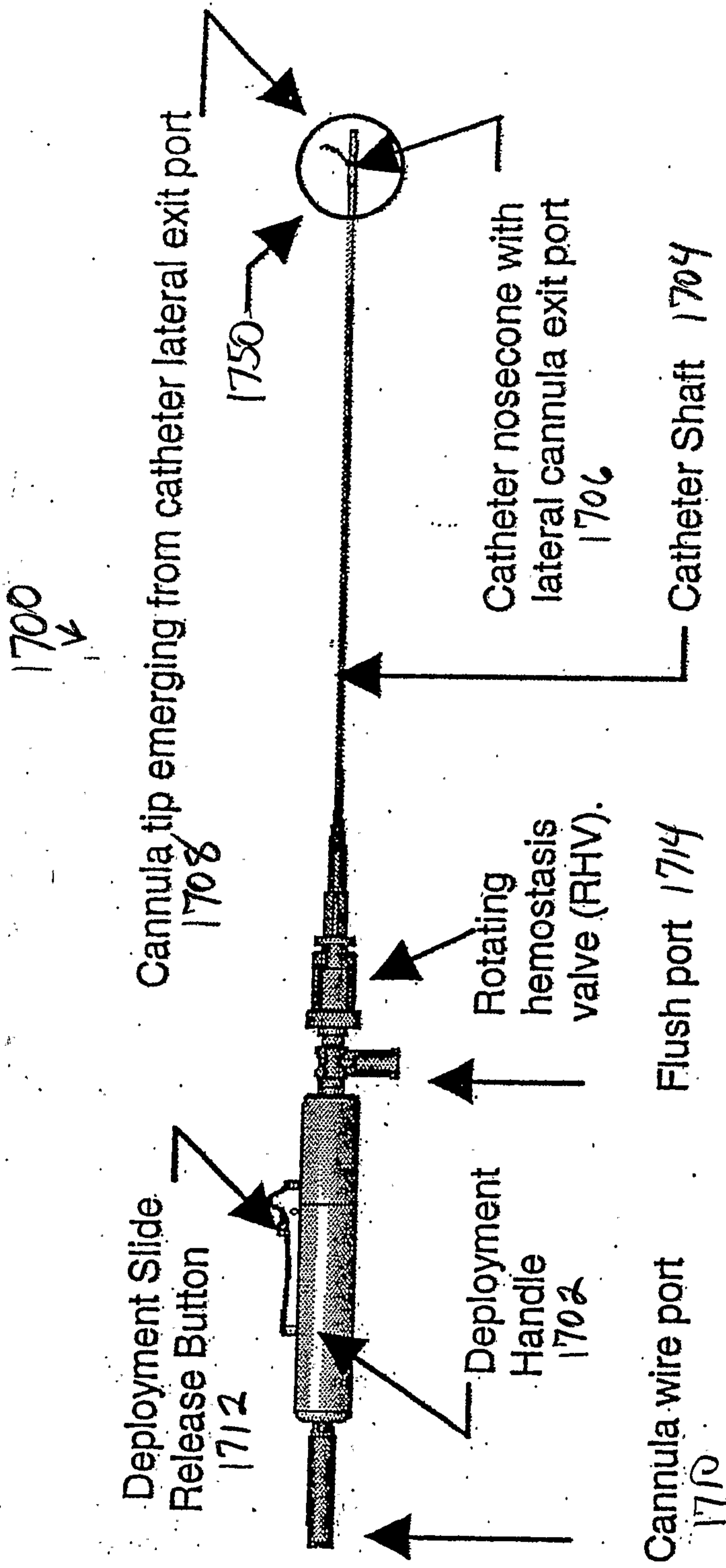


FIGURE 17

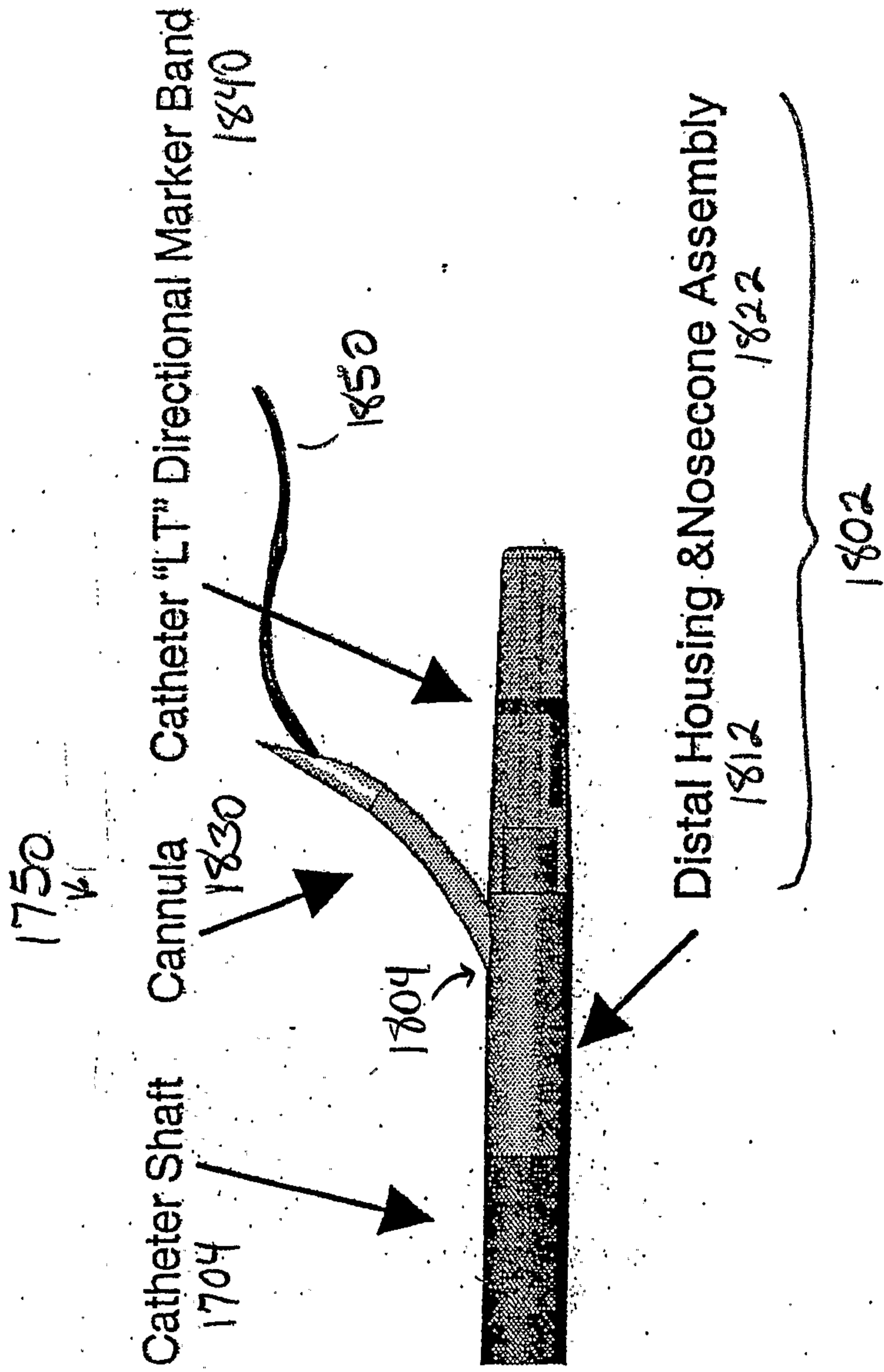
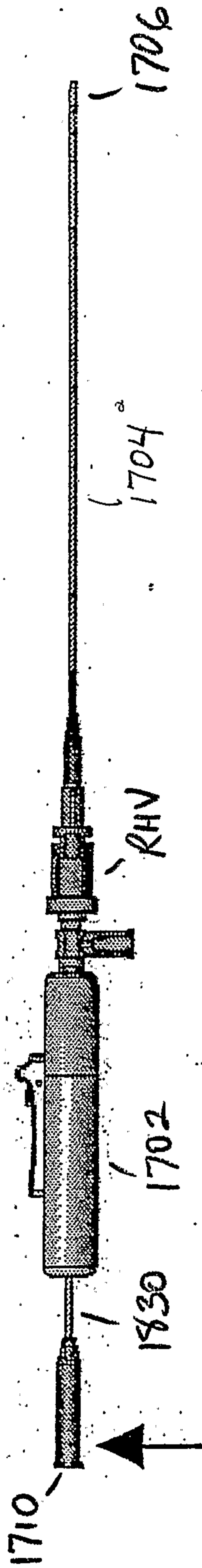


FIGURE 18



Cannula/Guide retracted position

FIGURE 19

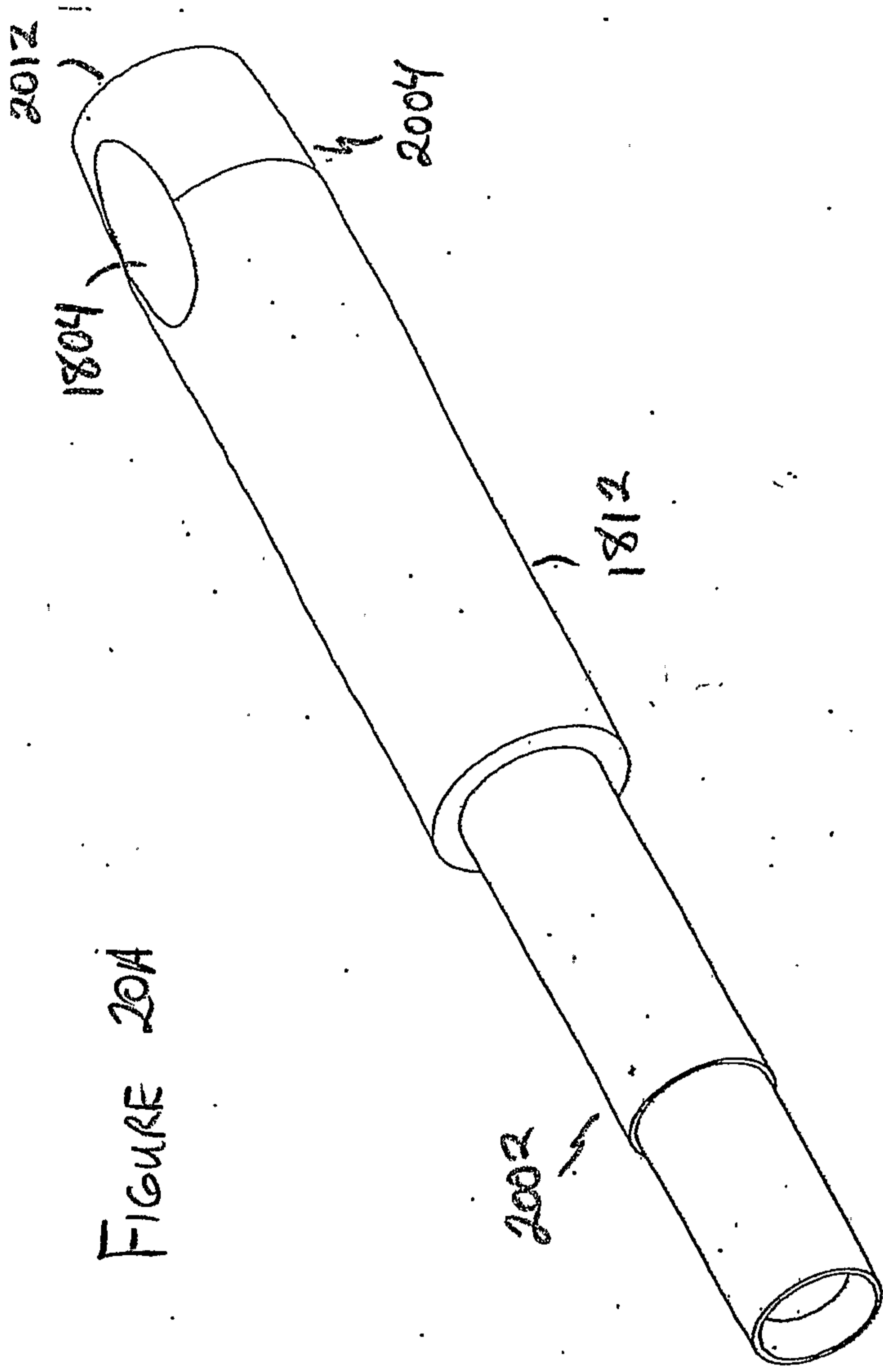


FIGURE 20A

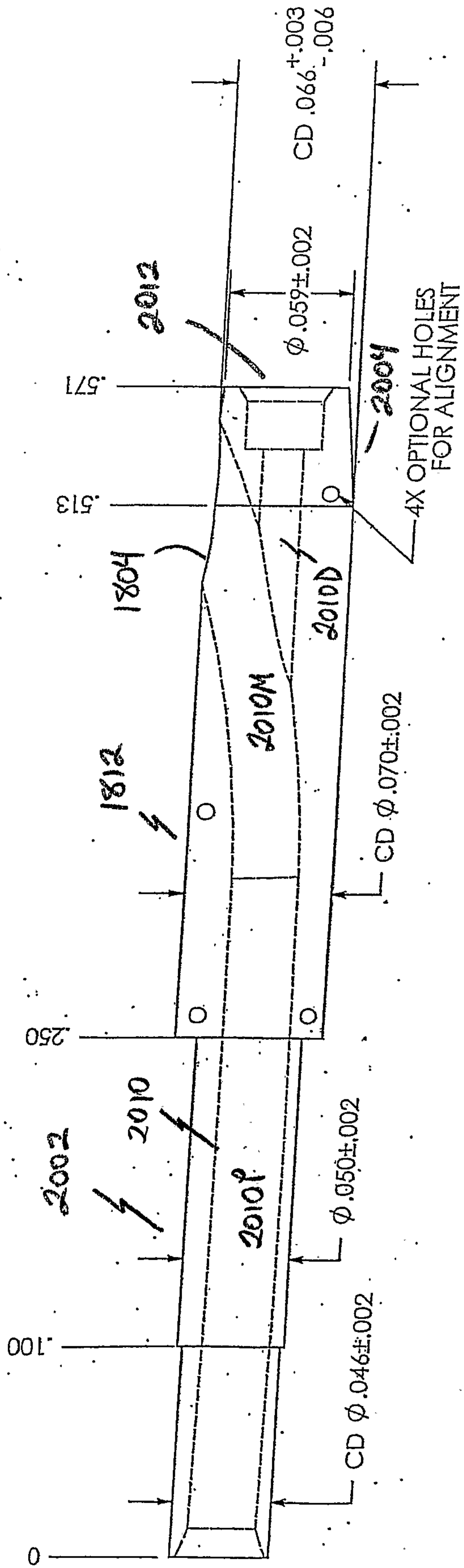
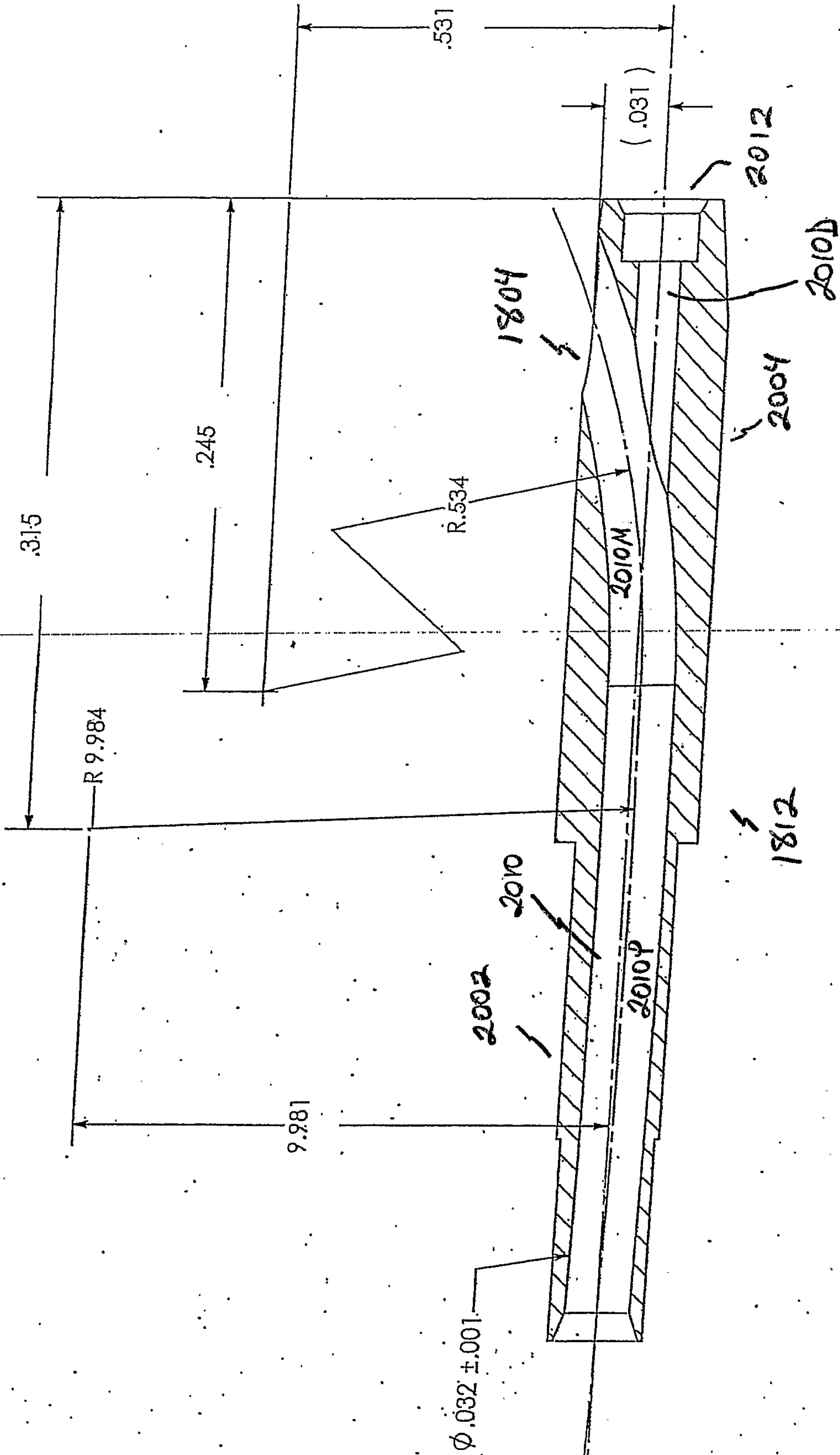


FIGURE 20B



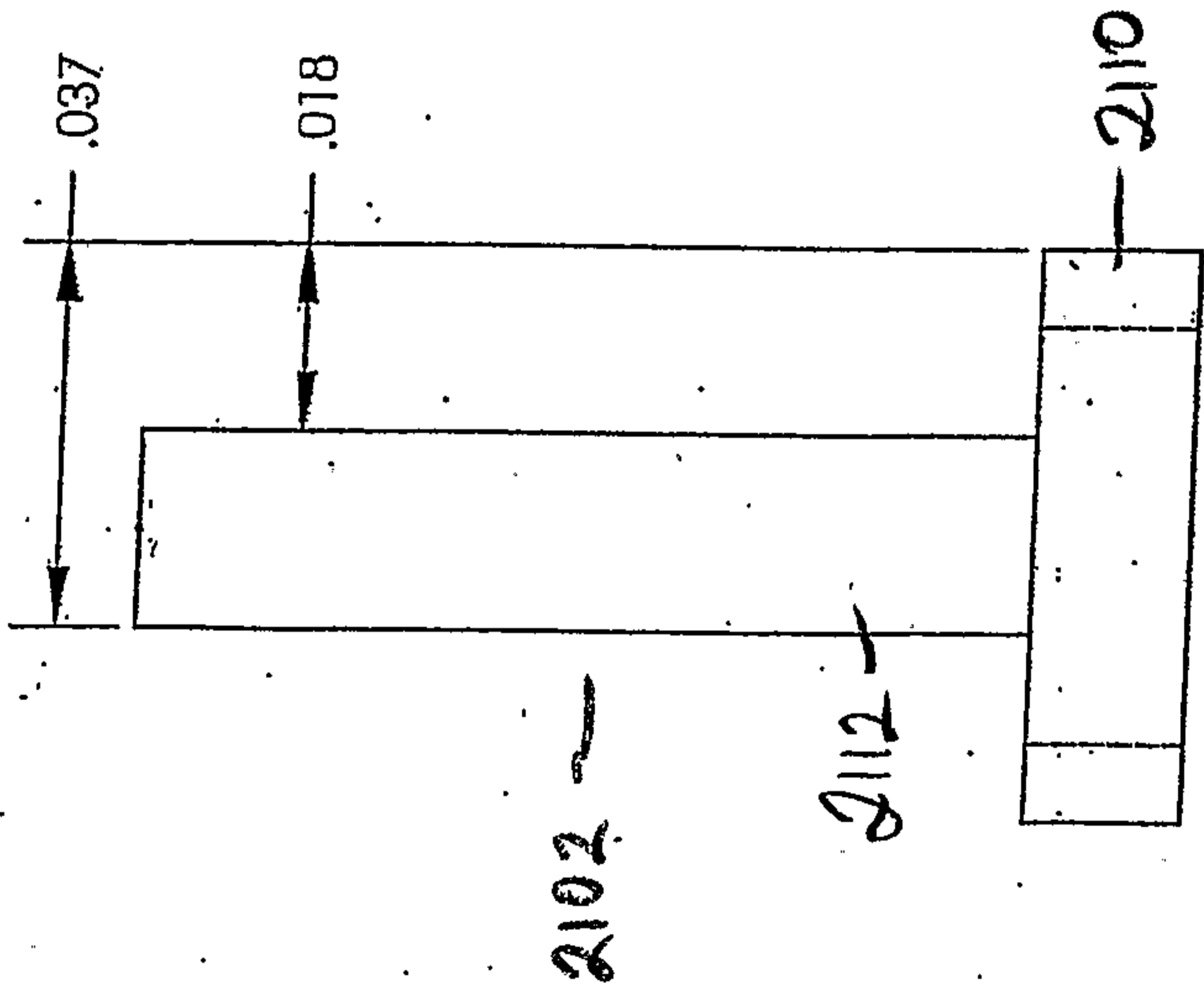


FIGURE 21B

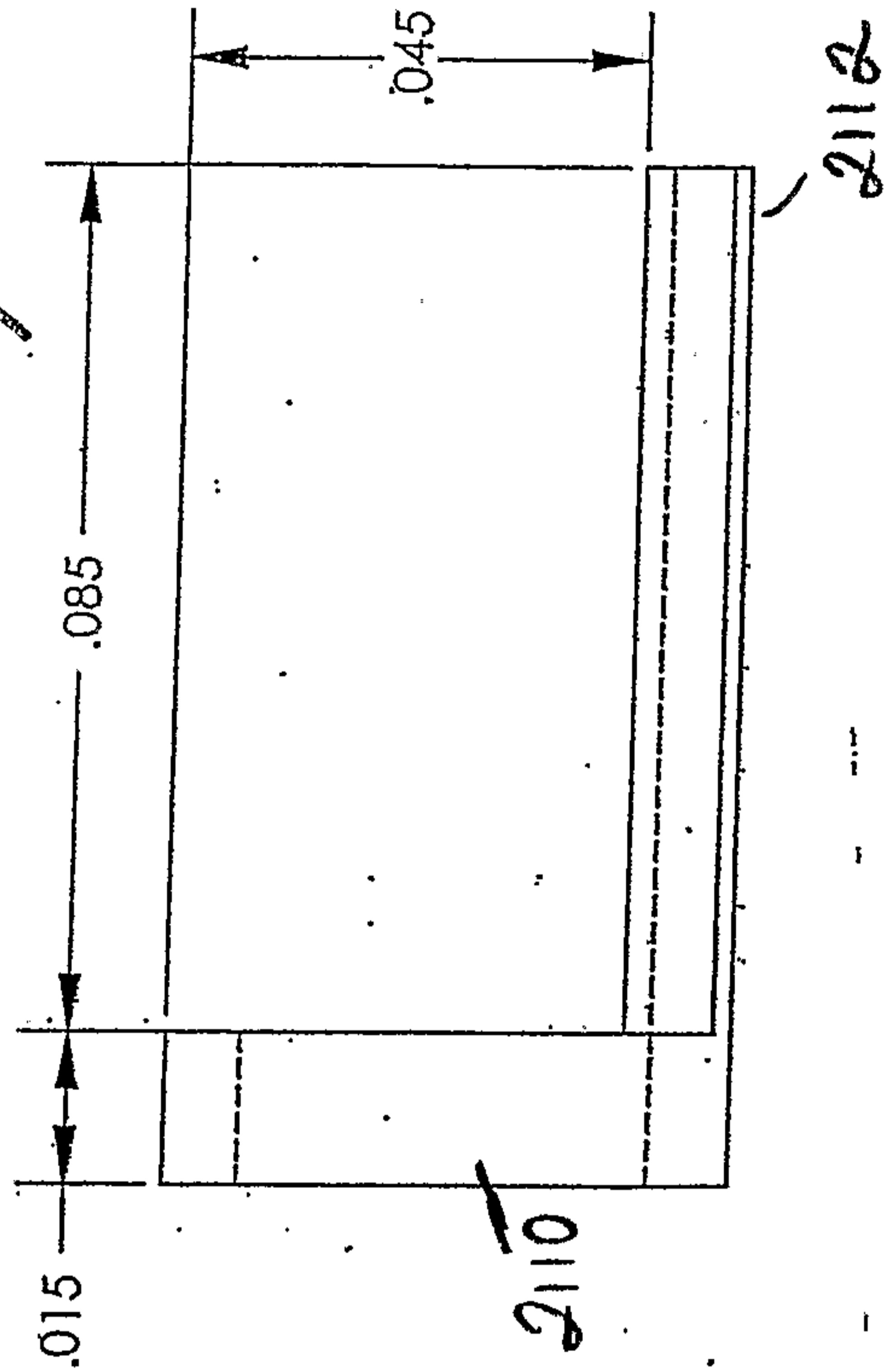


FIGURE 21A

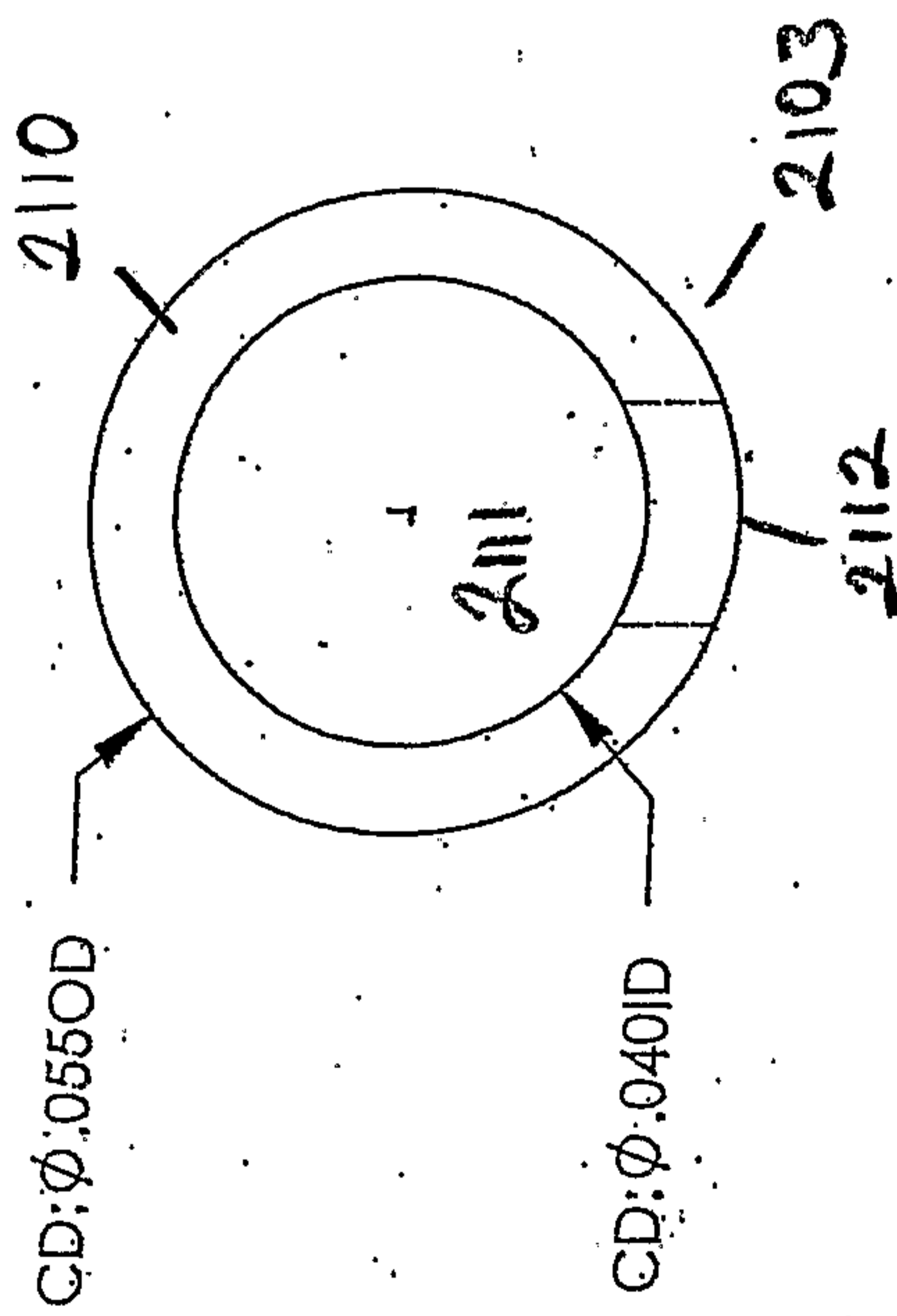


FIGURE 21C

2200

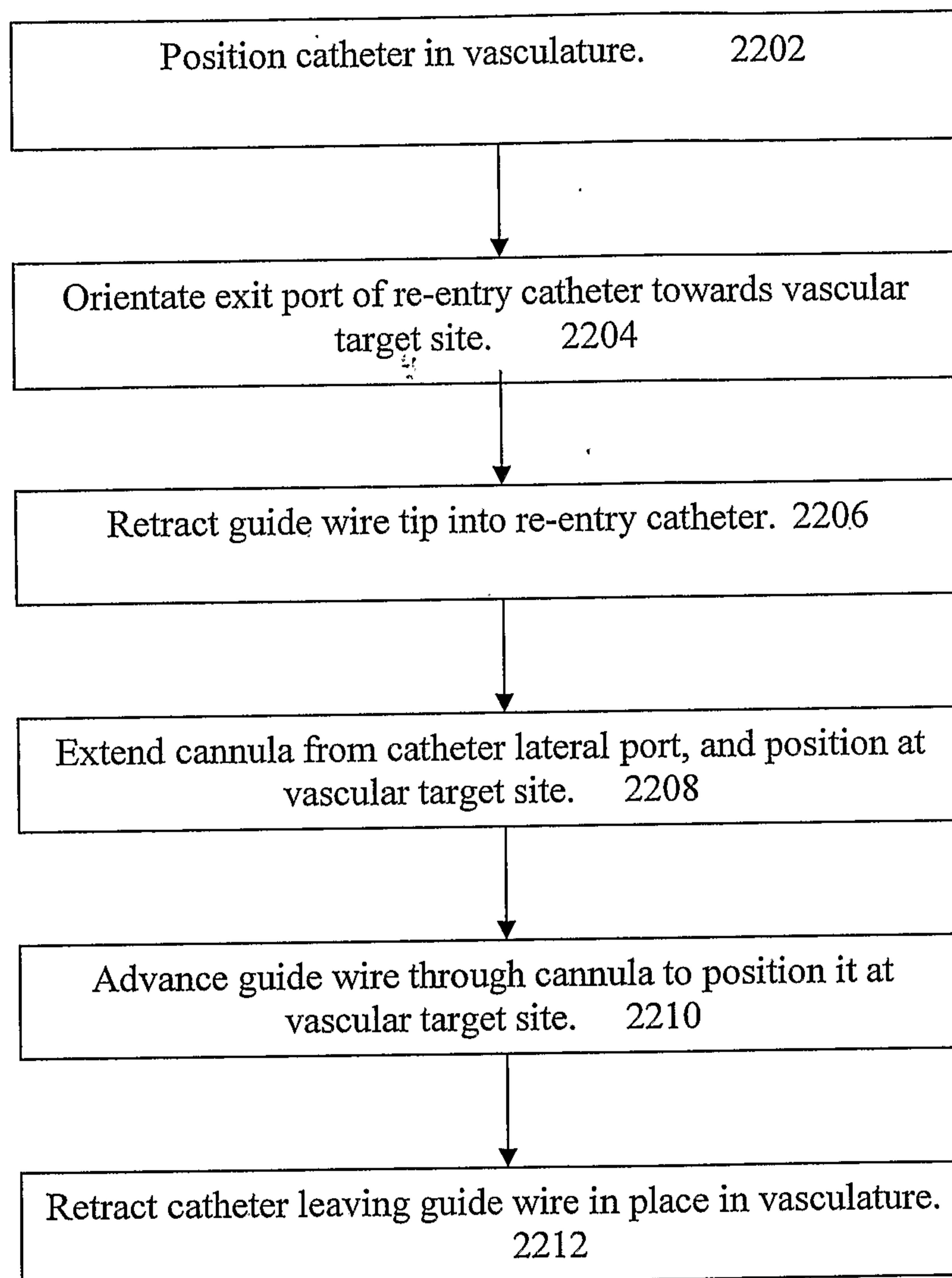
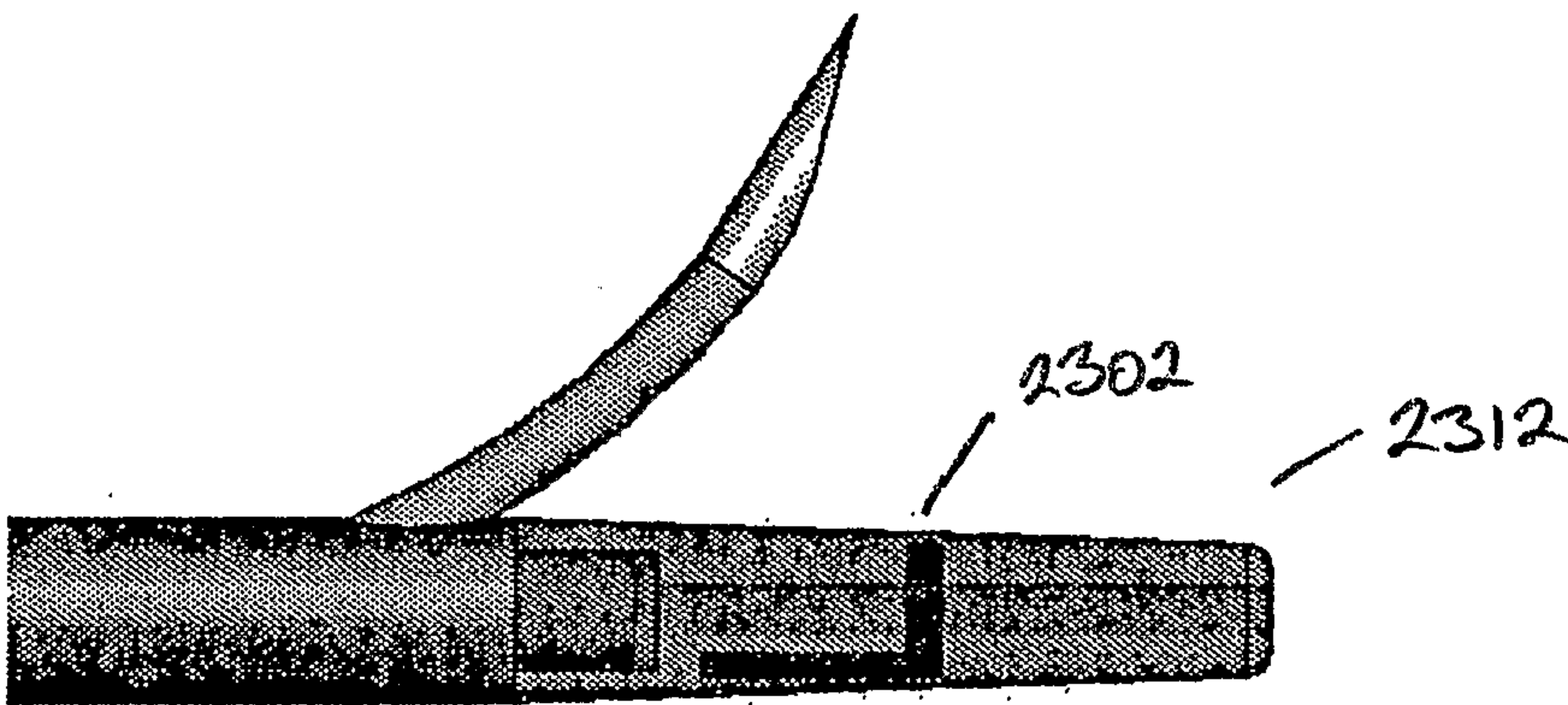
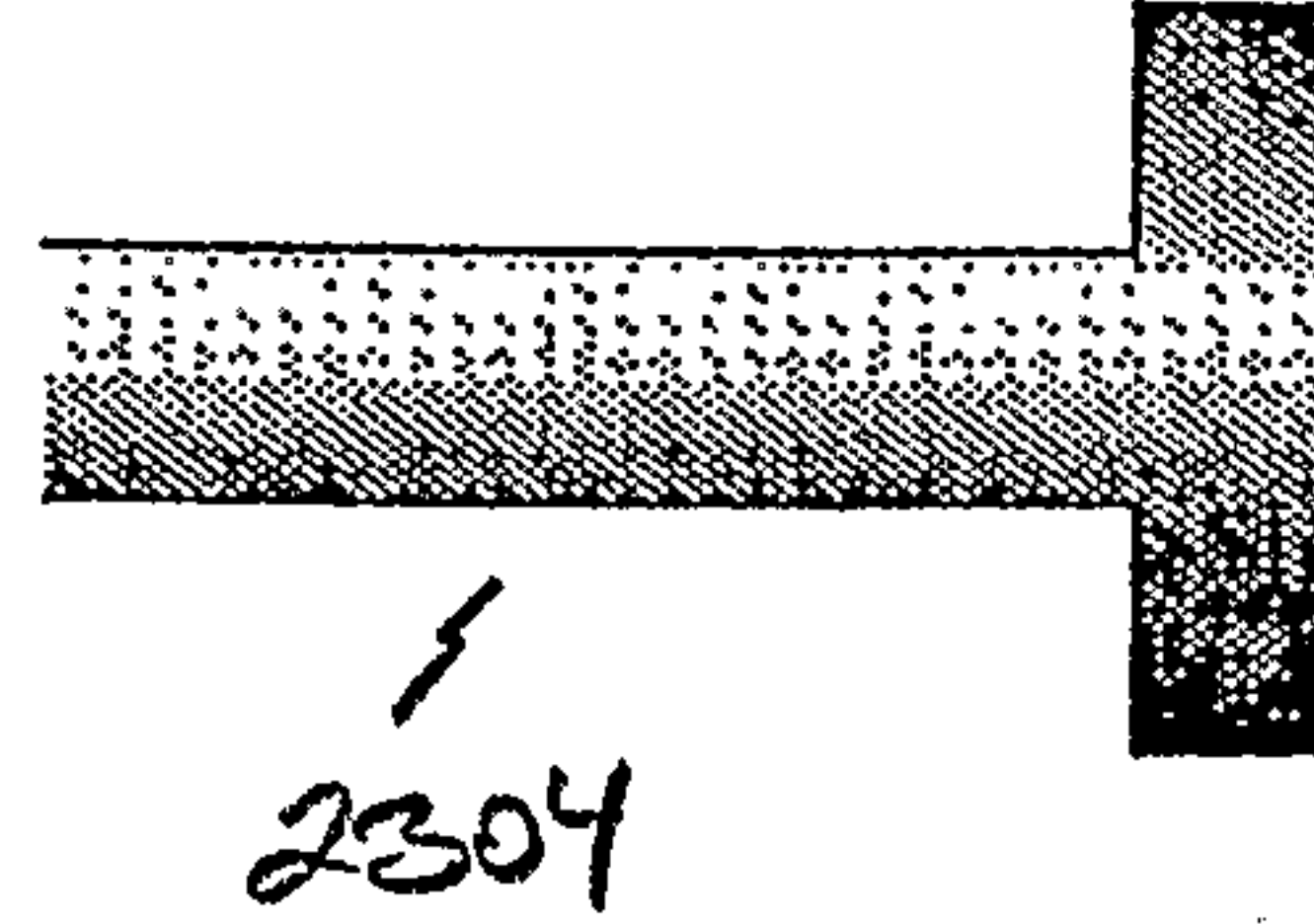
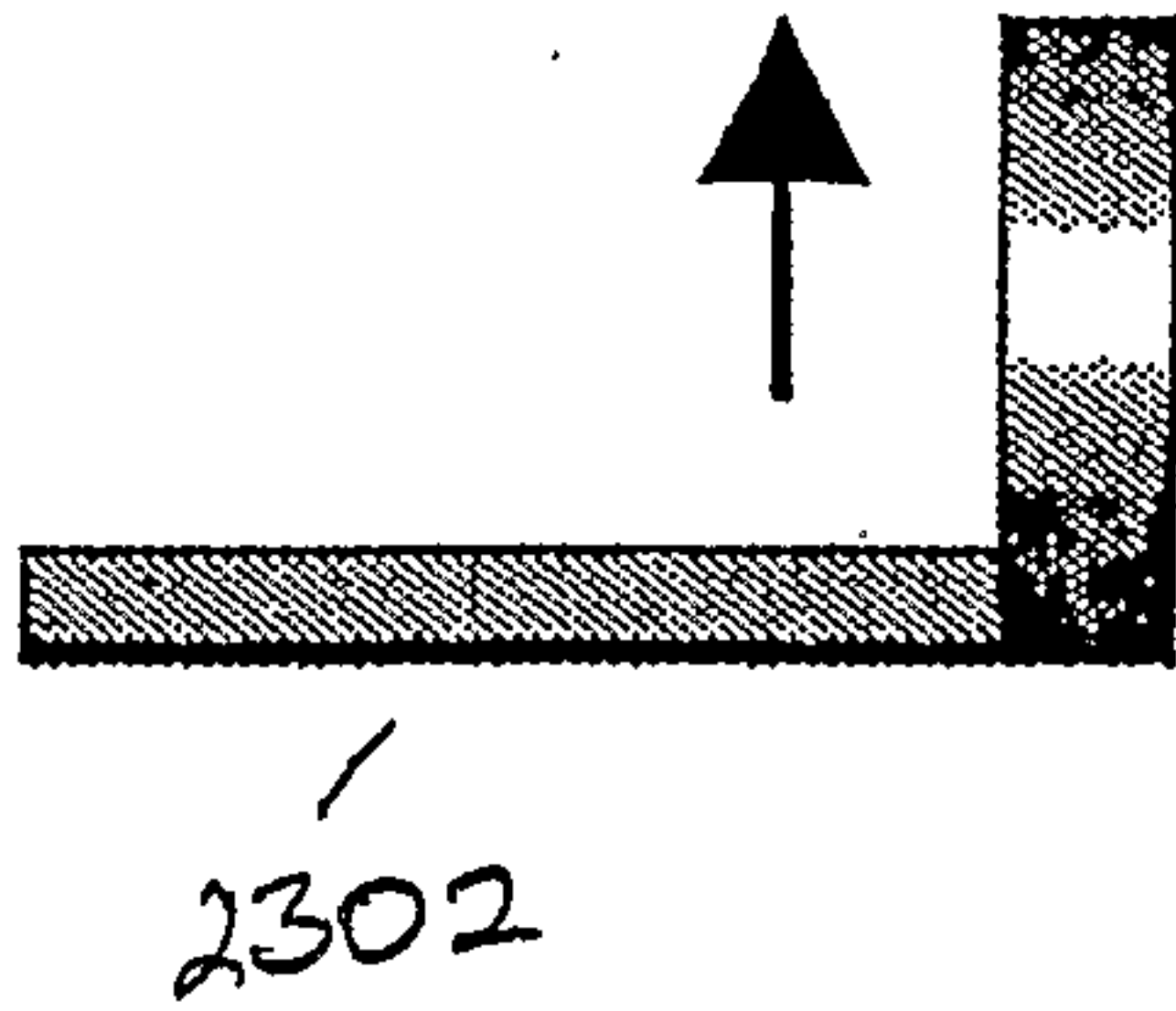


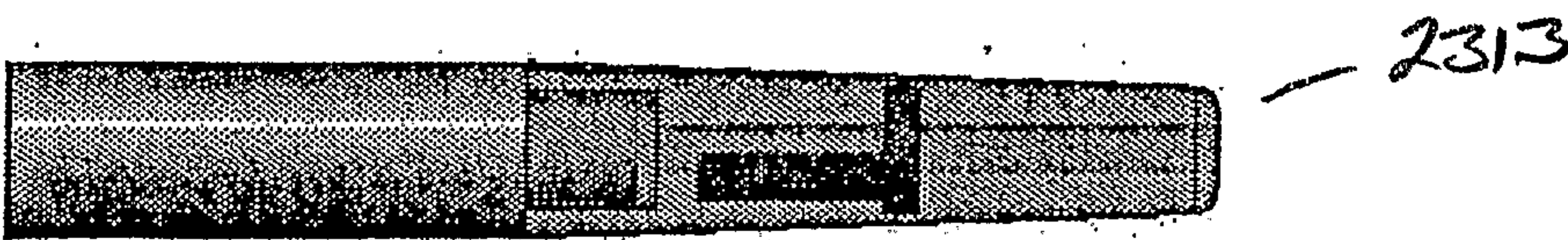
Figure 22

Use the "L" shape to locate cannula tip towards target site.

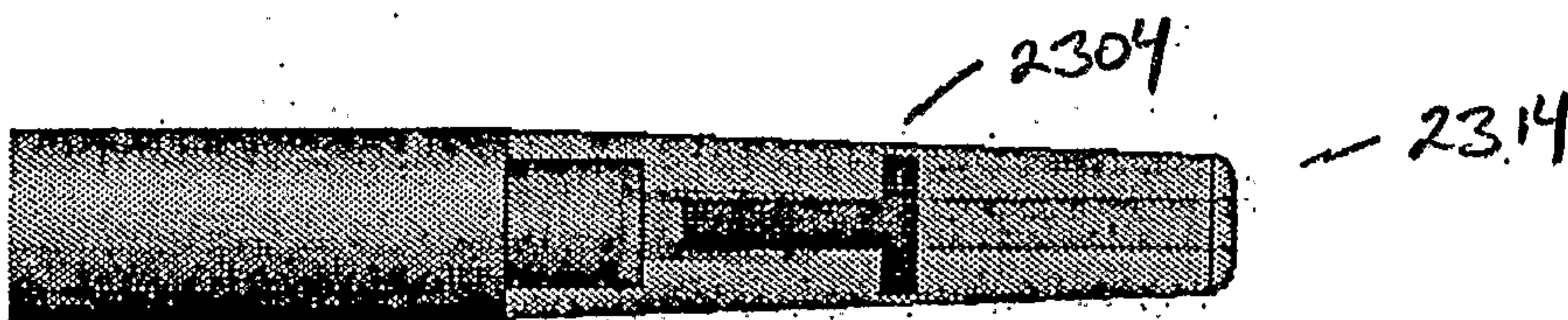
Use the "T" shape to fine tune cannula tip towards target site.



Locate needle direction in the "L" shape. (Note: extended needle in this illustration is for demonstration only. The needle should be in fully retracted position during the procedure).



Rotating catheter via the RHV for fine-tuning. (Note: Never rotate or advance catheter while cannula is deployed).



Fine tune needle direction in the "T" shape. (Note: Never rotate or advance catheter while cannula is deployed).

FIGURE 23

