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**DK/EP 2692865 T3**

**Description****BACKGROUND OF THE INVENTION**5 **Field of the Invention**

## (a) Technologies for the identification of specific functional and binding proteins

[0001] The discovery of target-specific proteins, including antibodies and fragments thereof, is of significant commercial interest, because the selection of highly selective functional proteins or binding proteins, including antibodies and fragments thereof, has a high potential for the development of new biological entities (NBEs) with novel therapeutic properties that very specifically integrate, or interfere with biological processes, and therefore are predicted to display lower side-effect profiles than conventional new chemical entities (NCEs). In that respect, particularly the development of highly target-specific, therapeutic antibodies, and antibody-based therapeutics, have paved the way to completely novel therapies with improved efficacy. As a consequence, therapeutic monoclonal antibodies represent the fastest growing segment in the development of new drugs over the last decade, and presently generate about USD 50 billion global revenues, which accounts for a significant share of the total global market of pharmaceutical drugs. Additional page 1a.

[0002] Therefore, efficient and innovative technologies, that allow the discovery of highly potent, but also well tolerated therapeutic proteins, in particular antibody-based therapeutics, are in high demand.

[0003] In order to identify a protein with a desired functionality or a specific binding property, as is the case for antibodies, it is required to generate, to functionally express and to screen large, diverse collections, or libraries of proteins, including antibodies and fragments thereof, for desired functional properties or target binding specificity. A number of technologies have been developed over the past twenty years, which allow expression of diverse protein libraries either in host cells, or on viral and phage particles and methods for their high-throughput screening and/or panning toward a desired functional property, or binding phenotype.

[0004] In Kempeni, Ann Rheum Dis 1999, 58 (suppl I) 170-2, the preliminary results of early clinical trials with fully human anti-TNF alpha monoclonal antibody D2E7 is discussed.

[0005] Furthermore, Urban et al., Nucleic Acids Research, 2005, Vol. 33, No.4 discusses the display of single chain Fv fragments expressed on the surface of retroviral particles by fusion to a retroviral envelope protein and selection of single chain Fv fragments binders by panning of retroviral particles to a desired binding protein. It is not related to the display of full-length antibodies on the surface of mammalian cells.

[0006] Standard, state-of-the-art technologies to achieve identification of target-specific binders or proteins with desired functional properties include, e.g. phage-display, retroviral display, bacterial display, yeast display and various mammalian cell display technologies, in combination with solid surface binding (panning) and/or other enrichment techniques. All of these technologies are covered by various patents and pending patent applications.

[0007] While phage and prokaryotic display systems have been established and are widely adopted in the biotech industry and in academia for the identification of target-specific binders, including antibody fragments (Hoogenboom, Nature Biotechnology 23, 1105-1116 (2005)), they suffer from a variety of limitations, including the inability to express full-length versions of larger proteins, including full-length antibodies, the lack of proper post-translational modification, the lack of proper folding by vertebrate chaperones, and, in the case of antibodies, an artificially enforced heavy and light chain combination. Therefore, in case of antibody discovery by these methods, "reformatting" into full-length antibodies and mammalian cell expression is required. Due to the above-mentioned limitations this frequently results in antibodies with unfavorable biophysical properties (e.g. low stability, tendency to aggregate, diminished affinity), limiting the therapeutic and diagnostic potential of such proteins. This, on one hand, leads to significant attrition rates in the development of lead molecules generated by these methods, and, on the other hand, requires significant effort to correct the biophysical and molecular liabilities in these proteins for further downstream drug development.

[0008] Therefore, protein and antibody discovery technologies have been developed using lower eukaryotic (e.g. yeast) and, more recently, also mammalian cell expression systems for the identification of proteins with desired properties, as these technologies allow (i) expression of larger, full-length proteins, including full-length antibodies, (ii) better or normal post-translational modification, and, (iii) in case of antibodies, proper heavy-light chain pairing (Beerli & Rader, mAbs 2, 365-378 (2010)). This, in aggregate, selects for proteins with favorable biophysical properties that have a higher potential in drug development and therapeutic use.

[0009] Although expression and screening of proteins in vertebrate cells would be most desirable, because vertebrate cells (e.g. hamster CHO, human HEK-293, or chicken DT40 cells) are preferred expression systems for the production of larger therapeutic proteins, such as antibodies, these technologies are currently also associated with a number of limitations, which has lead to a slow adoption of these technologies in academia and industry.

[0010] First, vertebrate cells are not as efficiently and stably genetically modified, as, e.g. prokaryotic or lower eukaryotic cells like yeast. Therefore, its remains a challenge to generate diverse (complex) enough vertebrate cell based proteins

libraries, from which candidates with desired properties or highest binding affinities can be identified. Second, in order to efficiently isolate proteins with desired properties, usually iterative rounds of cell enrichment are required. Vertebrate expression either by transient transfection of plasmids (Higuchi et al J. Immunol. Methods 202, 193-204(1997) ), or transient viral expression systems, like sindbis or vaccinia virus (Beerli et al. PNAS 105, 14336-14341 (2008), and WO02102885) do not allow multiple rounds of cell selection required to efficiently enrich highly specific proteins, and these methods are therefore either restricted to screening of small, pre-enriched libraries of proteins, or they do require tedious virus isolation/cell re-infection cycles.

5 [0011] In order to achieve stable expression of binding proteins and antibodies in vertebrate cells, that do allow multiple rounds of selections based on stable genotype-phenotype coupling, technologies have been developed, utilizing specific recombinases (flp/frt recombinase system, Zhou et al. mAbs 5, 508-518 (2010)), or retroviral vectors (WO2009109368). However, the flp/frt recombination is a low-efficient system for stable integration of genes into vertebrate host cell genomes and therefore, again, only applicable to small, pre-selected libraries, or the optimization of selected protein or antibody candidates.

10 [0012] In comparison to the flp/frt recombinase system, retroviral vectors allow more efficient stable genetic modification of vertebrate host cells and the generation of more complex cellular libraries. However, (i) they are restricted to only selected permissible cell lines, (ii) they represent a biosafety risk, when human cells are utilized, (iii) retroviral expression vectors are subject to unwanted mutagenesis of the library sequences due to low-fidelity reverse transcription, (iv) retroviral vectors do not allow integration of genomic expression cassettes with intact intron/exon structure, due to splicing of the retroviral genome prior to packaging of the vector into retroviral particles, (v) retroviruses are subject to uncontrollable and unfavorable homologous recombination of library sequences during packaging of the viral genomes, (vi) are subject to retroviral silencing, and (vii) require a tedious two-step packaging-cell transfection / host-cell infection procedure. All these limitations represent significant challenges and limitations, and introduce significant complexities for the utility of retroviral vector based approaches in generating high-quality/high complexity vertebrate cell libraries for efficient target-specific protein, or antibody discovery.

15 [0013] Therefore, clearly a need exists for a more efficient, more controllable and straightforward technology that allows the generation of high-quality and highly complex vertebrate cell based libraries expressing diverse libraries of proteins including antibodies and fragments thereof from which proteins with highly specific function and/or binding properties and high affinities can be isolated.

20 30 (b) Transposases/Transposition:

[0014] Transposons, or transposable elements (TEs), are genetic elements with the capability to stably integrate into host cell genomes, a process that is called transposition (Ivics et al. Mobile DNA 1, 25 (2010) . TEs were already postulated in the 1950s by Barbara McClintock in genetic studies with maize, but the first functional models for transposition have been described for bacterial TEs at the end of the 1970s (Shapiro, PNAS 76, 1933-1937 (1979).

25 [0015] Meanwhile it is clear that TEs are present in the genome of every organism, and genomic sequencing has revealed that approximately 45% of the human genome is transposon derived (International Human Genome Sequencing Consortium Nature 409: 860-921 (2001)). However, as opposed to invertebrates, where functional (or autonomous) TEs have been identified (Fig. 1a), humans and most higher vertebrates do not contain functional TEs. It has been hypothesized that evolutionary selective pressure against the mutagenic potential of TEs lead to their functional inactivation millions of years ago during evolution.

30 [0016] Autonomous TEs comprise DNA that encodes a transposase enzyme located in between two inverted terminal repeat sequences (ITRs), which are recognized by the transposase enzyme encoded in between the ITRs and which can catalyze the transposition of the TE into any double stranded DNA sequence (FIG. 1a). There are two different classes of transposons: class I, or retrotransposons, that mobilize via an RNA intermediate and a "copy-and-paste" mechanism (FIG. 2b), and class II, or DNA transposons, that mobilize via excision-integration, or a "cut-and-paste" mechanism (FIG. 2a) (Ivics et al. Nat. Methods 6, 415-422(2009)).

35 [0017] Bacterial, lower eukaryotic (e.g. yeast) and invertebrate transposons appear to be largely species specific, and cannot be used for efficient transposition of DNA in vertebrate cells. Only, after a first active transposon had been artificially reconstructed by sequence shuffling of inactive TEs from fish, which was therefore called "*Sleeping beauty*" (Ivics et al. Cell 91, 501-510 (1997)), did it become possible to successfully achieve DNA integration by transposition into vertebrate cells, including human cells. *Sleeping beauty* is a class II DNA transposon belonging to the Tc1/mariner family of transposons (Ni et al. Briefings Funct. Genomics Proteomics 7, 444-453 (2008)). In the meantime, additional functional transposons have been identified or reconstructed from different species, including Drosophila, frog and even human genomes, that all have been shown to allow DNA transposition into vertebrate and also human host cell genomes (FIG. 3). Each of these transposons, have advantages and disadvantages that are related to transposition efficiency, stability of expression, genetic payload capacity, etc.

40 [0018] To date, transposon-mediated technologies for the expression of diverse libraries of proteins, including anti-

bodies and fragments thereof, in vertebrate host cells for the isolation of target specific, functional binding proteins, including antibodies and fragments thereof, have not been disclosed in the prior art.

## BRIEF SUMMARY OF THE INVENTION

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**[0019]** The method as characterized in the claims and disclosed herein describes a novel technology offering unparalleled efficiency, flexibility, utility and speed for the discovery and optimization of polypeptides having a desired binding specificity and/or functionality, including antigen-binding molecules such as antibodies and fragments thereof, for desired functional and/or binding phenotypes. The novel method is based on transposable constructs and diverse DNA libraries cloned into transposable vectors and their transfection into host cells by concomitant transient expression of a functional transposase enzyme. This ensures an efficient, stable introduction of the transposon-based expression vectors into vertebrate host cells in one step, which can then be screened for a desired functional or binding phenotype of the expressed proteins, after which the relevant coding sequences for the expressed proteins, including antibodies and fragments thereof, can be identified by standard cloning and DNA sequencing techniques.

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**[0020]** In one embodiment, the invention is broadly directed to a method for identifying a polypeptide having a desired binding specificity or functionality, comprising:

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- (i) generating a diverse collection of polynucleotides encoding polypeptides having different binding specificities or functionalities, wherein said polynucleotides comprise a sequence coding for a polypeptide disposed between first and second inverted terminal repeat sequences that are recognized by and functional with at least one transposase enzyme;
- (ii) introducing the diverse collection of polynucleotides of (i) into host cells;
- (iii) expressing at least one transposase enzyme functional with said inverted terminal repeat sequences in said host cells so that said diverse collection of polynucleotides is integrated into the host cell genome to provide a host cell population that expresses said diverse collection of polynucleotides encoding polypeptides having different binding specificities or functionalities;
- (iv) screening said host cells to identify a host cell expressing a polypeptide having a desired binding specificity or functionality; and
- (v) isolating the polynucleotide sequence encoding said polypeptide from said host cell.

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**[0021]** In a preferred embodiment, the polynucleotides are DNA molecules. In one embodiment, the diverse collection of polynucleotides comprises a ligand-binding sequence of a receptor or a target binding sequence of a binding molecule. In a preferred embodiment, the polynucleotides comprise a sequence encoding an antigen-binding molecule, such as an antibody VH or VL domain, or an antigen-binding fragment thereof, or antibody heavy or light chains that are full-length (i.e., which include the constant region). In certain embodiments, the polynucleotides may comprise a sequence encoding both a VH and VL region, or both antibody heavy and light chains. In another embodiment, the polynucleotides comprise a sequence encoding a single-chain Fv or a Fab domain.

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**[0022]** In one embodiment, the diverse collection of polynucleotides is generated by subjecting V region gene sequences to PCR under mutagenizing conditions, for example, by PCR amplification of V region repertoires from vertebrate B cells. In another embodiment, the diverse collection of polynucleotides is generated by gene synthesis (e.g., by randomization of sequences encoding a polypeptide having known binding specificity and/or functionality). In one useful embodiment, the diverse collection of polynucleotides comprises plasmid vectors. In another useful embodiment, the diverse collection of polynucleotides comprises double-stranded DNA PCR amplicons. The plasmid vectors may comprise a sequence encoding a marker gene, such as a fluorescent marker, a cell surface marker, or a selectable marker. The marker gene sequence may be upstream or downstream of the sequence encoding the polypeptide having a binding specificity or functionality, but between the inverted terminal repeat sequences. Alternatively, the marker gene sequence may be downstream of said sequence encoding a polypeptide having binding specificity or functionality and separated by an internal ribosomal entry site.

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**[0023]** In some embodiments, the diverse collection of polynucleotides encode a plurality of antigen-binding molecules of a vertebrate, such as a mammal, e.g., a human.

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**[0024]** In one embodiment, step (ii) of the method comprises introducing into host cells polynucleotides comprising sequences encoding immunoglobulin VH or VL regions, or antigen-binding fragments thereof, and wherein said VH and VL region sequences are encoded on separate vectors. In another embodiment, step (ii) of the method of the invention comprises introducing into host cells polynucleotides comprising sequences encoding full-length immunoglobulin heavy or light chains, or antigen-binding fragments thereof, wherein said full-length heavy and light chain sequences are on separate vectors. The vectors may be introduced into the host cells simultaneously or sequentially. In another embodiment, sequences encoding VH and VL regions or full-length heavy and light chains are introduced into host cells on the same vector. In the event that the VH and VL sequences or the full-length antibody heavy and light chain sequences

are introduced into the host cells on different vectors, it is useful for the inverted terminal repeat sequences on each vector to be recognized by and functional with different transposase enzymes.

[0025] The host cells are preferably vertebrate cells, and preferably mammalian cells, such as rodent or human cells. Lymphoid cells, e.g., B cells, are particularly useful. B cells may be progenitor B cells or precursor B cells such as, for example, Abelson-Murine Leukemia virus transformed progenitor B cells or precursor B cells and early, immunoglobulin-null EBV transformed human proB and preB cells. Other useful host cells include B cell lines such as Sp2/0 cells, NSO cells, X63 cells, and Ag8653 cells, or common mammalian cell lines such as CHO cells, Per.C6 cells, BHK cells, and 293 cells.

[0026] In one embodiment of the method of the invention, the expressing step (iii) comprises introducing into said host cells an expression vector encoding a transposase enzyme that recognizes and is functional with at least one inverted terminal repeat sequence in the polynucleotides. The vector encoding the transposase enzyme may be introduced into the host cells concurrently with or prior or subsequent to the diverse collection of polynucleotides. In one embodiment, the transposase enzyme is transiently expressed in said host cell. Alternatively, the expressing step (iii) may comprise inducing an inducible expression system that is stably integrated into the host cell genome, such as, for example, a tetracycline-inducible or tamoxifen-inducible system. In a preferred embodiment, step (iii) comprises expressing in the host cell(s) a vector comprising a functional Sleeping Beauty transposase or a functional PiggyBac transposase. In one useful embodiment, step (iii) comprises expressing in said host cell a vector comprising SEQ ID NO:17. In another useful embodiment, the vector encodes SEQ ID NO:18, or a sequence with at least 95% amino acid sequence homology and having the same or similar inverted terminal repeat sequence specificity.

[0027] In one embodiment of the method of the invention, the screening step (iv) comprises magnetic activated cell sorting (MACS), fluorescence activated cell sorting (FACS), panning against molecules immobilized on a solid surface panning, selection for binding to cell-membrane associated molecules incorporated into a cellular, natural or artificially reconstituted lipid bilayer membrane, or high-throughput screening of individual cell clones in multi-well format for a desired functional or binding phenotype. In one embodiment, the screening step (iv) comprises screening to identify polypeptides having a desired target-binding specificity or functionality. In a preferred embodiment, the screening step (iv) comprises screening to identify antigen-binding molecules having a desired antigen specificity. In one useful embodiment, the screening step further comprises screening to identify antigen-binding molecules having one or more desired functional properties. The screening step (iv) may comprise multiple cell enrichment cycles with host cell expansion between individual cell enrichment cycles.

[0028] In one embodiment of the method of the invention, the step (v) of isolating the polynucleotide sequence encoding the polypeptide having a desired binding specificity or functionality comprises genomic or RT-PCR amplification or next-generation deep sequencing. In one useful embodiment, the polynucleotide sequence isolated in step (v) is subjected to affinity optimization. This can be done by subjecting the isolated polynucleotide sequence to PCR or RT-PCR under mutagenizing conditions. In another useful embodiment, the mutagenized sequence is then further subjected to steps (i)-(v) of the method of the invention. In a preferred embodiment, the polynucleotide sequence obtained in (v) comprises a sequence encoding a VH or VL region of an antibody, or an antigen-binding fragment thereof, and wherein said antibody optimization comprises introducing one or more mutations into a complementarity determining region or framework region of said VH or VL.

[0029] In one useful embodiment, the inverted terminal repeat sequences are from the PiggyBac transposon system and are recognized by and functional with the PiggyBac transposase. In one embodiment, the sequence encoding the upstream PiggyBac inverted terminal repeat sequence comprises SEQ ID NO:1. In another embodiment, the sequence encoding the downstream PiggyBac inverted terminal repeat sequence comprises SEQ ID NO:2.

[0030] In another useful embodiment, the inverted terminal repeat sequences are from the Sleeping Beauty transposon system and are recognized by and functional with the Sleeping Beauty transposase. In one embodiment, the sequence encoding the upstream Sleeping Beauty inverted terminal repeat sequence comprises SEQ ID NO:14. In another embodiment, the sequence encoding the downstream Sleeping Beauty inverted terminal repeat sequence comprises SEQ ID NO:15.

[0031] In one embodiment of the invention, the polynucleotides comprise VH or VL region sequences encoding a sequence derived from a human anti-TNF alpha antibody. In one embodiment, the human anti-TNF alpha antibody is D2E7.

[0032] In a useful embodiment, step (iii) comprises introducing into said host cell a vector comprising a sequence encoding a functional PiggyBac transposase. In one embodiment the vector comprises SEQ ID NO:11. In another embodiment, the vector encodes SEQ ID NO:12, or a sequence with at least 95% amino acid sequence homology and having the same or similar inverted terminal repeat sequence specificity.

[0033] In preferred embodiments, the inverted terminal repeat sequences are recognized by and functional with at least one transposase selected from the group consisting of: PiggyBac, Sleeping Beauty, Frog Prince, Himar1, Passport, Minos, hAT, Tol1, Tol2, Ac/Ds, PIF, Harbinger, Harbinger3-DR, and Hsmar1.

[0034] The present invention is further directed to a library of polynucleotide molecules encoding polypeptides having

different binding specificities or functionalities, comprising a plurality of polynucleotide molecules, wherein said polynucleotide molecules comprise a sequence encoding a polypeptide having a binding specificity or functionality disposed between inverted terminal repeat sequences that are recognized by and functional with at least one transposase enzyme. Preferably the polynucleotides are DNA molecules and comprise a ligand-binding sequence of a receptor or a target-binding sequence of a binding molecule. In a particularly preferred embodiment, the library comprises polynucleotides, wherein each polynucleotide comprises a sequence encoding an antigen-binding sequence of an antibody. In one embodiment, the library comprises polynucleotides encoding a VH or VL region of an antibody or an antigen-binding fragment thereof. Alternatively, the polynucleotides may encode a VH region and a VL region. In a preferred embodiment, the polynucleotides of the library comprise a sequence encoding a full-length antibody heavy or light chain (i.e., including the constant region) or an antigen-binding fragment thereof. Alternatively, the polynucleotides may encode both a full-length immunoglobulin heavy and light chain. In other embodiments, the polynucleotides of the library comprise a sequence encoding a single-chain Fv or a Fab domain. In preferred embodiments, the polynucleotides of the library are in the form of plasmids or double stranded DNA PCR amplicons. In certain embodiments, the plasmids of the library comprise a marker gene. In another embodiment, the plasmids comprise a sequence encoding a transposase enzyme that recognizes and is functional with the inverted terminal repeat sequences. In one embodiment, the library of the invention comprises polynucleotides that encode the full-length immunoglobulin heavy chain including the natural intron/exon structure of an antibody heavy chain. The full-length immunoglobulin heavy chain may comprise the endogenous membrane anchor domain.

[0035] The present invention is also directed to a method for generating a library of transposable polynucleotides encoding polypeptides having different binding specificities or functionality, comprising (i) generating a diverse collection of polynucleotides comprising sequences encoding polypeptides having different binding specificities or functionalities, wherein said polynucleotides comprise a sequence encoding polypeptide having a binding specificity or functionality disposed between inverted terminal repeat sequences that are recognized by and functional with a least one transposase enzyme.

[0036] The present invention is also directed to a vector comprising a sequence encoding a VH or VL region of an antibody, or antigen-binding portion thereof, disposed between inverted terminal repeat sequences that are recognized by and functional with at least one transposase enzyme. In certain embodiments, the vector encodes a full-length heavy or light chain of an immunoglobulin. Preferably, the sequence encoding the VH or VL or the heavy or light chain is a randomized sequence generated by, for example, PCR amplification under mutagenizing conditions or gene synthesis. In one embodiment, the vector comprises inverted terminal repeat sequences that are recognized by and functional with the PiggyBac transposase. In another embodiment, the inverted terminal repeat sequences are recognized by and functional with the Sleeping Beauty transposase. In one embodiment, the vector comprises a VH or VL region sequence derived from an anti-TNF alpha antibody such as, for example, D2E7. In certain embodiments, the vector comprises at least one sequence selected from the group consisting of: SEQ ID NO:5, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:13, SEQ ID NO:16, SEQ ID NO:17, and SEQ ID NO:19.

[0037] The present invention is also directed to a host cell comprising a vector of the invention as described above. In a preferred embodiment, the host cell further comprises an expression vector comprising a sequence encoding a transposase that recognizes and is functional with at least one inverted terminal repeat sequence in the vector encoding said VH or VL region sequence.

[0038] The present invention is still further directed to antigen-binding molecules, e.g., antibodies, produced by a method comprising claim 1.

[0039] The present invention is also directed to a method for generating a population of host cells capable of expressing polypeptides having different binding specificities or functionalities, comprising:

- 45 (i) generating a diverse collection of polynucleotides comprising sequences encoding polypeptides having different binding specificities or functionalities, wherein said polynucleotides comprise a sequence encoding a polypeptide having a binding specificity or functionality disposed between inverted terminal repeat sequences that are recognized by and functional with a least one transposase enzyme; and
- 50 (ii) introducing said diverse collection of polynucleotides into host cells.

#### BRIEF DESCRIPTION OF THE DRAWINGS/FIGURES

#### [0040]

55 FIG. 1: a.) This drawing depicts the configuration of an autonomous transposable element (TE), which can transpose or "jump" into any target DNA sequence. The key components of a TE are an active transposase enzyme that recognizes the inverted terminal repeats (ITRs) flanking the transposase enzyme itself up- and downstream of its sequence. TEs catalyze either the copying or the excision of the TE, and the integration in unrelated target DNA

5 sequences. b.) This drawing depicts the configuration of a transposon vector system, in which the expression of an active transposase enzyme is effected by an expression vector that is not coupled to the TE itself. Instead, the TE may contain any sequence(s), or gene(s) of interest that is/are cloned in between the up- and downstream ITRs. Integration of the TE containing any sequence(s), or gene(s) of interest (e.g. a DNA library encoding a library of proteins) may integrate into unrelated target DNA sequences, if the transposase enzyme expression is provided in *trans*, e.g. by a separate transposase expression construct, as depicted here.

10 FIG. 2: a) This drawing depicts the two different ways how TEs can "jump" or transpose into unrelated target DNA. For group II transposons the transposase enzyme in a first step recognizes the ITRs of the transposable element and catalyzes the excision of the TE from DNA. In a second step, the excised TE is inserted into unrelated target DNA sequence, which is also catalyzed by the transposase enzyme. This results in a "cut-and-paste" mechanism of transposition. For group I transposons (shown in b.) the coding information of the TE is first replicated (e.g. transcribed and reverse transcribed, in the case of retrotransposons) and the replicated TE then integrates into unrelated target DNA sequence, which is catalyzed by the transposase enzyme. This results in a "copy-and-paste" mechanism of transposition.

15 FIG. 3: This figure provides an overview of active transposase enzymes that have been identified and/or reconstructed from dormant, inactive TEs, and that have been shown to be able to confer transposition in various vertebrate and also human cells, as provided in the table. The table has been adapted from Table I of publication Ni et al. *Briefings Functional Genomics Proteomics* 7, 444-453 (2008).

20 FIG. 4: This figure outlines the principle of the method disclosed herein, for the isolation of coding information for proteins, including antibodies and fragments thereof, with a desired function, e.g. the binding to a target of interest, as depicted here. The gene(s) of interest, e.g. a diverse transposable DNA library encoding proteins, including antibody polypeptide chains or fragments thereof, that is cloned in between inverted terminal repeats (ITRs) of a transposable construct is introduced into a vertebrate host cell together with an expression vector for an active transposase enzyme (see top of the drawing). The expression of the transposon enzyme in said host cells *in trans* and the presence of the gene(s) of interest cloned in between ITRs that can be recognized by the transposase enzyme allows the stable integration of the ITR-flanked gene(s) of interest into the genome of the host cells, which can then stably express the protein(s) of interest encoded by the genes of interest. The cellular library expressing the protein(s) of interest can then be screened for a desired functionality of the expressed proteins, e.g., but not limited to the binding to a target protein of interest, as depicted here. By means of cell separation techniques known in the art, e.g. MACS or FACS, the cells expressing the protein(s) of interest with the desired phenotype and which therefore contain the corresponding genotype, can be isolated and the coding information for the gene(s) of interest can be retrieved from the isolated cells by cloning techniques known in the art, e.g. but not limited to genomic PCR cloning, as depicted here.

25 FIGs. 5a) and 5b): This drawing outlines the cloning strategy for the generation of a transposable human immunoglobulin (Ig) kappa light chain (LC) expression vector, as described in Example 1. FIG. 5 a.) depicts the cloning strategy for the insertion of 5'- and 3'-ITRs from the *PiggyBac* transposon into the mammalian expression vector pIRES-EGFP (Invitrogen, Carlsbad, CA, USA), which already contains the strong mammalian cell promoter element pCMV(IE) (immediate early promoter of CMV), and intron/polyA signals for strong mammalian host cell expression. In addition, downstream of the Clal, EcoRV, Notl, EcoRI containing multiple cloning site, into which gene(s) of interest can be cloned, pIRES-EGFP contains an internal ribosomal entry site (IRES) with a downstream ORF of enhanced green fluorescent protein (EGFP), which effects the coupling of expression of gene(s) of interest cloned upstream of the IRES. Bacterial functional elements (ampicillin resistance gene, amp<sup>R</sup>) and a bacterial origin of replication (Col E1) for amplification and selection of the plasmid in *E. coli* are depicted as well. The resulting *PiggyBac* ITRs containing plasmid is designated pIRES-EGFP-T1T2. FIG 5b) then depicts the insertion of a gene synthesized human Ig kappa LC into the unique EcoRV restriction enzyme site of pIRES-EGFP-T1-T2, which positions the human Ig kappa LC upstream of the IRES-EGFP cassette, and thereby couples the expression of the human Ig kappa LC to EGFP marker gene expression. The insertion of the human Ig kappa LC results in transposable human Ig kappa LC expression vector pIRES-EGFP-T1T2-IgL. The drawings show selected unique restriction enzyme sites in the plasmids, as well as selected duplicated sites resulting from cloning steps.

30 FIG. 6: This drawing outlines the cloning of a transposable human immunoglobulin (Ig) gamma 1 heavy chain (HC) expression vector, which can be generated by exchange of the human Ig kappa LC open reading frame (ORF) against the ORF for a human Ig gamma 1 HC ORF. The design of the final Ig gamma 1 HC ORF is similar, also with regard to the engineering of a unique Eco47III restriction enzyme site separating the variable (V) from the constant (C) coding regions, which allows the exchange of a single antibody V coding region against a diverse library of antibody V coding regions, as described in Example 3.

35 FIG. 7: This drawing depicts the cloning of a mammalian *PiggyBac* transposase enzyme expression vector, as described in the Example 4, using pCDNA3.1(+)-hygro as the backbone of the mammalian expression vector, into which the gene synthesized ORF from *PiggyBac* transposase is cloned into the unique EcoRV restriction enzyme

site of pCDNA3.1(+) hygro, resulting in PiggyBac transposon expressin vector pCDNA3.1(+) hygro-PB. expression vector pCDNA3.1(+) hygro-PB. Also in this drawing the relative position of other mammalian functional elements (CMV-IE promoter, BGH-polyA signal, SV40-polyA segment, hygromycinB ORF) and bacterial functional elements (ampicillin resistance gene, ampR, origin of replication, ColE1), as well as selected relevant restriction enzyme recognition sites are shown.

Fig. 8: This drawing depicts the cloning of a Sleeping Beauty transposable human immunoglobulin kappa light chain (Ig-kappa LC) expression vector, as described in Example 5. The cloning can be performed by sequentially replacing the PiggyBac 5' and 3' ITRs with Sleeping Beauty 5' and 3'ITRs in construct pIREs-EGFP-T1T2-IgL. Also in this drawing the relative position of other mammalian functional elements (CMV-IE promoter, BGH-polyA signal, SV40-polyA segment, hygromycinB ORF) and bacterial functional elements (ampicillin resistance gene, ampR, origin of replication, Co1E1), as well as selected relevant restriction enzyme recognition sites are shown.

[0052] Fig. 9: This drawing depicts the cloning of a mammalian Sleeping Beauty transposase enzyme expression vector, as described in the Example 6, using pCDNA3.1(+) hygro as the backbone of the mammalian expression vector, into which the gene synthesized ORF from Sleeping Beauty transposase is cloned into the unique EcoRV restriction enzyme site of pCDNA3.1(+) hygro, resulting in Sleeping Beauty transposon expression vector pCDNA3.1(+) hygro-SB. Also in this drawing the relative position of other mammalian functional elements (CMV-IE promoter, BGH-polyA signal, SV40-polyA segment, hygromycinB ORF) and bacterial functional elements (ampicillin resistance gene, ampR, origin of replication, ColE1), as well as selected relevant restriction enzyme recognition sites are shown.

## DETAILED DESCRIPTION OF THE INVENTION

### *Definitions*

[0041] As used herein, "diverse collection" means a plurality of variants or mutants of particular functional or binding proteins exhibiting differences in the encoding nucleotide sequences or in the primary amino acid sequences, which define different functionalities or binding properties.

[0042] As used herein, "library" means a plurality of polynucleotides encoding polypeptides having different binding specificities and/or functionalities. In certain embodiments, the library may comprise polynucleotides encoding at least  $10^2$ , at least  $10^3$ , at least  $10^4$ , at least  $10^5$ , at least  $10^6$ , at least  $10^7$ , at least  $10^8$ , or at least  $10^9$  unique polypeptides, such as, for example, full-length antibody heavy or light chains or VH or VL domains.

[0043] As used herein, "inverted terminal repeat sequence" or "ITR" means a sequence identified at the 5' or 3' termini of transposable elements that are recognized by transposases and which mediate the transposition of the ITRs including intervening coding information from one DNA construct or locus to another DNA construct or locus.

[0044] As used herein, "transposase" means an enzyme that has the capacity to recognize and to bind to ITRs and to mediate the mobilization of a transposable element from one target DNA sequence to another target DNA sequence.

[0045] As used herein, "antigen binding molecule" refers in its broadest sense to a molecule that specifically binds an antigenic determinant. A non-limiting example of an antigen binding molecule is an antibody or fragment thereof that retains antigen-specific binding. By "specifically binds" is meant that the binding is selective for the antigen and can be discriminated from unwanted or nonspecific interactions.

[0046] As used herein, the term "antibody" is intended to include whole antibody molecules, including monoclonal, polyclonal and multispecific (e.g., bispecific) antibodies, as well as antibody fragments having an Fc region and retaining binding specificity, and fusion proteins that include a region equivalent to the Fc region of an immunoglobulin and that retain binding specificity. Also encompassed are antibody fragments that retain binding specificity including, but not limited to, VH fragments, VL fragments, Fab fragments,  $F(ab')_2$  fragments, scFv fragments, Fv fragments, minibodies, diabodies, triabodies, and tetrabodies (see, e.g., Hudson and Souriau, Nature Med. 9: 129-134 (2003)).

[0047] An embodiment of the invention disclosed herein is a method as characterized in the claims for the identification of specific functional or binding polypeptides, including, but not limited to antibody chains or fragments thereof (FIG. 4), which comprises:

- i. cloning of diverse transposable DNA libraries encoding proteins, including antibody polypeptide chains or fragments thereof, in between inverted terminal repeats (ITRs) derived from transposable elements and recognizable by and functional with at least one transposase enzyme,
- ii. introduction of one or more diverse transposable DNA libraries of step (i) into vertebrate host cells by standard methods known in the art,
- iii. providing temporary expression of at least one functional transposase enzyme in said vertebrate host cells in

trans, such that said one or more diverse transposable DNA libraries are stably integrated into the vertebrate host cell genomes, thereby providing a vertebrate host cell population that then stably expresses diverse libraries of proteins, including antibody chains or fragments thereof,

- 5        iv. screening of said diverse cellular libraries, stably expressing proteins, including antibodies or fragments thereof, for a desired functional or binding phenotype by methods known in the art,
- 10      v. optionally, including iterative enrichment cycles with the stably genetically modified vertebrate host cells for a desired binding or functional phenotype, and
- 15      vi. isolation of the corresponding genes from the enriched host cells encoding the desired binding or functional phenotype by standard cloning methods, known in the art, for instance, but not limited to, PCR (polymerase chain reaction), using primers specific for the sequences contained in the one or more transposed DNA library constructs.

15      **[0048]** A preferred embodiment of step (i) is to generate diverse transposable DNA libraries either by gene synthesis, or by polymerase chain reaction (PCR) using appropriate primers for the amplification of diverse protein coding regions, and DNA templates comprising a diversity of binding proteins, including antibodies, or fragments thereof, by methods known in the art.

20      **[0049]** For the generation of diverse antibody libraries, a diverse collection of antibody heavy and light chain sequences may be generated by standard gene synthesis in which the V region coding sequences may be randomized at certain positions, e.g. but not limited to, any or all of the complementarity determining regions (CDRs) of the antibody heavy or light chain V-regions. The diversity can be restricted to individual CDRs of the V-regions, or to a particular or several framework positions, and/or to particular positions in one or more of the CDR regions. The V regions with designed variations, as described above, can be synthesized as a fragment encoding entire antibody heavy or light chains that are flanked by inverted terminal repeats functional for at least one desired transposase enzyme. Preferably, the DNA library containing diverse variable domains encoding V regions for antibody heavy or light chains is generated, and flanked by appropriate cloning sites, including but not limited to restriction enzyme recognition sites, that are compatible with cloning sites in antibody heavy or light chain expression vectors. Useful transposon expression systems for use in the methods of the invention include, for example, the PiggyBac transposon system as described, for example, in US Pat. Nos. 6,218,185; 6,551,825; 6,962,810; 7,105,343; and 7,932,088 and the Sleeping Beauty transposon system as described in US Pat. Nos. 6,489,458; 7,148,203; 7,160,682; US 2011 117072; US 2004 077572; and US 2006 252140.

30      **[0050]** Diverse antibody heavy and light chain libraries may also be obtained from B cell populations isolated from desired vertebrate species, preferably humans, and preferably from cellular compartments containing B cells, e.g., but not limited to peripheral or cord blood, and lymphoid organs like bone marrow, spleen, tonsils and lymph-node tissues. 35      In this case, diverse antibody V region sequences for antibody heavy and light chains can be isolated by RT-PCR or by genomic PCR using antibody heavy and light chain specific degenerate PCR primer pairs, that can amplify the majority of V-region families by providing upstream primers that bind to homologous sequences upstream of, or within leader sequences, upstream of or within V-region frameworks, and by providing downstream primers that bind in regions of homology within or downstream of the J joining gene segment of variable domain coding regions, or within or downstream 40      of the coding regions of the constant regions of antibody heavy or light chains.

45      **[0051]** The PCR primer sets utilized for the amplification of diverse variable coding regions may be flanked by appropriate cloning sites, e.g. but not limited to restriction enzyme recognition sites, that are compatible with cloning sites in antibody heavy or light chain expression vectors.

50      **[0052]** The transposable DNA libraries of step (i) encoding diverse proteins, including antibodies and antibody fragments thereof, can be provided in the form of plasmid libraries, in which the gene-synthesized or the PCR amplified transposable DNA libraries are cloned using appropriate cloning sites, as mentioned above. Alternatively, the transposable DNA libraries encoding diverse libraries of binding proteins, such as antibodies and fragments thereof, can be provided in form of linear, double-stranded DNA constructs, directly as a result of DNA synthesis, or as a result of PCR amplification. The latter approach of providing the transposable DNA libraries as linear double-stranded DNA PCR amplicons, that have not been cloned into expression vectors or plasmids (in comparison to all other vertebrate cell expression systems) has the advantage that the maximum molecular complexity of the transposable DNA libraries is maintained and not compromised by a limited cloning or ligation efficiency into an expression vector. In contrast, cloning by ligation, or otherwise, into plasmid expression or shuttle vectors is a necessary intermediate for all other plasmid-based or viral vector based vertebrate cell expression systems.

55      **[0053]** However, the use of plasmid-based transposon expression vectors containing the diverse transposable DNA libraries encoding diverse binding proteins, including antibodies and antibody fragments thereof, has the advantage that these expression vectors can be engineered to contain additional functional elements, that allow the screening, or, alternatively, the selection for stably transposed vertebrate host cells for the stable integration of the transposon expres-

sion vector in transposed vertebrate host cells.

[0054] This is achieved by providing in operable linkage to the diverse transposable DNA libraries, i.e. cloned into the transposon expression vectors in *cis*, expression cassettes for marker genes including., but not limited to, fluorescent marker proteins (e.g. green, yellow, red, or blue fluorescent proteins, and enhanced versions thereof, as known in the art), or expression cassettes for cell surface markers including, but not limited to, CD markers, against which specific diagnostic antibodies or other diagnostic tools are available.

[0055] Alternatively, expression cassettes for selectable markers, that allow selection of transposed vertebrate host cells for antibiotic resistance, including, but not limited to, puromycin, hygromycinB, bleomycin, neomycin resistance, can be provided in operable linkage to the diverse transposable DNA libraries, i.e. cloned into the transposon expression vectors in *cis*.

[0056] The operable linkage can be achieved by cloning of said expression cassettes for marker genes or antibiotic resistance markers, either up- or downstream of the coding regions comprising said diverse transposable DNA libraries, but within the inverted terminal repeats of the transposon vector.

[0057] Alternatively, the operable linkage can be achieved by cloning of the coding regions for said marker or antibiotic resistance genes downstream of the coding regions comprising said diverse transposable DNA libraries, but separated by internal ribosomal entry site (IRES) sequences, that ensure transcriptional coupling of the expression of said diverse transposable DNA libraries with said marker or antibiotic resistance genes, and thereby allowing the screening for or selection of stably transposed vertebrate host cells.

[0058] In step (ii) of the method disclosed herein, said diverse transposable DNA libraries encoding diverse libraries of proteins, including antibodies and fragments thereof, are introduced into desired vertebrate host cells by methods known in the art to efficiently transfer DNA across vertebrate cell membranes, including., but not limited to, DNA-transfection using liposomes, Calcium phosphate, DEAE-dextran, polyethylenimide (PEI) magnetic particles, or by protoplast fusion, mechanical transfection, including physical, or ballistic methods (gene gun), or by nucleofection. Any of the above-mentioned methods and other appropriate methods to transfer DNA into vertebrate host cells may be used individually, or in combination for step (ii) of the method disclosed herein.

[0059] In the case of dimeric proteins, including, but not limited to, antibodies and fragments thereof, it is a useful embodiment of the method disclosed herein to introduce diverse transposable DNA libraries and/or transposon vectors for antibody heavy or light chains contained in separate transposable vectors, which can independently be introduced into the vertebrate host cells. This either allows the sequential introduction of diverse transposable DNA libraries for antibody heavy or light chains into said cells, or their simultaneous introduction of diverse transposable DNA libraries for antibody heavy or light chains, which, in either case, allows the random shuffling of any antibody heavy with any antibody light chain encoded by the at least two separate diverse transposable DNA libraries.

[0060] Another useful embodiment of the previous embodiment is to utilize separate transposon vectors and/or diverse DNA transposable libraries for antibody heavy and light chains, where said constructs or libraries are contained on transposable vectors recognized by different transposase enzymes (Fig. 3). This allows the independent transposition of antibody heavy and antibody light chain constructs without interference between the two different transposase enzymes, as one transposable vector is only recognized and transposed by its specific transposase enzyme. In case of sequential transposition of transposable vectors or DNA libraries encoding antibody heavy or light chains, the advantage of utilizing different transposase enzymes with different ITR sequences is, that upon the second transposition event, the first already stably transposed construct is not again mobilized for further transposition.

[0061] This embodiment also allows the discovery of antibodies by the method of guided selection (Guo-Qiang et al. Methods Mol. Biol. 562, 133-142 (2009)). Guided selection can e.g. be used for the conversion of any non-human antibody specific for a desired target/epitope specificity and with a desired functionality into a fully human antibody, where the same target/epitope specificity and functionality is preserved. The principle of guided selection entails the expression of a single antibody chain (heavy or light chain) of a reference (the "guiding") antibody, in combination with a diverse library of the complementary antibody chains (i.e. light, or heavy chain, respectively), and screening of these heavy-light chain combinations for the desired functional or binding phenotype. This way, the first antibody chain, "guides" the selection of one or more complementary antibody chains from the diverse library for the desired functional or binding phenotype. Once the one or more novel complementary antibody chains are isolated, they can be cloned in expression vectors and again be used to "guide" the selection of the second, complementary antibody chain from a diverse antibody chain library. The end-result of this two-step process is that both original antibody heavy and light chains of a reference antibody are replaced by unrelated and novel antibody chain sequences from the diverse libraries, but where the novel antibody heavy-light chain combination exhibits the same, or similar functional or binding properties of the original reference antibody. Therefore, this method requires the ability to independently express antibody heavy and light chain constructs or libraries in the vertebrate host cells, which can be achieved by the preferred embodiment to provide antibody heavy and light chain expression cassettes in different transposable vector systems, recognized by different transposon enzymes.

[0062] However, diverse transposable DNA libraries can also be constructed in a way, that the coding regions for

multimeric proteins, including antibodies and fragments thereof, are contained in the same transposon vector, i.e. where the expression of the at least two different subunits of a multimeric protein, for example VH and VL regions or full-length heavy and light chains, is operably linked by cloning of the respective expression cassettes or coding regions into the same transposable vector.

5 [0063] Useful vertebrate host cells for the introduction of transposable constructs and/or transposable DNA libraries of step (ii) are cells from vertebrate species that can be or that are immortalized and that can be cultured in appropriate cell culture media and under conditions known in the art. These include, but are not limited to, cells from e.g. frogs, fish, avians, but preferably from mammalian species, including, but not limited to, cells from rodents, ruminants, non-human primate species and humans, with cells from rodent or human origin being preferred.

10 [0064] Useful cell types from the above-mentioned species include, but are not limited to cells of the lymphoid lineage, which can be cultured in suspension and at high densities, with B-lineage derived cells being preferred, as they endogenously express all the required proteins, factors, chaperones, and post-translational enzymes for optimal expression of many proteins, in particular of antibodies, or antibody-based proteins. Of B-lineage derived vertebrate cells, those are preferred that represent early differentiation stages, and are known as progenitor (pro) or precursor (pre) B cells, because said pro- or preB cells in most cases do not express endogenous antibody chains that could interfere with exogenous or heterologous antibody chain expression that are part of the method disclosed herein.

15 [0065] Useful pro- and pre- B lineage cells from rodent origin are Abelson-Murine Leukemia virus (A-MuLV) transformed proB and preB cells (Alt et al. Cell 27, 381-390(1981)) that express all necessary components for antibody expression and also for their proper surface deposition, including the B cell receptor components Ig-alpha (CD79a, or mb-1), and Ig-beta (CD79b, or B-29) (Hombach et al. Nature 343, 760-762 (1990)), but as mentioned above, mostly lack the expression of endogenous antibody or immunoglobulin chains. Here, A-MuLV transformed pro- and preB cells are preferred that are derived from mouse mutants, including, but not limited to, mouse mutants defective in recombination activating gene-1 (RAG-1), or recombination activating gene-2 (RAG-2), or animals carrying other mutations in genes required for V(D)J recombination, e.g. XRCC4, DNA-ligase IV, Ku70, or Ku80, Artemis, DNA-dependent protein kinase, catalytic subunit (DNA-PK<sub>cs</sub>), and thus lack the ability to normally express of endogenous antibody polypeptides.

20 [0066] Additional useful types of progenitor (pro) and precursor (pre) B lineage cells are early, immunoglobulin-null (Ig-null) EBV transformed human proB and preB cells (Kubagawa et al. PNAS 85, 875-879(1988)) that also express all the required factors for expression, post-translational modification and surface expression of exogenous antibodies (including CD79a and CD79b).

25 [0067] Other host cells of the B lineage can be used, that represent plasma cell differentiation stages of the B cell lineage, preferably, but not limited to Ig-null myeloma cell lines, like Sp2/0, NSO, X63, Ag8653, and other myeloma and plasmacytoma cells, known in the art. Optionally, these cell lines may be stably transfected or stably genetically modified by other means than transfection, in order to over-express B cell receptor components Ig-alpha (CD79a, or mb-1), and Ig-beta (CD79b, or B-29), in case optimal surface deposition of exogenously expressed antibodies is desired.

30 [0068] Other, non-lymphoid mammalian cells lines, including but not limited to, industry-standard antibody expression host cells, including, but not limited to, CHO cells, Per.C6 cells, BHK cells and 293 cells may be used as host cells for the method disclosed herein, and each of these cells may optionally also be stably transfected or stably genetically modified to over-express B cell receptor components Ig-alpha (CD79a, or mb-1), and Ig-beta (CD79b, or B-29), in case optimal surface deposition of exogenously expressed antibodies is desired.

35 [0069] Essentially, any vertebrate host cell, which is transfectable, can be used for the method disclosed herein, which represents a major advantage in comparison to any viral expression systems, such as., but not limited to vaccinia virus, retroviral, adenoviral, or sindbis virus expression systems, because the method disclosed herein exhibits no host cell restriction due to virus tropism for certain species or cell types, and furthermore can be used with all vertebrate cells, including human cells, at the lowest biosafety level, adding to its general utility.

40 [0070] Step (iii) of the method disclosed herein results in the stable genetic modification of desired vertebrate host cells with the transfected transposable constructs of step (ii) by temporary, or transient expression of a functional transposase enzyme, such that a stable population of vertebrate host cells is generated that expresses diverse libraries of proteins encoded by said constructs.

45 [0071] A useful embodiment of step (iii) is to transiently introduce into the host cells, preferably by co-transfection, as described above, a vertebrate expression vector encoding a functional transposase enzyme together with said at least one diverse transposable DNA library. It is to be understood that transient co-transfection or co-integration of a transposase expression vector can either be performed simultaneously, or shortly before or after the transfer of the transposable constructs and/or diverse transposable DNA libraries into the vertebrate host cells, such that the transiently expressed transposase can optimally use the transiently introduced transposable vectors of step (ii) for the integration of the transposable DNA library into the vertebrate host cell genome.

[0072] Another useful embodiment of step (iii) is to effect the stable integration of the introduced transposable vectors and/or transposable DNA libraries of step (ii) by transiently expressing a functional transposase enzyme by means of an inducible expression system known in the art, that is already stably integrated into the vertebrate host cell genome.

Such inducible and transient expression of a functional transposase may be achieved by e.g., tetracycline inducible (tet-on/tet-off) or tamoxifen-inducible promoter systems known in the art. In this case, only the one or more transposable vector or DNA library needs to be introduced into the host cell genome, and the stable transposition of the constructs and the stable expression of the proteins encoded by the one or more transposable vector or DNA library is effected by the transiently switched on expression of the functional transposase enzyme in the host cells.

5 [0073] Step (iv) of the method disclosed herein effects the isolation of transposed vertebrate host cells expressing proteins with a desired functionality or binding phenotype.

[0074] A preferred embodiment of step (iv) is to screen for and to isolate the transposed host cells of step (iii) expressing desired proteins, including antibodies and fragments thereof, with target-binding assays and by means of standard cell separation techniques, like magnetic activated cell sorting (MACS) or high-speed fluorescence activated cell sorting (FACS) known in the art. Especially, in a first enrichment step of a specific population of transposed vertebrate host cells, where large number of cells need to be processed, it is preferred to isolate target specific cells from a large number of non-specific cells by MACS-based techniques.

10 [0075] Particularly, for additional and iterative cell enrichment cycles, FACS enrichment is preferred, as potentially fewer numbers of cells need to be processed, and because multi channel flow cytometry allows the simultaneous enrichment of functionalities, including., but not limited to, binding to a specific target of more than one species, or the specific screening for particular epitopes using epitope-specific competing antibodies in the FACS screen.

15 [0076] If proteins, including antibodies and fragments thereof, are to be discovered that interact with soluble binding partners, these binding partners are preferably labeled with specific labels or tags, such as but not limited to biotin, myc, or HA-tags known in the art, that can be detected by secondary reagents, e.g. but not limited to, streptavidin or antibodies, that themselves are labeled magnetically (for MACS based cell enrichment) or with fluorochromes (for FACS based cell enrichment), so that the cell separation techniques can be applied.

20 [0077] If proteins, including antibodies and fragments thereof are to be discovered against membrane bound proteins, which cannot easily be expressed as soluble proteins, like e.g. but not limited to, tetraspannins, 7-transmembrane spanners (like G-coupled protein coupled receptors), or ion-channels, these may be expressed in viral particles, or overexpressed in specific cell lines, which are then used for labeling or panning methods known in the art, which can enrich the vertebrate host cells expressing the proteins from the transposed constructs, including antibodies and fragments thereof.

25 [0078] Due to the stable genotype-phenotype coupling in the stably transposed vertebrate host cell population, a useful embodiment of step (v) is to repeat cell enrichment cycles for a desired functional or binding phenotype, until a distinct population of cells is obtained that is associated with a desired functional or binding phenotype. Optionally, individual cell clones can be isolated e.g., but not limited to, by single-cell sorting using flow cytometry technology, or by limiting dilution, in order to recover the transposed DNA information from individual cell clones that are coupled to a particular, desired functional or binding phenotype.

30 [0079] For the identification of functional target-specific antibodies it is often favorable to not only screen and to select for a particular binding phenotype, but to additionally screen for additional functional properties of target specific antibodies, in particular antagonistic or agonistic effects in biological assay.

35 [0080] Therefore, it is desirable to be able to efficiently "switch" cell membrane bound antibody expression to secreted antibody expression in the vertebrate host cells with sufficient yields, in order to produce enough quantity of a particular antibody clone for functional assays.

40 [0081] In natural B lineage cells the switch from membrane bound to secreted antibody expression occurs via a mechanism of alternative splicing, in which in preB and B cells an alternative splice donor near the 3'end of the last heavy chain constant region exon is preferentially spliced to a splice acceptor of a membrane anchor exon downstream of the heavy chain constant regions exons. This way, an antibody heavy chain is produced in B cells with an extended C-terminal, membrane spanning domain, that anchors the heavy chain and thereby the entire heavy-light chain containing antibody in the cell membrane. The C-terminal, membrane spanning domain also interacts non-covalently with the membrane spanning components Ig-alpha (CD79a or mb-1) and Ig-beta (CD79b or B29), which likely results in better membrane anchoring and higher surface immunoglobulin expression in B lineage cells.

45 [0082] Once, a B cell differentiates further to the plasma cell stage, the alternative splicing does not occur any more and the alternative splice donor near the 3' end of the last heavy chain constant region is no longer recognized or utilized, and the mRNA template is terminated downstream of the heavy chain constant region stop codon, and a heavy chain of a secreted antibody is translated.

50 [0083] In order to exploit this natural mechanism of alternative splicing and "switching" from membrane bound to secreted expression of expressed antibodies, it is a useful embodiment of the method disclosed herein to construct the transposable vectors and diverse DNA libraries encoding proteins, including antibodies or fragments thereof, in such a way that the natural intron/exon structure of a constant antibody heavy chain, including the exons encoding the membrane spanning domains is maintained. This embodiment represents a clear advantage against retroviral expression systems, as the retroviral vector genome is already spliced before it is packaged into a retroviral particle and stably transduced

into the host cell genome.

[0084] Other viral vector systems may be restricted in the length of the DNA insert that can be incorporated into the vectors, thereby precluding the cloning of larger genomic regions into such expression vectors and thereby preventing the exploitation of the natural "switching" from membrane-bound to secreted antibody expression by alternative splicing.

5 Certain transposons (e.g. *Tol2*, see Fig. 3), have been characterized to be able to efficiently transpose more than 10 kb DNA fragments into vertebrate host cells without any loss in transposition efficiency (Kawakami Genome Biol. 8, Suppl I, S7 (2007)) Therefore, it is a useful embodiment of the method disclosed herein to construct transposable expression vectors comprising genomic exon/intron structures for better and proper expression and for the natural regulation switching from membrane bound to secreted antibody expression. The methods of the invention are useful to transpose DNA

10 fragments at least 5kb, at least 6kb, at least 7kb, at least 8kb, at least 9 kb, at least 10 kb in size into host cell genomes.

[0085] The differentiation of earlier B lineage differentiation stage that favors membrane bound antibody expression, to a later, plasma cell stage, that favors secreted antibody expression can be induced by B cell differentiation factors, such as, but not limited to, CD40/IL4 triggering, or stimulation by mitogens, such as, but not limited to, lipopolysaccharide (LPS), or other polyclonal activators, *Staph. aureus* Cowan (SAC) strain activators, and CpG nucleotides, or any combination thereof.

15 [0086] Preferably, this differentiation is effected in transformed cells, in which the proliferation can artificially be inhibited, such that proper B cell differentiation can again occur, as it has been described for A-MuLV transformed murine preB cells, in which the Abelson tyrosine kinase is specifically inhibited by the tyrosine inhibitor Gleevec (Muljo et al. Nat. Immunol. 4, 31-37 (2003)). Therefore, it is a preferred embodiment to utilize Ig-null A-MuLV transformed murine 20 preB cells for the method, which by treatment with Gleevec, can again differentiate to more mature B cell stages, including plasma cells, which then secrete sufficient amounts of secreted antibody for additional functional testing on the basis of alternative splicing of genomic heavy chain expression constructs. It is a preferred embodiment of the method disclosed herein, to further improve such B-lineage cell differentiation by stable overexpression of anti-apoptotic factors, known in the art, including, but not limited to, bcl-2 or bcl-x<sub>L</sub>.

25 [0087] After step (iv), the enrichment of transposed vertebrate host cells as described above has been performed, optionally, additional cell enrichments according to the above-mentioned methods may be performed (step (v)), until cell populations, or individual cells are isolated expressing proteins, including antibodies and fragments thereof, with desired functional and/or binding properties.

30 [0088] Step (vi) of the method disclosed herein is then performed in order to isolate the relevant coding information contained in the transposed vertebrate host cells, isolated for a desired functional and/or binding property.

35 [0089] A useful embodiment of step (vi) for the isolationm cloning and sequencing of the relevant coding information for a desired functional or binding protein, including an antibody or antibody fragment thereof, contained in the isolated cells, is to utilize genomic or RT-PCR amplification with specific primer pairs for the relevant coding information comprised in the transposed DNA constructs, and to sequence the genomic or RT-PCR amplicons either directly, or after sub-cloning into sequencing vectors, known in the art, e.g., but not limited into TA- or Gateway-cloning vectors.

40 [0090] Another useful embodiment of step (vi) is to subject the enriched cell populations of steps (iv) or (v), which exhibit a desired functional or binding phenotype to next-generation ("deep") sequencing (Reddy et al. Nat. Biotech. 28, 965-969 (2010)), in order to retrieve directly and in one step a representative set of several thousands of sequences for the coding information contained in the transposed DNA constructs. Based on a bioinformatics analysis of the relative frequency of sequences identified from the enriched cell populations, it allows a prediction about which sequences encoded a functional or binding protein, including an antibody or fragment thereof (Reddy et al. Nat. Biotech. 28, 965-969 (2010)). Statistically overrepresented sequences are then resynthesized and cloned into expression vector for expression as recombinant proteins, antibodies or fragments thereof, in order to characterize them functionally and for their binding properties. This method can significantly accelerate the identification of relevant sequences within a functionally and phenotypically enriched cell population, that expresses proteins with functional or target specific properties.

45 [0091] Yet another useful embodiment of the method disclosed herein is to utilize transposition-mediated vertebrate cell expression of proteins, including antibodies or fragments thereof, for the mutagenesis and optimization of desired proteins, including the affinity optimization of antibodies and fragments thereof.

50 [0092] This can be achieved by isolating the genes encoding the proteins, including antibody chains or fragments thereof, from transposed vertebrate cell populations enriched for a desired binding or functional phenotype according to the methods disclosed in step (iv), such as but not limited to, by genomic PCR or RT-PCR amplification under mutagenizing conditions, know in the art. The mutagenized sequences can then be re-cloned into transposition vectors and then again be transposed into vertebrate host cells, in order to subject them to screening according to the methods disclosed herein, for improved functional or binding properties.

55 [0093] In one useful embodiment of this approach, specific primers are used that allow the PCR amplification under mutagenizing conditions of complete transposed constructs, including the flanking ITRs.

[0094] By this method a mutagenized PCR amplicon containing a defined average frequency of random mutations is generated from the functionally or phenotypically selected transposed cells. Said PCR amplicon with controlled mutations

(variations) of the original templates can now directly be re-transposed into new vertebrate host cells, according to preferred embodiments disclosed in the methods applicable in step (ii).

[0095] The main advantage of this method over other approaches of genetically modifying vertebrate cells is, that with this technology no time-consuming re-cloning of the mutagenized PCR amplicons and time consuming quality control of the mutagenized sequences into expression vectors is required, which is a mandatory requirement in all other plasmid-based or viral expression systems, if a mutagenized sequence shall be subjected to another round of screening.

[0096] Because transposition of DNA only requires the presence of ITRs flanking the coding region of genes of interest, PCR-amplified mutagenized PCR amplicons can directly be reintroduced and re-transposed into novel vertebrate host cells for expression and screening for improved properties and/or affinity matured mutants.

[0097] Taken together, the methods disclosed herein, of utilizing TEs for the stable genetic modification of vertebrate host cells with transposable constructs and/or diverse transposable DNA libraries encoding proteins, including antibodies and fragments thereof, offers unparalleled efficiency, flexibility, utility and speed for the discovery and optimization of said proteins for optimal desired functional or binding phenotypes.

15 Examples:

**Example 1:** Cloning of transposable light chain expression vector for human antibody kappa light chains compatible with the PiggyBac transposase enzyme

[0098] A transposable expression vector for human kappa light chains can be generated by cloning of the ITRs from the *PiggyBac* transposon system up and downstream of a human immunoglobulin kappa light chain expression cassette.

[0099] For this, as a first step, the minimal sequences for the up- and downstream ITRs of the *PiggyBac* transposon can be derived from pXLBaclI (published in US 7,105,343 and can be gene synthesized with flanking restriction enzyme sites for cloning into the mammalian expression vector pIRES-EGFP (PT3157-5, order #6064-1, Invitrogen, Carlsbad, CA, USA)

[0100] The upstream *PiggyBac* ITR sequence with the 5' terminal repeat has to be gene synthesized with flanking MunI restriction enzyme sequence (boldface and underlined), compatible with a unique MunI restriction enzyme site in pIRES-EGFP, and additional four random nucleotides (in lowercase letters) allowing proper restriction enzyme digestion. This sequence is as follows:

30 Seq-ID1:

5' -

35 **atatTTGTTAACCTAGAAAGATAGTCTGCGTAAATTGACGCATGCATTCTGAAATATTGCTCTC**  
TTTCTAAATAGCGCGAATCCGTCGCTGTGCATTAGGACATCTCAGTCGCCGCTGGAGCTCCGTGAGGCG  
TGCTTGTCAATGCCGTAAAGTGTCACTGATTTGAECTATAACGACCGCGTGAGTC  
40 AAAATGACGCATGATTA  
TCTTTTACGTGACTTTAAGATTTAACTCATACGATAATTATATTGTTATTCATGTTCTACTTACGTGATA  
ACTTATTATATATATTTCTTGTATACAATTGat at - 3'

[0101] The MunI restriction enzyme sites at each end are underlined and typed in boldface print, the random nucleotide additions at the termini are printed in lowercase.

[0102] The downstream *PiggyBac* ITR sequence with the 3' terminal repeat has to be gene synthesized with flanking Xhol restriction enzyme sequence(boldface and underlined)compatible with a unique Xhol restriction enzyme site in pIRES-EGFP, and additional four random nucleotides (in lowercase letters) allowing proper restriction enzyme digestion. This sequence is as follows:

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## Seq-ID2:

5'-

5 atatCTCGAGTAAACCCTAGAAAGATAATCATATTGTGACGTACGTTAAAGATAATCATGCGAAAATTGAC  
 GCATGTGTTTATCGGTCTGTATATCGAGGTTATTATTAAATTGAATAGATATTAAAGTTTATTATATT  
 ACACTTACATACTAATAATAAAATTCAACAAACAATTATTATTTATGTTATTATTATAA  
 10 ACTCAAAATTCTTCTATAAAGTAACAAAACTTTATCCTCGAGata - 3'

[0103] Upon MunI restriction enzyme digestion of the gene synthesized Seq-ID1, the DNA fragment can be ligated into MunI linearized pIRES, generating pIRES-EGFP-TR1 according to standard methods, known in the art. The proper orientation of the insert can be verified by diagnostic restriction enzyme digestion, and/or by DNA sequencing of the cloned construct (Fig. 5a).

[0104] In a next step gene synthesized and Xhol digested DNA fragment Seq-ID2, can be ligated into Xhol linearized pIRES-EGFP-T1 (Fig. 5a) by standard methods known in the art in order to generate pIRES-EGFP-T1T2, containing both *PiggyBac* ITRs up and downstream of the IRES-EGFP expression cassette (Fig. 5b). The proper orientation of the insert can be verified by diagnostic restriction enzyme digestion, and/or by DNA sequencing of the cloned construct (Fig. 5b).

[0105] For the cloning of a human immunoglobulin kappa light chain into the vector pIRES-EGFP-T1T2, the human Ig kappa light chain from human anti-TNF-alpha specific antibody D2E7 can be synthesized, which can be retrieved from European patent application EP 1 285 930.

[0106] The V region of D2E7 fused in frame to a Vk1-27 leader sequence (Genbank entry: X63398.1), which is the closest germ-line gene V-kappa family member V-kappa of D2E7, and to the human kappa constant region (Genbank entry: J00241) has the following nucleotide sequence:

## Seq-ID3:

30 5'-atggacatgagggtccctgctcagctcctggactcctgctgctctggctcccagggtgccagatgtGACATCCA  
 GATGACCCAGTCTCCATCCTCCCTGTCTGCATCTGTAGGGGACAGAGTCACCATCACTGTCGGCAAGTCAGGGC  
 ATCAGAAAATTACTTAGCCTGGTATCAGCAAAACCAGGGAAAGCCCCATAAGCTCCTGATCTATGCTGCATCCACTT  
 35 TGCAATCAGGGTCCCCTCGGTTCACTGGCAGTGATCTGGACAGATTCACTCTCACCATCAGCAGCCTACA  
 GCCTGAAGATGTTGCAACTTATTACTGTCAAAGGTATAACCGTGCACCGTATACTTTGCCAGGGACCAAGGTG  
 GAAATCAAGCGCTCTGTGGCTGCACCATCTGTCTTCATCTCCGCCATCTGATGAGCAGTTGAAATCTGGAAC  
 40 CCTCTGTTGTGCTGCTGAATAACTCTATCCCAGAGAGGCCAAAGTACAGTGGAGGTGGATAACGCCCTCCA  
 ATCGGGTAACTCCCAGGAGAGTGTACAGAGCAGGACAGCAAGGACAGCACCTACAGCCTCAGCAGCACCC  
 CTGAGCAAAGCAGACTACGAGAACACAAAGTCTACGCCCTGCGAAGTCACCCATCAGGGCCTGAGCTGCCGTCA  
 CAAAGAGCTTCAACAGGGAGAGTGTAA-3'

45 [0107] This translates in the following amino acid sequence:

## Seq-ID4:

50 MDMRVPQLLGLLLLWLPGARCDIQMTQSPSSLSASVGDRVTITCRASQGIRNYLAWYQQKPGKAPKLLIYA  
 ASTLQSGVPSRFSGSGTDFLTLSLQPEDVATYYCQRYNRAPYTFQGQTKVEIKRSVAAPSVFIFPPSD  
 EQLKSGTASVVCLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYSLSSTTLSKADYEKHKVYAC  
 55 EVTHQGLSSPVTKSFNRGEC

[0108] The DNA fragment Seq-ID3 encoding the D2E7 Ig kappa light chain of Seq-ID4 can be gene synthesized and directly ligated by blunt-ended ligation into the unique EcoRV restriction enzyme site (which is also a blunt cutter), by

methods known in the art, resulting in construct pIRES-EGFP-T1T2-IgL (Fig. 5b)

5 [0109] Seq-ID3 has been engineered to contain a unique Eco47III restriction enzyme site in between the V-kappa and the C-kappa coding regions (highlighted in boldface and underlined), which allows the replacement of V-kappa regions in this construct against other V-kappa regions or V-kappa libraries, using a unique restriction enzyme upstream of V-kappa coding region in the construct, together with Eco47III. The proper orientation of the kappa light chain insert can be verified by diagnostic restriction enzyme digestion, and/or by DNA sequencing of the cloned construct (Fig. 5b).

[0110] The sequence for the transposable human antibody kappa light chain vector pIRES-EGFP-T1T2-IgL is provided in sequence Seq-ID5:

10 Seq-ID5:

5'-

15 GACGGATCGGGAGATCTCCGATCCCCTATGGTCGACTCTCAGTACAATCTGCTTGATGCCGCATAGTTAA

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5           GCCAGTATCTGCTCCCTGCTTGTGTTGGAGGTGCGTGAGTAGTCGCGAGCAAATTAAAGCTACAACAA  
 GGCAAGGCTTGACCGACAATTGTTAACCCCTAGAAAGATAGTCTGCGTAAATTGACGCATGCATTCTGAAA  
 TATTGCTCTCTTTCTAAATAGCGCGAATCCGTCGCTGTGCATTAGGACATCTCAGTCGCCGTTGGAGC  
 10          TCCCCTGAGGGCGTGTGCAATCGCGTAAGTGTCACTGATTTGAACATATAACGACCGCGTGAGTCAGGAA  
 GACGCATGATTATCTTTACGTGACTTTAACGTTAACTCATACGATAATTATATTGTTATTTCATGTTCT  
 ACTTACGTGATAACTTATTATATATATTGTTATACAATTGCAAGAATCTGCTTAGGGTTAGG  
 15          CGTTTGCGCTGCTTCGCGATGTACGGGCCAGATATACGCGTTGACATTGATTATTGACTAGTTATTAAATAG  
 TAATCAATTACGGGTCAATTAGTTCATAGCCCATAATGGAGTTCCGCGTTACATAACTTACGGTAAATGGC  
 CCGCCTGGCTGACCGCCCAACGACCCCCGCCATTGACGTCAATAATGACGTATGTTCCATAGTAACGCCA  
 ATAGGGACTTCCATTGACGTCAATGGGTGGACTATTTACGGTAAACTGCCACTTGGCAGTACATCAAGTG  
 20          TATCATATGCCAAGTACGCCCTATTGACGTCAATGACGGTAAATGGCCGCTGGCATTATGCCAGTAC  
 ATGACCTTATGGACTTCTACTTGGCAGTACATCTACGTATTAGTCATCGCTATTACCATGGTATGCGG  
 TTTTGGCAGTACATCAATGGCGTGGATAGCGGTTGACTCACGGGATTCCAAGTCTCCACCCATTGAC  
 25          GTCAATGGGAGTTGTTGGCACCAAATCAACGGACTTCCAAAATGCGTAACAACACTCCGCCATTG  
 ACGCAAATGGGCGGTAGGCGTGTACGGTGGAGGTCTATATAAGCAGAGCTCTGGCTAACTAGAGAACCC  
 ACTGCTTACTGGTTATCGAAATTAAATACGACTCACTATAGGGAGACCAAGCTGGTACCGAGCTGGATC  
 GATatggacatgagggtccctgctcagctcctggactcctgctgctctggctccaggtgccagatgtGAC  
 30          ATCCAGATGACCCAGTCTCCATCCTCCGTCTGCATCTGTAGGGACAGAGTCACCACACTGTCGGGCA  
 AGTCAGGGCATCAGAAATTACTTAGCCTGGTATCAGCAAAACCAGGGAAAGCCCTAACGCTCTGATCTAT  
 GCTGCATCCACTTGCATCAGGGTCCCCTCGGTTAGTGGCAGTGGATCTGGACAGATTCACTCTC  
 ACCATCAGCAGCCTACAGCCTGAAGATGTTCAACTTATTACTGTCAAAGGTATAACCGTGCACCGTATACT  
 35          TTTGGCAGGGACCAAGGTGGAAATCAAGCGCTCTGTGGCTGCACCACCTGTCTTCATCTCCGCCATCT  
 GATGAGCAGTTGAAATCTGAACTGCCTCTGTGTTGTGCCTGCTGAATAACTCTATCCCAGAGAGGCCAA  
 GTACAGTGGAAAGGTGGATAACGCCCTCCAATCGGGTAACCTCCAGGAGAGTGTACAGAGCAGGACAGCAAG  
 GACAGCACCTACAGCCTCAGCAGCACCTGACGCTGAGCAAAGCAGACTACGAGAACACAAAGTCTACGCC  
 40          TGCAGTCACCCATCAGGGCTGAGCTCGCCGTACAAAGAGCTAACAGGGAGAGTGTAAATCTGC  
 GGCGCGTCAGGAAATTAGTGGATCCACTAGTAACGGCCAGTGTGCTGAAATTACGCTGTCTGC  
 GAGGGCCAGCTGTTGGGTGAGTACTCCCTCTCAAAAGCGGGCATGACTCTGCGCTAACGATTGTCAGTT  
 CAAAAACGAGGAGGATTGATATTCACCTGGCCCGGGTATGCCTTGAGGGTGGCGTCCATCTGGTC  
 AGAAAAGACAATTTTGTCAAGCTTGAGGTGTGGCAGGCTTGAGATCTGGCATAACACTGAGTGAC  
 AATGACATCCACTTGCCTTCTCCACAGGTGTCCACTCCCAGGTCAAACGTCAGGTCGAGCATGCATCT  
 45          AGGGCGGCAATTCCGCCCTCTCCCTCCCCCCCCCTAACGTTACTGCCGAAGCCGCTGGAAATAAGGCC  
 GGTGTGCGTTGTCTATATGTGATTTCCACCATATTGCCGTCTTGGCAATGTGAGGGCCGGAAACCTG  
 GCCCTGTCTTGTACGAGCATTCTAGGGTCTTCCCTCTGCCAAAGGAATGCAAGGTCTGTTGAATG  
 TCGTGAAGGAAGCAGTTCTCTGGAAGCTTCTGAAAGACAAACAGCTGTAGCGACCCCTTGCAGGCAGC  
 50          GGAACCCCCCAGCTGGCGACAGGTGCCCTGCGGCCAAAGCCACGTGTATAAGATAACACCTGCAAAGGCC  
 CACAACCCCAGTGCACGTTGTGAGTGGATAGTGTGAAAGAGTCAAATGGCTCTCCTCAAGCGTATTCA  
 ACAAGGGCTGAAGGATGCCAGAAGGTACCCATTGTATGGATCTGATCTGGGCCCTCGGTGCACATGCT  
 TTACATGTGTTAGTCGAGGTTAAAAAAACGTCTAGGCCCCCGAACCACGGGACGTGGTTTCTTGAA  
 AAACACGATGATAAGCTTGCCACAACCCGGATCCACCGTCGCCACCATGGTGAGCAAGGGCGAGGAGCTG

TTCACCGGGGTGGTGCCCATCCTGGTCAGCTGGACGGCAGCTAACGGCCACAAGTTAGCGTGTCGGC  
 GAGGGCGAGGGCGATGCCACCTACGGCAAGCTGACCTGAAGTTCATCTGCACCACCGGCAAGCTGCCGTG  
 CCCTGGCCCACCCCTGTGACCACCTGACCTACGGCGTGCAGTGCTCAGCCGCTACCCGACCACATGAAG  
 5 CAGCACGACTTCAAGTCCGCATGCCGAAGGCTACGTCAGGAGCGACCCATCTTCAAGGACGAC  
 GGCAACTACAAGACCCGCGCCGAGGTGAAGTTCGAGGGCGACACCCCTGGTAACCGCATCGAGCTGAAGGGC  
 ATCGACTTCAAGGAGGACGGCAACATCCTGGGGACAAGCTGGAGTACAACACTACAACAGCCACAACGTCTAT  
 10 ATCATGGCCGACAAGCAGAAGAACGGCATCAAGGTGAACCTCAAGATCCGCCACAACATCGAGGACGGCAGC  
 GTGCAGCTGCCGACCACTACCAGCAGAACACCCCCATCGGCACGGCCCGTGCCTGCCGACAACCAC  
 TACCTGAGCACCAGTCCGCCCTGAGCAAAGACCCAACGAGAAGCGCATCACATGGCCTGCTGGAGTTC  
 15 GTGACCGCCGCCGGATCACTCTGGCATGGACGAGCTGTACAAGTAAAGCGGCCCTAGAGCTCGCTGATCA  
 GCCTCGACTGTGCCCTAGTTGCCAGCCATCTGTGTTGCCCTCCCCGTGCCTCCTTGACCCCTGGAAG  
 GTGCCACTCCCAGTGCCTTCCTAATAAAATGAGGAAATTGCATCGATTGCTGAGTAGGTGTCATTCTA  
 TTCTGGGGGTGGGTGGGCAGGACAGCAAGGGGAGGATTGGGAAGACAATAGCAGGCATGCTGGGATG  
 20 CGGTGGCTCTATGGCTTCTGAGGCGGAAAGAACAGCTGGGCTCGAGGATAAAAGTTTGTACTTTATA  
 GAAGAAAATTTGAGTTTTGTTTAAATAAAATAAAACATAAAATAATTGTTGTTGAATTTATA  
 TTAGTATGTAAGTGTAAATATAAAACTTAATATCTATTCAAATTAAATAAAACCTCGATATACAGAC  
 CGATAAAACACATGCGTCAATTTACGCATGATTATCTTAACGTACGTACAATATGATTATCTTCTAGG  
 25 GTTAACTCGAGTCGATTCTAGTTGTGGTTGTCCAAACTCATCAATGTATCTTATCATGTCTGTATACCGTC  
 GACCTCTAGCTAGAGCTTGGCGTAATCATGGTCAAGCTGTTCTGTGAAATTGTTATCCGCTACAAT  
 TCCACACAACATACGAGCCGGAAGCATAAAAGTGTAAAGCCTGGGTGCTTAATGAGTGAGCTAACTCACATT  
 30 AATTGCGTTGCGCTCACTGCCGTTCCAGTCGGAAACCTGCGTGCAGCTGCATTAATGAATCGGCCA  
 ACGCGGGGGAGAGGCCTTGCCTTGGCGCTCTCCGCTCGCTCACTGACTCGCTGCGCTCGGT  
 CGTTGGCTGCGCGAGCGGTATCAGCTCACTCAAAGGCGGTAAACGGTTATCCACAGAAATCAGGGGATAA  
 CGCAGGAAAGAACATGTGAGCAAAGGCCAGCAAAGGCCAGGAACCGTAAAAGGCCGTTGCTGGCGTT  
 35 TTTCCATAGGCTCCGCCCCCTGACGAGCATCACAAAATCGACGCTCAAGTCAGAGGTGGCGAAACCGAC  
 AGGACTATAAGATAACCAGGCCTTCCCCCTGGAAGCTCCCTGCGCTCTCTGTTCCGACCCCTGCCGCT  
 TACCGGATACCTGTCGCCTTCTCCCTGGGAAGCGTGGCGCTTCTCAATGCTCACGCTGTAGGTATCT  
 40 CAGTCGGTAGGTGTTCGCTCCAGCTCAAGCTGGCTGTGACGAACCCCCGTTAGCCGACCGCTGCGC  
 CTTATCCGTAACATCGTCTTGAGTCCAACCCGTAAGACAGACTTATGCCACTGGCAGCAGCCACTGG  
 TAACAGGATTAGCAGAGCGAGGTATGTAAGCGGTGCTACAGAGTTCTGAAGTGGTGGCTAACTACGGCTA  
 CACTAGAAGGACAGTATTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTCGGAAAAAGAGTTGGTAGCTC  
 45 TTGATCCGGAAACAAACCACCGCTGGTAGCGGTGGTTTTGTTGCAAGCAGCAGATTACGCCAGAAA  
 AAAAGGATCTCAAGAAGATCCTTGATCTTCTACGGGTCTGACGCTCAGTGGAAACGAAAACCTCACGTTA  
 AGGGATTTGGTCATGAGATTATCAAAAGGATCTCACCTAGATCCTTAAATTAAAAATGAAGTTTAA  
 ATCAATCTAAAGTATATGAGTAAACTTGGTCTGACAGTTACCAATGTTAATCAGTGAGGCACCTATCTC  
 50 AGCGATCTGTCTATTCGTTCCATAGTTGCCTGACTCCCCGTCGTGAGATAACTACGATACGGGAGGG  
 CTTACCATCTGGCCCCAGTGTGCAATGATACCGCGAGACCCACGCTCACCGCTCCAGATTATCAGCAAT  
 AAACCGCCAGCCGGAAGGGCGAGCGCAGAAGTGGCTCTGCAACTTATCCGCTCCATCCAGTCTATTAA  
 55 TTGTTGCCGGAAAGCTAGAGTAAGTAGTTGCCAGTTAATAGTTGCGCAACGTTGCCATTGCTACAGG  
 CATCGTGGTGTACGCTCGTGTGGTATGGCTTCATTAGCTCCGGTCCACGATCAAGGCGAGTTAC

ATGATCCCCATGTTGTGCAAAAAGCGGTTAGCTCCTCGTCCTCCGATCGTTGTCAGAAGTAAGTTGGC  
 CGCAGTGTATCACTCATGGTTATGGCAGCAGCTGCATAATTCTCTACTGTCATGCCATCCGTAAGATGCTT  
 TTCTGTGACTGGTGAGTACTCAACCAAGTCATTCTGAGAATAGTGTATGCGGCGACCGAGTTGCTCTGCC  
 5 GGCCTCAATACGGATAATACCGCGCCACATAGCAGAACTTAAAAGTGTCATCATTGGAAAACGTTCTTC  
 GGGCGAAAACCTCAAGGATCTTACCGCTGTTGAGATCCAGTTCGATGTAACCCACTCGTGCACCCAAC  
 10 ATCTTCAGCATCTTTACTTCAACCAGCGTTCTGGTGAGCAAAAACAGGAAGGAAAATGCCGAAAAAA  
 GGGATAAGGGCGACACGGAAATGTTGAATACTCATACTCTCCTTTCAATATTATTGAAGCATTATCA  
 GGGTTATTGTCTCATGAGCGGATACATATTGAATGTATTAGAAAATAACAAATAGGGTTCCGCGCAC  
 ATTCCCCGAAAAGTGCCACCTGACGTC-3'

15 **Example 2:** Cloning of a transposable heavy chain expression vector for human antibody gamma1 heavy chains:

[0111] In order to clone a transposable heavy chain vector, the ORF from pIRES-EGFP-T1T2-IgL simply needs to be exchanged with a fully human IgG1 heavy chain coding region.

[0112] For the replacement of the human kappa light chain in vector pIRES-EGFP-T1T2-IgL by a human immunoglobulin gamma-1 heavy chain, the VH region of antibody D2E7, which is specific for human TNF-alpha (see: EP 1 285 930 A2) can be synthesized and fused in frame to the leader sequence of a close germ-line VH3-region family member and in addition fused in frame to the coding region of a human gamma1 constant region (Genbank: J00228) including the membrane spanning exons (Genbank: X52847). In order to be able to replace the human Ig kappa light chain from pIRES-EGFP- T1T2-IgL unique ClaI and NotI restriction enzyme sites need to be at the 5' and the 3' end of the sequence (underlined), respectively, including four nucleotides flanking the restriction enzyme sites (highlighted in lowercase letters at the ends of the sequence), allowing proper restriction enzyme digestion of the gene-synthesized DNA fragment and ligation into the ClaI-NotI linearized pIRES-EGFP- T1T2-IgL backbone, according to standard methods. The sequence that needs to be gene synthesized is the following sequence:

30 Seq-ID6:  
 5'-

aattATCGATATGGAGTTGGGCTGAGCTGGTTTCCTTGTGCGATTTAGAAGGTGTCCAGTGTGAGGT  
 35 GCAGCTGGTGGAGTCTGGGGAGGCTTGGTACAGCCCGCAGGTCCCTGAGACTCTCCTGTGCGGCCTCTGG

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ATT CAC CTT GAT GATT ATGCC ATGC ACTGGT CCG CAAGCT CCAG GG AAGG GCT GG AATGGT CT CAG C  
 5 TAT CACT TGA ATAG TGGT CAC ATAG ACT ATGC GG ACT CT GT GG AGGG CG ATT CACC ATCT CCAG AGA CAA  
 CGCC AAGA ACT CCT GTAT CTG CAA ATGA AAC AGT CTG AGAG AGT GAGG ATAC GG CG TATA TT ACT GT GCG AA  
 AGT CTC GTAC CT TAGC ACC CGT CCT CCT TG ACT ATT GGGG CAAGG TACCC TGGT CAC CGT CT CG **AAGCGC**  
 10 **T**TCC ACCA AAGG GCCC ATCG GTCT CCC CTGG CACC CT CC AAAG GAC CCT CTGGGG CAAG CAG CC  
 GGG CT GC CT GG TCAAGG ACTACT TCCC GAAC CGT GAC GTG TG GA ACT CAGG CG CC TG ACC AG CG  
 CGT GC ACAC CT CCC GG CT GT CCT ACAGT CCT CAGG ACT CT ACT CC CT CAG CAG CG TG TG ACC GT GCC CT  
 CAG CAG CT TGG GACCC AGAC CT ACAT CTG CAAC GT GAAT CACA AGCC AG CAAC ACCA AAGG TG GACA AGAA  
 AGTT GAGCC AAAT CTT GTG AC AAA ACTCAC ACAT GCC CAC CGT GCC CAG CAC CT GA ACT CCT GGGGG ACC  
 15 GTC AGT CT CTC CT TCC CCCCC AAAACCCA AAGG AC ACC CT CATG AT CT CCG GACCC CTG AGGT CAC AT GCG T  
 GGT GGT GG AC GTG AGCC AC GAAG ACC CTG AGGT CAAG TT CA ACT GG TAC GT GG AC GG CG TG AGGT GC ATAA  
 TGCC AAG AC AAAG CG CGG AGG AGC AGT ACA AC AGC AC GT ACC GT TG GT CAG CGT CCT CAC CGT CCT GCA  
 20 CCAGG ACT GG CT GAAT GG CAAGG AGT ACA AGT GCA AGGT CT CC AAAC AAG CC CT CC AGG CCCC AT CG AGAA  
 AACCAT CT CCA AAG CCA AAGGG CAG CCCC GAGA ACCAC AGGT GT AC ACC CT G CCCC AT CCGG AT GAG CT  
 GACCA AAG ACCAGG TCAG CCT GAC CT GG TCAA AGG CT CT AT CC CAG CG AC AT CG CG TG GG AGT GGG A  
 25 GAG CAAT GGG CAG CGG AGA ACA ACT ACA AG ACCAC G C C T C C G T G C T G ACT CC GAC GG C T C T C T C C T  
 CTAC AGC AAG CT ACC GT GG AC AAG AGC AGG GT GG CAG CAG GGA AC GT CT T C T CAT G C T C C G T G AT G C AT G A  
 GG CT CT G CAC AACC ACT ACAC ACAGA AG AGC CT C C C T G T C C G G AG C T G C A A C T G G AGG AG AG C T G T G C  
 GGAGG CG CAGG AC GGG GAG CT GG AC GGG CT GT GG AC GACC AT CACC AT TT CAT CAC ACT CTT C C T G T T AAG  
 30 CGT GT GCT ACAGT GCC ACC GT CAC CT T C T CAAG GT GAAGT GG AT CTT C C T C G G T G G AC CT G AAG C A  
 GACC AT CAT CCCC GACT ACAGG AAC ATG AT CGG ACAG GGG GCT **AGG GGG CG Cg tc g -3'**

**[0113]** From the start codon in position 11 of Seq-ID6, this nucleotide sequence translates to the human IgG1 heavy chain of anti-TNF-alpha specific clone D2E7 (see: EP 1 285 930 A2), but including the human gamma1 transmembrane exons M1 and M2. The protein translation is provided in Seq-ID7 below.
   
 35

#### Seq-ID7:

40 MEFGLSWVFLVAILEGVQCEVQLVESGGLVQPGRSRLSCAASGFTFDDYAMHWVRQAPGKGLEWVSAITW  
 NSGHIDYADSVEGRFTISRDNAKNSLYLQMNSLRAEDTAVYYCAKVSYLSTASSLDYWQGTLVTVSSASTK  
 GPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVWSNSGALTSGVHTFPAPLQSSGLYSLSVVTPSSL  
 45 GTQTYICNVNHKPSNTKVDKKVEPKSCDKTHTCPPCPAPELLGGPSVLFPPKPDTLMISRTPEVTCVVVD  
 VSHEDEPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTS  
 KAKGQPREPQVYTLPPSRDELTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTPPVLDSDGSFFLYSK  
 LTVDKSRWQQGNVFCSVMHEALHNHYTQKSLSLSPELQLEESCAEAQDGELDGLWTTITIFITLFLLSVCY  
 50 SATVTFVKVWKWIFSSVVDLKQTIIIPDYRNMIGQGA

**[0114]** The DNA fragment Seq-ID6 encoding the D2E7 Ig gamma-1 heavy chain can then be double-digested by Clal and NotI restriction enzymes and directionally ligated into Clal and NotI linearized pIREs-EGFP- T1T2-IgL, resulting in construct pIREs-EGFP- T1T2-IgL (Fig. 6)
   
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**[0115]** Seq-ID6 has also been engineered to contain a unique Eco47III restriction enzyme site in between the V-heavy variable and the C-gamma1 constant coding regions (highlighted in boldface and underlined), which allows the replacement of V-heavy regions in this construct against other V-heavy regions or V-heavy libraries, using a unique restriction enzyme upstream of V-heavy coding region in the construct, together with Eco47III. The correct ligation of the insert

can be verified by diagnostic restriction enzyme digestion, and/or by DNA sequencing of the cloned construct (Fig. 6).  
**[0116]** The sequence for the transposable human antibody gamma-1 heavy chain vector pIRES-EGFP- T1T2-IgH is provided in sequence Seq-ID8.

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## Seq-ID8:

5' -

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GACGGATCGGGAGATCTCCGATCCCCTATGGTCGACTCTCAGTACAATCTGCTCTGATGCCGCATAGTTAA  
 GCCAGTATCTGCTCCCTGCTTGTGTGGAGGTCGCTGAGTAGTGCGCGAGCAAATTAAAGCTACAACAA  
 GGCAAGGCTTGACCGACAATTGTTAACCCCTAGAAAGATAGTCTGCGTAAATTGACGCATGCATTCTTGGAAA  
 TATTGCTCTCTTTCTAAATAGCGCGAATCCGTCGCTGTGCATTTAGGACATCTCAGTCGCCGCTTGGAGC  
 15 TCCCGTGAGGCGTGTCAATGCGGTAAAGTGTCACTGATTTGAACATAACGACCGCGTGAGTCAAAAT  
 GACGCATGATTATCTTTACGTGACTTTAACATACGATAATTATATTGTTATTCATGTTCT  
 ACTTACGTGATAACTTATTATATATTTCTGTTACAAATTGCATGAAGAATCTGCTTAGGGTTAGG  
 20 CGTTTGCCTGCTTCGCGATGTACGGGCCAGATATACGCGTTGACATTGATTATGACTAGTTATTAAATAG  
 TAATCAATTACGGGTCAATTAGTCATAGCCCATAATGGAGTTCCGCGTTACATAACTACGGTAAATGGC  
 CCGCCTGGCTGACGCCAACGACCCCCGCCATTGACGTCAATAATGACGTATGTTCCCATAGTAACGCCA  
 ATAGGGACTTCCATTGACGTCAATGGTGGACTATTACGGTAAACTGCCCACCTGGCAGTACATCAAGTG  
 25 TATCATATGCCAAGTACGCCCTATTGACGTCAATGACGGTAAATGCCCGCCTGGCATTATGCCAGTAC  
 ATGACCTTATGGGACTTCCACTTGGCAGTACATCTACGTATTAGTCATCGCTATTACCATGGTATGCCG  
 TTTGGCAGTACATCAATGGCGTGGATAGCGGTTGACTCACGGGATTCCAAGTCTCCACCCATTGAC  
 30 GTCAATGGGAGTTGTTTGGCACCAAAATCAACGGACTTCCAAAATGCGTAACAACCTCCGCCCCATTG  
 ACGCAAATGGCGTAGGCGTGTACGGTGGAGGTCTATATAAGCAGAGCTCTGGCTAACAGAGAACCC  
 ACTGCTTACTGGCTATCGAAATTAAATACGACTCACTATAGGGAGACCAAGCTGGTACCGAGCTGGATC  
 GATATGGAGTTGGCTGAGCTGGTTCCCTGCGATTAGAAGGTGTCCAGTGTGAGGTGCAGCTG  
 35 GTGGAGTCTGGGGAGGCTTGGTACAGCCCGCAGGTCCCTGAGACTCTCCTGTGCGGCCTGGATTCA  
 TTTGATGATTATGCCATGCACTGGTCCGGCAAGCTCCAGGGAAAGGGCCTGGAATGGTCTCAGCTATCA  
 TGGAATAGTGGTCACATAGACTATGCGGACTCTGTGGAGGGCGATTACCACCTCCAGAGACAACGCCAAG  
 AACCTCCCTGTATCTGCAAATGAACAGTCTGAGAGCTGAGGATACGGCCGTATATTACTGTGCGAAAGTCTCG  
 40 TACCTTAGCACC CGTCTCCCTGACTATTGGGCCAAGGTACCGTCCCTGAGCGCTTCACC  
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AAGGGCCCATCGTCTTCCCCCTGGCACCCCTCCTCCAAGAGCACCTCTGGGGCACAGCAGCCCTGGCTGC  
 CTGGTCAAGGACTACTTCCCCGAACCGGTGACGGTGTGAACTCAGGCGCCCTGACCAGCGCGTGCAC  
 ACCTTCCCGCTGTCCTACAGTCCTCAGGACTCTACTCCCTCAGCAGCGTGGTGACCGTGCCCTCAGCAGC  
 5 TTGGGCACCCAGACCTACATCTGCAACGTGAATCACAAGCCCAGCAACACCAAGGTGGACAAGAAAAGTTGAG  
 CCCAAATCTTGTGACAAAACACACATGCCAACCGTGCCAGCACCTGAACCTCTGGGGGACCGTCAGTC  
 TTCCTCTCCCCAAAACCCAAGGACACCCCTCATGATCTCCGGACCCCTGAGGTACATGCGTGGTGGTG  
 10 GACGTGAGCCACGAAGACCCGTAGGTCAAGTTCACTGGTACGTGGACGGCGTGGAGGTGCATAATGCCAAG  
 ACAAAAGCCGCGGGAGGAGCAGTACAACACAGCACGTACCGTGTGGTCAGCGTCCTCACCGTCCTGCACCAGGAC  
 TGGCTGAATGGCAAGGAGTACAAGTCAAGGTCTCAACAAAGCCCTCCAGCCCCATCGAGAAAACCATC  
 15 TCCAAAGCCAAGGGCAGCCCCGAGAACACCAGGTGTACACCCGTCCCCATCCGGATGAGCTGACCAAG  
 AACCAAGGTCACTGCCGTGCTGGTCAAAGGTTCTATCCAGCGACATGCCGTGGAGTGGGAGAGCAAT  
 GGGCAGCCGGAGAACAACTACAAGACCAACGCCTCCCGTGTGGACTCCGACGGCTCCTTCTCCTACAGC  
 AAAGCTACCGTGGACAAGAGCAGGTGGCAGGGGAACGTCTCTCATGCTCCGTGATGCATGAGGCTCTG  
 20 CACAACCAACTACACACAGAACAGGCTCTCCCTGTCTCCGGAGCTGCAACTGGAGGAGAGCTGTGCGGAGGCG  
 CAGGACGGGGAGCTGGACGGCTGTGGACGACCATCACCATTTCATCACACTCTTCTGTTAAGCGTGTGC  
 TACAGTGCCACCGTCACCTCTTCAAGGTGAAGTGGATCTTCTCTCGTGGTGGACCTGAAGCAGACCATC  
 ATCCCCGACTACAGAACATGATCGGACAGGGGCTAGGCGCCGCTGACGGAATTCACTGGATCCACT  
 25 AGTAACGGCCGCCAGTGTGCTGGAATTAAATTGCGCTGTGCGAGGGCCAGCTGTTGGGTGAGTACTCCCTC  
 TCAAAAGCGGGCATGACTCTGCGCTAAGATTGTCAGTTCAAAAACGAGGAGGAGTTGATATTCACCTGG  
 CCCGCGGTGATGCCCTTGAGGGTGGCCCGTCCATCTGGTCAAGAAAGACAATCTTTGTTGTCAGCTTG  
 30 AGGTGTGGCAGGCTTGAGATCTGCCATACACTTGAGTGACAATGACATCCACTTGCCTTCTCCACAG  
 GTGTCCACTCCAGGTCAACTGCAGGTGAGCATGCTAGGGCGCCAATTCCGCCCTCTCCCTCCCC  
 CCCCCCTAACGTTACTGGCGAAGCCGCTTGGAAATAAGGCCGTGTGCGTTGTCTATATGTGATTTCCAC  
 CATATTGCCGTCTTGCAATGTGAGGGCCGAAACCTGCCCTGTCTTGTGACGAGCATTCTAGGG  
 35 TCTTCCCTCTGCCAAAGGAATGCAAGGTCTGTTGAATGCGTAAGGAAGCAGTCCTCTGGAGCTTC  
 TTGAAGACAAACAACGTCTGTAGCGACCTTGCAGGCAGCGGAACCCCCACCTGGCGACAGGTGCCTCTG  
 CGGCCAAAGCCACGTGTATAAGATAACACCTGCAAAGGCCACAACCCAGTGCCACGTTGTGAGTTGGAT  
 40 AGTTGTGGAAAGAGTCAAATGGCTCTCTCAAGCGTATTCAACAAGGGCTGAAGGATGCCAGAGTAC  
 CCATTGTATGGGATCTGATCTGGGCTCGGTGACATGCTTACATGTGTTAGTCGAGGTTAAAAAAACG  
 TCTAGGCCCGGAACCACGGGACGTGGTTTCTTGAAAAACACGATGATAAGCTGCCACAACCCGG  
 ATCCACCGGTGCCACCATGGTGAGCAAGGGCGAGGAGCTGTTACCGGGTGGTGCCTCCTGGTCAGC  
 45 TGGACGGCGACGTAAACGCCACAAGTCAGCGTGTCCGGCGAGGGCGAGGGCGATGCCACCTACGGCAAGC  
 TGACCTGAAAGTTCATCTGACCAACCGCAAGCTGCCGTGCCCTGGCCACCCCTCGTGACCAACCTGACCT  
 ACGGCGTGCAGTGCTTCAGCCGCTACCCGACCACATGAAGCAGCAGCAGACTCTCAAGTCCGCCATGCCG  
 AAGGCTACGTCCAGGAGCGCACCATCTCTCAAGGACGACGGCAACTACAAGACCCGCCAGGTGAAGT  
 50 TCGAGGGCGACCCCTGGTGAACCGCATCGAGCTGAAGGGCATGACTCAAGGAGGACGGCAACATCCTGG  
 GCCACAAGCTGGAGTACAACACAGCCACAACGTCTATATCATGGCGACAAGCAGAAGAACGGCATCA  
 AGGTGAACCTCAAGATCCGCCACAACATCGAGGACGGCAGCGTGCAGCTCGCCGACCAACTACCAGCAGAAC  
 55 CCCCCATCGCGACGGCCCCGTGCTGCTGCCGACAACCAACTACCTGAGCACCCAGTCCGCCCTGAGCAAAG  
 ACCCAACGAGAAGCGCGATCACATGGCCTGCTGGAGTTCGTGACCGCCGGGATCACTCGGCATGG

ACGAGCTGTACAAGTAAAGCGGCCCTAGAGCTCGTGATCAGCCTCGACTGTGCCCTAGTTGCCAGCCATC  
 5 TGTTGTTGCCCTCCCCCTGCCTTCCTGACCGTGGAAAGGTGCCACTCCCCTGCTCTTCTAATAAAAA  
 TGAGGAAATTGCATCGCATTGTCTGAGTAGGTGTCATTCTATTCTGGGGGTGGGGCAGGACAGCAA  
 GGGGGAGGATTGGGAAGACAATAGCAGGCATGCTGGGATGCGGTGGCTCTATGGCTCTGAGGCGGAAAG  
 AACCAGCTGGGCTCGAGGATAAAAGTTGTTACTTATAGAAGAAATTTGAGTTTGTTTTTAA  
 TAAATAAATAAACATAAAATAATTGTTGTAATTATTAGTATGTAAGTGTAAATATAATAAAAACCT  
 10 AATATCTATTCAAATTAAATAAAACCTCGATATACAGACCGATAAAACACATGCGTCAATTTCAGCATG  
 ATTATCTTAACGTACGTACAATATGATTATCTTAGGGTTAECTCGAGTGCATTCTAGTTGTGGTTG  
 TCCAAACTCATCAATGTATCTTATCATGTCATACCGTCACCTCTAGCTAGAGCTTGGCTAATCATGG  
 15 TCATAGCTGTTCCGTGAAATTGTTATCCGCTCACAAATTCCACACAACATACGAGCCGAAGCATAAAG  
 TGTAAGCCTGGGTGCCTAATGAGTGAAGCTAACACATTAATTGCGTGCCTACTGCCGTTTCAG  
 TCGGGAAACCTGTCGTGCCAGCTGCATTAATGAATCGGCCAACGCGGGGAGAGGCGGTTGCATTGG  
 CGCTCTCCGCTTCCTCGCTACTGACTCGCTCGCTCGGCTGCGGAGCGGTATCAGCTCAC  
 20 TCAAAGGCGTAATACGGTTATCCACAGAATCAGGGATAACGCAGGAAAGAACATGTGAGCAAAGGCCAG  
 CAAAAGGCCAGGAACCGTAAAAGGCCGCGTTGCTGGCGTTTCCATAGGCTCCGCCCCCTGACGAGCAT  
 CACAAAATCGACGCTCAAGTCAGAGGTGGCAGACAGGACTATAAGATAACCAGGCGTTCCAG  
 25 GGAAGCTCCCTCGCGCTCCCTGTTCCGACCCCTGCCCTACCGGATACCTGTCGCCCTTCCTCG  
 GGAAGCGTGGCGCTTCTCAATGCTCACGCTGTAGGTATCTCAGTCGGTAGGTCGCTCCAAGCTG  
 GGCTGTGTCACGAACCCCCCGTTCACGGCAGCGCCTATCCGGTAACTATCGTCTGAGTCAAC  
 CCGGTAAGACACGACTTATGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGAGG  
 30 GGTGCTACAGAGTCTTGAAGTGGTGGCTAACTACGGCTACACTAGAAGGACAGTATTGGTATCTCGCCT  
 CTGCTGAAGCCAGTTACCTCGGAAAAGAGTTGGTAGCTCTGATCCGCAAACAAACCACCGCTGGTAGC  
 GGTGGTTTTTGTGCAAGCAGCAGATTACGCGCAGAAAAAGGATCTAAGAAGATCCTTGATCTT  
 TCTACGGGTCTGACGCTCAGTGGAACGAAAACACGTTAAGGGATTGGTCATGAGATTATCAAAAAGG  
 35 ATCTCACCTAGATCCTTAAATTAAAAATGAAGTTAAATCAATCTAAAGTATATGAGATAACTTGG  
 TCTGACAGTTACCAATGCTTAATCAGTGAGGCACCTATCTCAGCGATCTGCTATTCGTTCATCCATAGTT  
 GCCTGACTCCCCGTCGTAGATAACTACGATAACGGGAGGGCTTACCATCTGGCCCCAGTGCTGCAATGATA  
 40 CCGCGAGACCCACGCTCACCGGCTCCAGATTATCAGCAATAAACCAGCCAGCCGAAGGGCCAGCGCAGA  
 AGTGGTCTGCAACTTATCCGCTCCATCCAGTCTATTAAATTGCGCCAGGATCTGTTGAGTACTCAACCAAGTCA  
 CCAGTTAATAGTTGCGAACGTTGCGCATTGCTACAGGCATCGTGGTAGCAGCTCGTGTGGTATG  
 GCTTCATTGAGCTCCGGTCCCAACGATCAAGGCGAGTTACATGATCCCCATGTTGCAAAAAAGCGGTT  
 45 AGCTCCTTCGGCCTCCGATCGTGTAGAAGTAAGTTGGCCAGTGTATCACTCATGGTATGGCAGCA  
 CTGCATAATTCTTACTGTCATGCCATCCGTAAGATGCTTTCTGACTGGTAGTACTCAACCAAGTCA  
 TTCTGAGAATAGTGTATGCGCGACCGAGTTGCTCTGCCGGCGTCAATACGGATAATACCGGCCACAT  
 50 AGCAGAACTTAAAGTGTCTCATCATTGAAAACGTTCTCAGGCGAAACTCTCAAGGATCTTACCGCTG  
 TTGAGATCCAGTTGATGTAACCCACTCGTGCACCCACTGATCTCAGCATCTTACTTACCGCTT  
 TCTGGGTGAGCAAAACAGGAAGGCAAAATGCCGAAAAAGGAATAAGGGCGACACGGAAATGTTGAATA  
 55 CTCATACTCTCCTTTCAATATTATGAGCATTATCAGGGTTATGTCATGAGCGGATAACATATT  
 GAATGTATTAGAAAAATAACAAATAGGGTTCCGCGCACATTCCCCGAAAGTGCCACCTGACGTC-3'

[0117] Examples 1 and 2 provide the basic human transposable expression vectors for human antibody kappa light

and human gamma-1 light chains and therefore for full-length, membrane bound human IgG1.

**Example 3:** Generation of transposable DNA libraries encoding human antibody heavy and light chain libraries.

[0118] In order to generate diverse transposable DNA libraries encoding human antibody heavy and light chain libraries, only the VL and VH regions from transposable vectors pIRES-EGFP-T1T2-IgL of Example 2 and pIRES-EGFP- T1T2-IgH of Example 3, respectively, need to be replaced. This can be done by gene synthesizing of human VH and VL coding regions flanked by Clal and Eco47III restriction enzyme sites, and by allowing nucleotide variations in certain HCDR and LCDR positions, as provided in Seq-ID9, which encodes libraries for variable heavy chain domains, and Seq-ID10, which encodes libraries for variable light chain domains, because both of them contain N-sequences in the HCDR3 (boldface), and LCDR3 (boldface), respectively. Both Seq-ID9 and Seq-ID10 sequences are flanked by C1al and Eco47III restriction enzymes (underlined), respectively, including four nucleotides flanking the restriction enzyme sites (highlighted in lowercase letters at the ends of the sequence), allowing proper restriction enzyme digestion of the gene-synthesized DNA fragments and directed ligation into Clal-Eco47III linearized pIRES-EGFP- T1T2-IgH and pIRES-EGFP- T1T2-IgL backbones, respectively.

Seq-ID9:

Seq-ID10:

35 5'-  
atggacatgagggtccctgctcagctcctggactcctgctgctggctccaggtgccagatgtGACATC  
CAGATGACCCAGTCTCCATCCTCCCTGTCTGCATCTGTAGGGGACAGAGTCACCACACTGTGCGGGCAACT  
CAGGGCATCAGAAATTACTTAGCCTGGTATCAGCAAAACCAGGGAAAGCCCCTAACGCTCCTGATCTATGCT  
40 GCATCCACTTGCAATCAGGGTCCCATCTGGTTAGTGGCAGTGGATCTGGGACAGATTCACTCTCAC  
ATCAGCAGCCTACAGCCTGAAGATGTTGCAACTTATTACTGTCAAAGGNNNNNNNNNNNNNNNNTATACTTT  
GGCCAGGGGACCAAGGTGAAATCAAGCGCTgcatt-3'

**[0119]** This way, diverse transposable DNA libraries, encoding antibody heavy and light chains on separate vectors, in which the expression of the antibody chains are transcriptionally and therefore operably linked to a green fluorescent marker protein can be generated.

## 50 Example 4: Cloning of a *PiggyBac* transposase expression vector

**[0120]** The ORF of functional *PiggyBac* transposase enzyme can be retrieved from US Patent US 7,105,343 B1 and is provided in Seq-ID11.

Seq-ID11:

5' -

5 ATGGGTAGTTCTTAGACGATGAGCATATCCTCTGCTCTGCAAAGCGATGACGAGCTTGGTGAG  
 GATTCTGACAGTGAATATCAGATCACGTAAGTGAAGATGACGCCAGAGCGATAAGAAGCGTTATA  
 GATGAGGTACATGAAGTGCAGCCAACGTCAGCGTAGTGAATATTAGACGAACAAATGTTATTGAAACA  
 10 CCAGGTTCTCATTGGCTCTAACAGAAATCTGACCTTGCCACAGAGGACTATTAGAGGTAAGAATAAACAT  
 TGTGGTCAACTCAAAGTCCACGAGCGTAGCCAGTCTGCACTGAACATTGTCAGATCTCAAAGAGGT  
 CCGACCGTATGTGCCCAATATATGACCCACTTTATGCTCAAACATTTTACTGATGAGATAATT  
 TCGGAAATTGTAATGGACAAATGCTGAGATATCATTGAAACGTCGGAATCTATGACAGGTGCTACATT  
 15 CGTGACACGAATGAAGATGAAATCTATGCTTCTTGGTATTCTGGTAATGACAGCAGTGAGAAAAGATAAC  
 CACATGTCCACAGATGACCTTTGATCGATCTTGTCAATGGTGTACGTCTGTAATGAGTCGTGATCGT  
 TTTGATTTTTGATACGATGCTTAGAATGGATGACAAAGTATACGGCCACACTTCGAGAAAACGATGTA  
 20 TTGACCATAGATGAAACAGTTACTTGGTTTAGAGGACGGTGTCCGTTAGGATGTATATCCAAACAAGCCA  
 AGTAAGTATGGAATAAAATCCTCATGATGTTGACAGTGGTACGAAGTATATGATAATGGAATGCCTTAT  
 TTGGGAAGAGGAACACAGACCAACGGAGTACCACTCGGTGAATACTACGTGAAGGAGTTATCAAAGCCTGTG  
 25 CACGGTAGTTGTCGAATATTACGTGTGACAATTGGTCACCTCAATCCCTTGGAAAAACTTACTACAA  
 GAACCGTATAAGTTAACCATGTGGAACCGTGGATCAAACAAACGCGAGATACCGGAAGTACTGAAAAAC  
  
 30 AGTCGCTCCAGGCCAGTGGAACATCGATGTTTGTGACGGACCCCTACTCTCGTCTCATATAAACCG  
 AAGCCAGCTAAGATGGTATACTTATTATCATCTGTGATGAGGATGCTCTATCAACGAAAGTACCGGTAAG  
 CCGCAAATGGTTATGTATTATAATCAAACAAAGGCGGAGTGGACACGCTAGACCAAATGTGTTCTGTGATG  
 ACCTGCAGTAGGAAGACGAATAGGTGGCTATGGCATTATTGTACGGAAATGATAAACATTGCCTGCATAAA  
 35 TCTTTATTATACAGCCATAATGTCAGTAGCAAGGGAGAAAAGGTTCAAAGTCGAAAAAATTATGAGA  
 AACCTTTACATGAGCCTGACGTACGTTATGCGTAAGCGTTAGAAGCTCTACTTGAAGAGATATTG  
 CGCGATAATATCTAAATTTGCCAAATGAAGTGCCTGGTACATCAGATGACAGTACTGAAGAGCCAGTA  
 40 ATGAAAAAAACGTACTTACTGTACTGCCCCCTCTAAATAAGGCAGAAGGCAAATGCATCGTGCAAAAAA  
 TGCAAAAAAGTTATTGTCGAGAGCATAATATTGATATGTGCCAAAGTTGTTTAG-3'

[0121] This sequence translates into the amino acid Seq-ID12.

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## Seq-ID12:

MGSSLDEHILSALLQSDDELVGEDSDSEISDHVSEDDVQSDTEEAFIDEVHEVQPTSSGSEILDEQN  
5 VIEQPGSSLASNRLTLPQRTIRGKNCWSTSKSTRRSRVSLNIVRSQRGPTRMCRNIYDPLLCF  
KLFFTDEIISEIVKWTNAEISLKRRESMTGATFRDTNEDEIYAFFGILVMTAVRKDNHMSTDLLFDRS  
LSMVYVSVMRSRDRFDLIRCLRMDDKSIRPTLRENDVFTPVRKIWDLFIHQCIQNYTPGAHLTIDEQ  
10 LLGFRGRCPFRMYIPNKPSKYGIKILMMCDSGTKYMINGMPYLGRGTQTNGVPLGEYYVKELSKP  
VHGSCRNITCDNWFTSIPLAKNLLQEPIKLTIVGTVRSNKREIPEVLKNSRSRPVGTSMFCFDGPLT  
LVSYKPKPAKMYVLLSSCDEDASINESTGKPQMVMYYNQTKGGVDTLDQMCsvMTCSRKTNRWP  
15 MALLYGMINIACINSFIIYSHNVSSKGKEVQSRKKFMRNLYMSLTSSFMRKLEAPTLKRYLRDNISNI  
LPNEVPGTSDDSTEPPVMKKRTYCTYCP SKIRRKANASCKKCKVICREHNIDMCQSC

[0122] In order to generate a vertebrate cell expression vector for the *PiggyBac* transposase enzyme, this ORF can be gene synthesized and cloned as a blunt ended DNA into the unique, blunt-cutting restriction enzyme site EcoRV in the standard vertebrate cell expression vector pCDNA3.1-hygro(+) (catalogue # V870-20, Invitrogen, Carlsbad, CA, USA), by methods known in the art. The correct ligation of the *PiggyBac* ORF, relative to the pCDNA3 promoter can be verified by diagnostic restriction enzyme digestion, and/or by DNA sequencing of the cloned *PiggyBac* expression construct pCDNA3.1-hygro(+-PB (Fig. 7).

[0123] The sequence of the *PiggyBac* expression construct pCDNA3.1-hygro(+-PB is provided in Seq-ID 13, below:

## Seq-ID13:

25 5' - GACGGATCGGGAGATCTCCGATCCCCTATGGTCGACTCTCAGTACAATCTGCTCTGAT  
GCCGCATAGTTAACGCCAGTATCTGCTCCCTGCTTGTGTTGGAGGTGCTGAGTAGTGCGCGAGCAAAATT  
30 TAAGCTACAACAAGGCAAGGCTTGACCGACAATTGCATGAAGAATCTGCTTAGGGTTAGGCCTTTGCGCTG  
CTTCGCGATGTACGGGCCAGATATACCGTTGACATTGATTATTGACTAGTTATTAGTAATAGCAATTACG

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GGGTCATTAGTCATAGCCCCATATATGGAGTCCCGCGTTACATAACTACGGTAAATGGCCCGCCTGGCTGA  
 CCGCCCAACGACCCCCGCCATTGACGTCAATAATGACGTATGTTCCCATAGTAACGCCAATAGGGACTTTC  
 CATTGACGTCAATGGGTGGACTATTTACGGTAAACTGCCACTGGCAGTACATCAAGTGTATCATATGCCA  
 5 AGTACGCCCTATTGACGTCAATGACGGTAAATGGCCGCCTGGCATTATGCCAGTACATGACCTTATGG  
 GACTTCCCTACTGGCAGTACATCTACGTATTAGTCATCGCTATTACCATGGTATGCCAGTACATGACCTTATGG  
 ATCAATGGCGTGGATAGCGGTTGACTCACGGGGATTCCAAGTCTCCACCCCATTGACGTCAATGGGAGT  
 10 TTGTTTGGCACAAAATCAACGGACTTCCAAAATGTCGTAACAACCTCGCCCCATTGACGCAAATGGGC  
 GGTAGGCGTGTACGGTGGAGGTCTATATAAGCAGAGCTCTGGCTAAGTAGAGAACCCACTGCTTACTGG  
 CTTATCGAAATTAAACGACTCACTATAGGGAGACCCAAGCTGGCTAGCGTTAAACTTAAGCTTGGTACCG  
 15 AGCTCGGATCCACTAGTCCAGTGTGGTGAATTCTGCAGATATGGTAGTTAGACGATGAGCATATCC  
 TCTCTGCTCTTGCAAGCGATGACGAGCTTGTGGTGAGGATTCTGACAGTGAAATATCAGATCACGTA  
 GTGAAGATGACGTCCAGAGCGATACAGAAGAAGCGTTATAGATGAGGTACATGAAGTGCAGCCAACGTCAA  
 GCGGTAGTGAATATTAGACGAACAAAATGTTATGAAACAACCAGGTTCTCATTGGCTTAACAGAATCT  
 20 TGACCTTGCCACAGAGGACTATTAGAGGTAAAGAATAAACATTGTTGGTCAACTTCAAAGTCCACGAGGCGTA  
 GCCGAGTCTCTGCACTGAACATTGTCAGATCTCAAAGAGGTCGCAGCGTATGTGCCGAATATATGACC  
 CACTTTATGCTCAAATTTTACTGATGAGATAATTGGAAATTGAAATGGACAATGCTGAGA  
 TATCATTGAAACGTCGGAATCTATGACAGGTCTACATTCGTGACACGAATGAAGATGAAATCTATGCTT  
 25 TCTTTGGTATTCTGGAATGACAGCAGTGAGAAAAGATAACCACATGTCACAGATGACCTTTGATCGAT  
 CTTTGTCAATGGGTACGTCTGTAAATGAGTCGTGATGTTGATTTTGTACGATGTCTTAGAATGG  
 ATGACAAAAGTATACGGCCACACTCGAGAAAACGATGTATTACTCCTGTTAGAAAAATATGGGATCTCT  
 30 TTATCCATCAGTGCATAACAAATTACACTCCAGGGCTCATTTGACCATAGATGAACAGTTACTGGTTTA  
 GAGGACGGTGTCCGTTAGGATGTATATCCAAACAAGCCAAGTAAGTATGAAATAAAATCCTCATGATGT  
 GTGACAGTGGTACGAAGTATATGATAAAATGGAATGCCTTATTGGAGAGGAACACAGACCAACGGAGTAC  
 CACTCGGTGAATACTACGTGAAGGAGTTACAAAGCCTGTGCACGGTAGTTGTCGTAATATTACGTGTC  
 35 ATTGGTTCACCTCAATCCATTGGCAAAACTACTACAAGAACCGTATAAGTTAACCAATTGTGGAACCG  
 TGCGATCAAACAAACGCGAGATACCGGAAGTACTGAAAAACAGTCGCTCCAGGCCAGTGGAACATCGATGT  
 TTTGTTTGACGGACCCCTTACTCTGCTCATATAACCGAAGCCAGCTAAGATGGTATACTTATTATCAT  
 40 CTTGTGATGAGGATGCTTCTATCACGAAAGTACCGTAAACCGCAAATGGTTATGTATTATAATCAAAC  
 AAGGCGGAGTGGACCGCTAGACCAATGTGTTCTGTGATGACCTGCACTAGGAAGACGAATAGGTGGCCTA  
 TGGCATTATTGTACGGAATGATAAACATTGCTGCATAAATTCTTTATTATACAGCCATAATGTCAGTA  
 GCAAGGGAGAAAAGGTTCAAAGTCGAAAAAATTATGAGAAACCTTACATGAGCCTGACGTACGTT  
 45 TGCCTAAGCGTTAGAAGCTCCTACTTGAAGAGATATTGCGCGATAATATCTCTAATATTGCAAATG  
 AAGTGCCTGGTACATCAGATGACAGTACTGAAGAGCCAGTAATGAAAAACGTAATTACTGTACTTACTGCC  
 CCTCTAAAATAAGCGAAAGGCAAATGCATCGTCAAAAGTCAAAAGTTATTGTCGAGAGCATAATA  
 TTGATATGTGCCAAAGTTGTTAGATCCAGCACAGTGGCGCCGCTCGAGTCTAGAGGGCCGTTAAACC  
 50 CGCTGATCAGCCTCGACTGTGCCTCTAGTTGCCAGCCATCTGTTGCCCCCTCCCCGTGCCCTCCTTG  
 ACCCTGGAAAGGTGCCACTCCACTGTCTTCTAATAAAATGAGGAAATTGCATCGCATTGTCAGTAGG  
 TGTCATTCTATTCTGGGGGTGGGGCAGGACAGCAAGGGGGAGGATTGGAAAGACAATAGCAGGCAT  
 55 GCTGGGGATGCGGTGGCTATGGCTCTGAGGCGAAAGAACAGCTGGGCTCTAGGGGTATCCCCAC  
 GCGCCCTGTAGCGCGCATTAGCGCGCGGGTGTGGTACGCGCAGCGTACCGCTACACTGCCAGC

GCCCTAGCGCCCGCTCTTCGCTTCTTCCCTTCGCCACGGTCGCCGGCTTCCCCGTCAAGCT  
 CTAAATCGGGGCATCCCTTAGGGTTCCGATTAGTGCTTACGGCACCTGACCCAAAAACTGATTAG  
 GGTGATGGTTACGTAGTGGCCATGCCCTGATAGACGGTTTCGCCCTTGACGGAGTCCACGTT  
 5 TTTAATAGTGGACTCTGTTCCAAACTGGAACAACACTCAACCCATCTCGGTCTATTCTTTGATTATAA  
 GGGATTTGGGGATTTOGGCCTATTGGTAAAAATGAGCTGATTAACAAAATTAACGCGAATTAAATTC  
 TGTGAAATGTGTGTCAGTTAGGGTGTGAAAGTCCCCAGGCCTCCCAGCAGGAGAAGTATGCAAAGCATG  
 10 CATCTCAATTAGTCAGCAACCAGGTGTGAAAGTCCCCAGGCCTCCCAGCAGGAGAAGTATGCAAAGCATG  
 CATCTCAATTAGTCAGCAACCATAGTCCC GCCCTAACCTCCGCCATCCGCCCTAACCTCCGCCAGTCC  
 GCCCATTCTCCGCCCATGGCTGACTAATTTTTATTATGAGAGGCCAGGGCCCTGCCTCTGAG  
 15 CTATTCCAGAAGTAGTGAGGAGGCTTTGGAGGCCTAGGCTTGCAAAAGCTCCGGAGCTGTATA  
 TCCATTTCGGATCTGATCAGCACGTGATGAAAAGCCTGAACCTACCGCGACGCTGTGAGAAGTTCTG  
 ATCGAAAAGTCGACAGCGTCTCGACCTGATGCAGCTCTGGAGGGCGAAGAATCTGTGCTTCAGCTTC  
 GATGTAGGAGGGCGTGGATATGTCCTGCGGGTAAATAGCTGCGCCGATGGTTCTACAAAGATCGTTATGTT  
 20 TATCGGCACTTGCATCGGCCGCGCTCCGATTCCGAAAGTGCTTGACATTGGGAATTCAAGCAGAGCCTG  
 ACCTATTGCATCTCCGCCGTGCACAGGGTGTACGTTGCAAGACCTGCTGAAACCGAACTGCCGCTGTT  
 CTGCAGCCGGTCCGGAGGCCATGGATGCGATCGTGCAGGCGATCTAGCCAGACGAGCAGGGTCGGCCA  
 TTCGGACCGCAAGGAATCGGTCAATACACTACATGGCGTGAATTGATATGCGCATTGCTGATCCCCATGTG  
 25 TATCACTGGCAAACACTGTGATGGACGACACCGTCAGTGCCTCGCAGGCTCTCGATGAGCTGATGCTT  
 TGGCCGAGGACTGCCCGAAGTCCGGCACCTCGTCACGCGGATTGGCTCAACATGCTCTGACGGAC  
 AATGGCCGCATAACAGCGGTCACTGGAGCGAGGCGATGGCTGGGGATTCCAATACGAGGTCGCCAAC  
 30 ATCTTCTCTGGAGGCCGTGGTTGGCTTGATGGAGCAGCAGACGCGCTACTTCGAGCGGAGGCATCGGAG  
 CTTGCAGGATCGCCGCGCTCCGGCGTATATGCTCCGCATTGGCTTGACCAACTCTATCAGAGCTGGTT  
 GACGGCAATTGATGATGCAGCTGGCGCAGGGTCGATGCGACGCAATGCTCCGATCCGGAGCCGGACT  
 GTCGGGCGTACACAAATOGCCCGCAGAAGCGCGGCCGTGACCGATGGCTGTAGAAGTACTGCCGAT  
 35 AGTGGAAACCGACGCCAGCACTCGTCCGAGGGAAAGGAATGACACGTGCTACGAGATTGATCCACC  
 GCCGCCTCTATGAAAGGTTGGCTCGGAATCGTTCCGGACGCCGCTGGATGATCCTCAGCGCGG  
 GATCTCATGCTGGAGTCTCGCCCACCCAACTGTTATTGAGCTTACGCTATAATGGTACAAATAAGCAAT  
 40 AGCATCACAAATTTCACAAATAAGCATTGACTGCTAGAGCTGGCGTAATCATGGTCAAGCTGTTCT  
 GTGTGAAATTGTTATCCGCTCACAAATTCCACACAAACATACGAGCCGGAAGCATAAAGTGTAAAGCCTGGGT  
 GCCTAATGAGTGGACTACACATTAATTGCGCTCGCTACTGCCGCTTCCAGTCGGAAACCTGTCG  
 45 TGCCAGCTGCATTAATGAACTGGCAACGCCGGGGAGAGGGGTTGGCTATTGGCGCTCTCCGCTTCC  
 TCGCTCACTGACTCGCTCGCCTCGGTGCGCTGCGCGAGCGGTATCAGCTCAACTCAAAGCGGTAATA  
 CGGTTATCCACAGAATCAGGGATAACGCAGGAAGAACATGTGAGCAAAAGGCCAGCAAAAGGCCAGGAAC  
 CGTAAAAGGCCGCTGGCTGGCTTTCCATAGGCTCCGCCCTGACGAGCATCACAAAATGACGC  
 50 TCAAGTCAGAGGTTGGCGAAACCGACAGGACTATAAAGATACCGAGGCTTCCCGTGGAGCTCCCTCGT  
 CGCTCTCTGTTCCGACCTGCCGCTTACGGATACCTGTCGCCCTTCTCCCTCGGAAGCGTGGCGCTT  
 TCTCAATGCTCACGCTGTAGGTATCTCAGTCGGTAGGTCGCTCCAGCTGGCTGTGACGAA  
 55 CCCCCCGTTAGCCGACCGCTGCCCTATCGGTAACTATCGTCTTGAGTCCAACCGGTAAAGACACGAC  
 TTATGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTAGGCGGTGCTACAGAGTTC

TTGAAGTGGTGGCCTAACTACGGCTACACTAGAAGGACAGTATTGGTATCTGCCTCTGCTGAAGCCAGTT  
 5 ACCTTCGGAAAAAGAGTTGGTAGCTCTGATCCGGCAAACAAACCACCGCTGGTAGCGGTGGTTTTTGTGTT  
 TGCAAGCAGCAGATTACGCCAGAAAAAAAGGATCTAAGAAGATCCTTGATCTTCTACGGGTCTGAC  
 GCTCAGTGGAACGAAAACACGTTAACGGGATTGGCATGAGATTATCAAAAGGATCTCACCTAGATC  
 CTTTAAATTAAAAATGAAGTTAAATCAATCTAAAGTATATGAGTAAACTGGTCTGACAGTTACCAA  
 TGCTTAATCAGTGAGGCACCTATCTCAGCGATCTGCTATTGCTCATCCATAGTTGCCTGACTCCCCGTC  
 GTGTAGATAACTACGATACGGGAGGGCTTACCATCTGGCCCAGTGCTGCAATGATACCGCGAGACCCACGC  
 10 TCACCGGCTCCAGATTATCAGCAATAAACCAGCCAGCCGAAGGGCCAGCGCAGAAGTGGCCTGCAACT  
 TTATCCGCTCCATCCAGTCTATTAAATTGTTGCCGGAAAGCTAGAGTAAGTAGTCGCCAGTTAATAGTTG  
 CGCAACGTTGTTGCCATTGCTACAGGCATCGTGGTGTACGCTCGTCTGGTATGGCTTCATTAGCTCC  
 15 GGTTCCTAACGATCAAGGCAGTTACATGATCCCCATGTTGCAAAAAAGCGGTTAGTCCTTCGGTCT  
 CCGATCGTTGTCAGAAGTAAGTTGCCGCAGTGTATCACTCATGGTTATGGCAGCACTGCATAATTCTCTT  
 ACTGTCATGCCATCCGTAAGATGCTTCTGTGACTGGTGAGTACTCAACCAAGTCATTCTGAGAATAGTGT  
 20 ATGCGGCACCGAGTTGCTCTGCCGGTCAATACGGATAATACCGGCCACATAGCAGAACTTAAAA  
 GTGCTCATCATTGAAAACGTTCTCGGGCGAAAACCTCAAGGATCTTACCGCTGTTGAGATCCAGTTG  
 ATGTAACCCACTCGTGCACCCAACTGATCTTCAGCATCTTCTACCGCTGTTCTGGGTGAGCAAAA  
 25 ACAGGAAGGCAAAATGCCGAAAAAAGGAATAAGGGCACACGGAAATGTTGAATACTCATACTCTTC  
 TTTCAATATTATTGAAGCATTATCAGGGTATTGCTCATGAGCGGATACATATTGAATGTATTTAGAAA  
 AATAAACAAATAGGGTTCCGCGCACATTCCCGAAAAGTGCCACCTGACGTC - 3'

[0124] With the completion of Example 4, all genetic constructs are available to realize the method disclosed herein.

**Example 5:** Cloning of transposable light chain expression vector for human antibody kappa light chains compatible with the *Sleeping Beauty* Transposase enzyme

[0125] In order to transpose human immunoglobulin heavy and light chain expression vectors contained in a transposable vector independently into host cells, a transposable immunoglobulin light chain construct with different inverted terminal repeat (ITR) sequences can be constructed that are recognized by the *Sleeping Beauty* transposase.

[0126] For this, the human Ig-kappa light chain expression vector pIRES-EGFP-T1T2-IgL (Seq-ID-5) of example 1 can be used to replace the 5' and 3' ITRs of the *PiggyBac* transposon system, contained in this vector, with the 5' and 3' ITRs of the *Sleeping Beauty* transposon system. The sequences for the *Sleeping Beauty* 5'ITR and 3'ITR, recognized and functional with the *Sleeping Beauty* transposase, can be retrieved from patent document US7160682B1.

[0127] The upstream *Sleeping beauty* ITR sequence with the 5' terminal repeat has to be gene synthesized with flanking MunI restriction enzyme sequences, allowing the replacement of the MunI flanked *PiggyBac* 5'ITR in construct pIRES-EGFP-T1T2-IgL (Seq-ID-5) of example 1 by the *Sleeping beauty* 5'ITR sequence. This sequence is as follows (Mun I restriction enzyme sites are highlighted in boldface print and 4 additional flanking random nucleotides, allowing proper restriction enzyme digestion of the gene synthesized fragment, are indicated in lowercase letters):

Seq-ID14:

5'-

50 atatCAATTGAGTTGAAAGTCGGAAGTTACATACACTTAAGTGGAGTCATTAA  
 AACTCGTTTCAACTACACCACAAATTCTGTTAACAAACAATAGTTGG  
 55 CAAGTCAGTTAGGACATCTACTTGTGCATGACACAAGTCATTTCACAA  
 TTGTTACAGACAGATTATTCACTTATAATTCACTGTATCACAATTCCAGTGG  
 GTCAGAAGTTACATACACTAACAATTGatat-3'

[0128] The downstream *Sleeping beauty* ITR sequence with the 3' terminal repeat (also published in US7160682B1) has to be gene synthesized with flanking Xhol restriction enzyme sequences, allowing the replacement of the Xhol flanked *PiggyBac* 3'ITR in construct pIRES-EGFP-T1T2-IgL (Seq-ID-5) of example 1 by the *Sleeping beauty* 3'ITR sequence. This sequence is as follows (Xhol restriction enzyme sites are highlighted in boldface print and 4 additional flanking random nucleotides, allowing proper restriction enzyme digestion of the gene synthesized fragment, are indicated in lowercase letters):

5 Seq-ID15:

10 5'-

15 atat**CTCGAG**TGAGTGTATGTTAACTTCTGACCCACTGGGAATGTGATGAAAG  
TCTTAAAATAAAAGTGGTGATCCTAACTGACCTTAAGACAGGGAATCTTACTC  
GGATTAAATGTCAGGAATTGTGAAAAAGTGAGTTAAATGTATTGGCTAAG  
GTGTATGTAAACTCCGACTTCAACT**CTCGAG**atat-3'

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[0129] In a first step, the MunI-flanked *PiggyBac* 5'ITR of construct pIRES-EGFP-T1T2-IgL (Seq-ID-5) has to be replaced by the *Sleeping Beauty* 5'ITR by digesting pIRES-EGFP-T1T2-IgL (Seq-ID-5) with MunI restriction enzyme and by ligating the MunI digested gene-synthesized fragment from Seq-ID-14 into the MunI linearized vector backbone of pIRES-EGFP-T1T2-IgL (Seq-ID-5). The proper replacement and correct orientation of *Sleeping Beauty* 5'ITR can be checked by diagnostic restriction enzyme digestions and/or DNA sequencing. The resulting plasmid is called pIRES-EGFP-sbT1-pbT2-IgL (Fig. 8).

[0130] In a second step, the Xhol-flanked *PiggyBac* 3'ITR of construct still contained in pIRES-EGFP-sbT1-pbT2-IgL has to be replaced by the *Sleeping Beauty* 3'ITR by digesting pIRES-EGFP-sbT1-pbT2-IgL with Xhol restriction enzyme and by ligating the Xhol digested gene-synthesized fragment from Seq-ID-15 into the Xhol linearized vector backbone of pIRES-EGFP-sbT1-pbT2-IgL. The proper replacement and correct orientation of *Sleeping Beauty* 3'ITR can be checked by diagnostic restriction enzyme digestions and/or DNA sequencing. The resulting plasmid is called pIRES-EGFP-sbT1T2-IgL (Fig. 8).

[0131] The entire sequence of the human Ig-kappa LC expression vector pIRES-EGFP-sbT1T2-IgL transposable by the *Sleeping Beauty* transposase is provided in Seq-ID-16:

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Seq-ID16:

5'-

5 GACGGATCGGGAGATCTCCGATCCCCTATGGTCGACTCTCAGTACAATCTGC  
TCTGATGCCGCATAGTTAACGCCAGTATCTGCTCCCTGCTTGTGTGGAGGT  
CGCTGAGTAGTGCGCGAGCAAAATTAAAGCTACAACAAGGCAAGGCTTGACC  
10 GACAATTGAGTTGAAGTCGGAAGTTACATACACTTAAGTTGGAGTCATTAA  
AACTCGTTTCAACTACACCAACAAATTCTGTTAACAAACAATAGTTTGG  
15 CAAGTCAGTTAGGACATCTACTTGTGCATGACACAAGTCATTTCACAA  
TTGTTACAGACAGATTATTCACTTATAATTCACTGTATCACAATTCCAGTGG  
GTCAGAAGTTACATACACTAACAAATTGCATGAAGAATCTGCTTAGGGTTAG  
20 GCGTTTGCCTGCTCGCGATGTACGGGCCAGATATACGCGTTGACATTGAT  
TATTGACTAGTTATTAAATAGTAATCAATTACGGGTCATTAGTCATAGCCCA  
TATATGGAGTTCCCGCGTTACATAACTACGGTAAATGGCCCGCTGGCTGACC

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GCCCAACGACCCCCGCCATTGACGTCAATAATGACGTATGTTCCCAGTAGTAA  
CGCCAATAGGGACTTCCATTGACGTCAATGGGTGGACTATTTACGGTAAACT  
5     GCCCACTTGGCAGTACATCAAGTGATCATATGCCAAGTACGCCCTATTGA  
CGTCAATGACGGTAAATGGCCCGCCTGGCATTATGCCAGTACATGACCTTAT  
GGGACTTTCTACTTGGCAGTACATCTACGTATTAGTCATCGCTATTACCATG  
10    GTGATGCGGTTTGGCAGTACATCAATGGCGTGGATAGCGGTTGACTCACG  
GGGATTTCAGTCTCCACCCATTGACGTCAATGGGAGTTGTTGGCACC  
15    AAAATCAACGGGACTTCAAAATGTCGTAACAACACTCCGCCATTGACGCA  
AATGGGCGGTAGGCGTGTACGGTGGGAGGTCTATATAAGCAGAGCTCTGG  
CTAACTAGAGAACCACTGCTTACTGGCTTATCGAAATTAAATACGACTCACTA  
20    TAGGGAGACCCAAGCTTGGTACCGAGCTGGATCGAtggacatgagggtccctgctag  
ctcctggactcctgctgctggctccagggtgccagatgtGACATCCAGATGACCCAGTCTCCATCC  
TCCCTGTCTGCATCTGTAGGGGACAGAGTCACCACACTGTCGGCAAGTCA  
25    GGGCATCAGAAATTACTTAGCCTGGTATCAGCAAAAACCAGGGAAAGCCCT  
AAGCTCCTGATCTATGCTGCATCCACTTGCAATCAGGGTCCATCTCGTT  
CAGTGGCAGTGGATCTGGACAGATTCACTCTCACCACAGCAGCCTACAG  
30    CCTGAAGATGTTGCAACTTATTACTGTCAAAGGTATAACCGTGCACCGTATAC  
TTTGGCCAGGGACCAAGGTGGAAATCAAGCGCTCTGTGGCTGCACCATCT  
GTCTTCATCTCCGCCATCTGATGAGCAGTTGAAATCTGAACTGCCTCTGT  
35    TGTGTGCCTGCTGAATAACTCTATCCCAGAGAGGCCAAAGTACAGTGGAAAG  
GTGGATAACGCCCTCCAATCGGTAACTCCCAGGAGAGTGTACAGAGCAGG  
ACAGCAAGGACAGCACCTACAGCCTCAGCAGCACCCGTACGCTGAGCAAAGC  
40    AGACTACGAGAAACACAAAGTCTACGCCTGCGAAGTCACCCATCAGGGCTG  
AGCTGCCCGTCACAAAGAGCTTCAACAGGGGAGAGTGTAAATCTGCGGCC  
GCGTCGACGGAATTCACTGGATCCACTAGTAACGGCCGCCAGTGTGCTGGAA  
45    TTAATTGCTGTGCGAGGGCCAGCTGTTGGGTGAGTACTCCCTCTCAAAA  
GCGGGCATGACTCTGCGCTAACAGATTGTCAGTTCAAAACGAGGAGGATT  
TGATATTCACCTGGCCCGGGTGTGCTTGTGAGGGTGGCCCGTCCATCTGG  
50    TCAGAAAAGACAATCTTTGTTGTCAAGCTGAGGTGTGGCAGGCTTGAGAT  
CTGGCCATACACTGAGTGACAATGACATCCACTTGCCCTCTCCACAGG  
TGTCCACTCCCAGGTCCAACTCAGGTCGAGCATGCATCTAGGGCGGCCAATT  
55    CCGCCCTCTCCCTCCCCCCCCCTAACGTTACTGGCGAAGCCGCTTGGAAAT

AAGGCCGGTGTCCGTTGTCATATGTGATTTCACCATATTGCCGTCTTG  
GCAATGTGAGGGCCCGAACCTGCCCTGCTTCTGACGAGCATTCTAGG  
5 GGTCTTCCCCCTCTGCCAAAGGAATGCAAGGTCTGTTGAATGTCGTGAAGGA  
AGCAGTTCCTCTGGAAGCTTCTGAAGACAAACAACGTCTGACCGCCCTT  
GCAGGCAGCGAACCCCCCACCTGGCGACAGGTGCCTCTGCGGCCAAAAGCC  
10 ACGTGTATAAGATAACACCTGCAAAGCGGCACAACCCCAGTGCCACGTTGTG  
AGTTGGATAGTTGTGGAAAGAGTCAAATGGCTCTCCTCAAGCGTATTCAACA  
AGGGGCTGAAGGATGCCAGAAGGTACCCATTGTATGGATCTGATCTGGG  
15 GCCTCGGTGCACATGCTTACATGTGTTAGTCGAGGTTAAAAAAACGTCTAG  
GCCCGCGAACACCGGGACGTGGTTCTTGAACACGATGATAAGC  
20 TTGCCACAACCGGGATCCACCGGTGCCACCATGGTGAGCAAGGGCGAGGA  
GCTGTTACCAGGGTGGTGCCATCCTGGTCAGCTGGACGGCGACGTAAAC  
GGCCACAAGTTCAGCGTGTCCGGCGAGGGCGAGGGCGATGCCACCTACGGCA  
25 AGCTGACCTGAAGTTCATCTGCACCACCGCAAGCTGCCGTGCCCTGGCCC  
ACCCTCGTGACCACCTGACCTACGGCGTGCAGTGCTCAGCCGCTACCCGA  
CCACATGAAGCAGCACGACTTCTCAAGTCCGCCATGCCGAAGGCTACGTC  
30 CAGGAGCGCACCATCTTCTCAAGGACGACGGCAACTACAAGACCCGCGCCG  
AGGTGAAGTCGAGGGCGACACCCCTGGTAACCGCATCGAGCTGAAGGGCAT  
CGACTTCAAGGAGGACGGCAACATCCTGGGGACAAGCTGGAGTACAAC  
35 AACAGCCACAACGTCTATATCATGGCCGACAAGCAGAAGAACGGCATCAAGG  
TGAACCTCAAGATCCGCCACAACATCGAGGACGGCAGCGTGCAGCTGCCGA  
CCACTACCAGCAGAACACCCCCATCGCGACGGCCCCGTGCTGCTGCCGAC  
40 AACCACTACCTGAGCACCCAGTCCGCCCTGAGCAAAGACCCAACGAGAAGC  
GCGATCACATGGCCTGCTGGAGTTGTCAGCCGCCGGGATCACTCTCGGC  
ATGGACGAGCTGTACAAGTAAAGCGGCCCTAGAGCTCGCTGATCAGCCTCGA  
45 CTGTGCCCTAGTTGCCAGCCATCTGTTGCTGCCCTCCCCGTGCCCTCCTT  
GACCCCTGGAAGGTGCCACTCCACTGTCCTTCTTAATAAAATGAGGAAATTG  
CATCGCATTGTCTGAGTAGGTGTCATTCTATTCTGGGGGTGGGTGGGGCAG  
50 GACAGCAAGGGGGAGGATTGGAAGACAATAGCAGGCATGCTGGGATGCG  
GTGGGCTCTATGGCTCTGAGGCGGAAAGAACCAAGCTGGGCTCGAGTTGAG  
TGTATGTTAACTCTGACCCACTGGGAATGTGATGAAAGAAATAAAAGCTGA  
55 AATGAATCATTCTCTACTATTATTCTGATATTACACATTCTAAAATAAGT

GGTGATCCTAACTGACCTAACAGACAGGAACTTACTCGGATTAAATGTCAG  
GAATTGTAAAAAGTGAGTTAAATGTATTGGCTAAGGTGTATGTAAACTC  
5 CGACTTCAACTCTCGAGTGCATTCTAGTTGGTTGTCCAAACTCATCAATG  
TATCTTATCATGTCTGTATAACCGTCGACCTCTAGCTAGAGCTGGCGTAATCA  
TGGTCATAGCTGTTCCGTGAAATTGTTATCCGCTCACAAATTCCACACAA  
10 CATACTCACATTAATTGCGTTGCGCTCACTGCCGCTTCCAGTCGGAAACCT  
GTCGTGCCAGCTGCATTAATGAATCGGCCAACGCGCGGGAGAGGCGGTTG  
15 CGTATTGGCGCTCTCCGCTTCGCTCACTGACTCGCTGCGCTCGTCGTT  
CGGCTGCGCGAGCGGTATCAGCTCACTCAAAGGCGGTAAACGGTTATCCA  
CAGAATCAGGGATAACGCAGGAAAGAACATGTGAGCAAAAGGCCAGCAA  
20 AGGCCAGGAACCGTAAAAAGGCCGTTGCTGGCGTTTCCATAGGCTCCG  
CCCCCCTGACGAGCATCACAAAATCGACGCTCAAGTCAGAGGTGGCGAAAC  
25 CCGACAGGACTATAAAGATACCAGGCCTTCCCCCTGGAAGCTCCCTCGTGC  
GCTCTCCTGTTCCGACCCCTGCCGCTTACCGGATACCTGTCCGCTTCTCCCT  
CGGAAGCGTGGCGTTCTCAATGCTCACGCTGTAGGTATCTCAGTCGGTG  
30 TAGTCGTTCGCTCCAAGCTGGCTGTGCACGAACCCCCGTTAGCCGAAAC  
CCGCTGCGCCTTATCCGTAACTATCGTCTGAGTCCAACCCGTAAGACACG  
ACTTATGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTA  
35 TGTAGGCGGTGCTACAGAGTTCTGAAGTGGTGGCCTAACTACGGCTACACTA  
GAAGGACAGTATTGGTATCTGCGCTTGCTGAAGCCAGTTACCTTCGGAAAA  
AGAGTTGGTAGCTCTGATCCGGAAACAAACCACCGCTGGTAGCGGTGGTT  
40 TTTTGGTGCAGCAGATTACGCGCAGAAAAAAAGGATCTCAAGAAGA  
TCCTTGATCTTCTACGGGCTGACGCTCAGTGGAACGAAAACACTACGTT  
AAGGGATTTGGTCATGAGATTATAAAAGGATCTCACCTAGATCCTTTA  
45 AATTAAAAATGAAGTTAAATCAATCTAAAGTATATGAGTAAACTTGGTC  
TGACAGTTACCAATGCTTAATCAGTGAGGCACCTATCTCAGCGATCTGTCTAT  
TTCGTTCATCCATAGTTGCCTGACTCCCCGTGCTAGATAACTACGATAACGG  
50 GAGGGCTTACCATCTGGCCCCAGTGCTGCAATGATAACCGCGAGACCCACGCT  
CACCGGCTCCAGATTATCAGCAATAAACCAGCCAGCCGGAAAGGGCCGAGCG  
CAGAAGTGGCCTGCAACTTATCCGCCTCCATCCAGTCTATTAAATTGTTGCC  
55 GGGAAAGCTAGAGTAAGTAGTCGCCAGTTAATAGTTGCGCAACGTTGTTGCC

ATTGCTACAGGCATCGTGGTCACGCTCGTCTGGTATGGCTTCATTCA  
CTCCGGTTCCAACGATCAAGGCGAGTTACATGATCCCCATGTTGTGCAAAA  
AAGCGGTTAGCTCCTCGTCCTCCGATCGTGTAGAAGTAAGTGGCCGCA  
5 GTGTTATCACTCATGGTTATGGCAGCACTGCATAATTCTTACTGTCATGCC  
ATCCGTAAGATGCTTTCTGTGACTGGTGAAGTACTCAACCAAGTCATTCTGAG  
10 AATAGTGTATGCGCGACCGAGTTGCTCTGCCCGCGTCAATACGGATAAA  
TACCGCGCCACATAGCAGAACCTTAAAAGTGCTCATCATTGGAAAACGTTCTT  
15 CGGGCGAAAACCTCAAGGATCTTACCGCTGTTGAGATCCAGTCGATGTA  
ACCCACTCGTGCACCCAACTGATCTCAGCATCTTACTTCACCAGCGTTTC  
TGGGTGAGCAAAACAGGAAGGCAAAATGCCGAAAAAAGGGATAAGGGC  
20 GACACGGAAATGTTGAATACTCATACTCTTCAATATTATTGAAGCA  
TTTATCAGGGTTATTGTCTCATGAGCGGATACATATTGAATGTATTAGAAA  
AATAAACAAATAGGGGTTCCCGCACATTCCCCGAAAAGTGCCACCTGACG  
25 TC-3'

**Example 6:** Cloning of a *Sleeping Beauty* expression vector

[0132] The open reading frame (ORF) of the *Sleeping Beauty* transposase enzyme can also be found in patent reference US7160682B 1/US2003154500A1.  
30 [0133] This sequence is provided in Seq-ID17, below:

Seq-ID17:

35 5'-  
ATGGGAAAATCAAAGAAATCAGCCAAGACCTCAGAAAAAAAAATTGTAGAC  
CTCCACAAGTCTGGTCATCCTGGAGCAATTCCAAACGCCTGAAAGTACC  
40 ACGTTCATCTGTACAAACAATAGTACGCAAGTATAAACACCATGGGACCACG  
CAGCCGTATACCGCTCAGGAAGGAGACCGTTCTGTCTCCTAGAGATGAAC  
GTACTTGGTGCAGAAAGTCAAATCAATCCCAGAACACAGCAAAGGACCT  
45 TGTGAAGATGCTGGAGGAAACAGGTACAAAGTATCTATATCCACAGTAAAA  
CGAGTCCTATATCGACATAACCTGAAAGGCCGCTCAGCAAGGAAGAAGCCAC  
TGCTCCAAAACCGACATAAGAAAGCCAGACTACGGTTGCAACTGCACATGG  
50 GGACAAAGATCGTACTTTGGAGAAATGTCCTCTGGTCTGATGAAACAAAAA

ATAGAACTGTTGCCATAATGACCATCGTTATGTTGGAGGAAGAAGGGGG  
AGGCTTGCAAGCCGAAGAACACCATCCCAACCGTGAAGCACGGGGTGGCA  
5 GCATCATGTTGTGGGGTGCTTGCTGCAGGAGGGACTGGTGCACTCACAA  
AATAGATGGCATCATGAGGAAGGAAAATTATGTGGATATATTGAAGCAACAT  
CTCAAGACATCAGTCAGGAAGTTAAAGCTTGGTCGCAAATGGGTCTCCAAA  
10 TGGACAATGACCCAAGCATACTTCAAAGTTGTGGCAAAATGGCTTAAGGA  
CAACAAAGTCAAGGTATTGGAGTGGCCATCACAAAGCCCTGACCTCAATCCT  
15 ATAGAAAATTGTGGCAGAACTGAAAAAGCGTGTGCGAGCAAGGAGGCCT  
ACAAACCTGACTCAGTTACACCAGCTCTGTCAGGAGGAATGGGCCAAATTC  
ACCCAACCTATTGTGGAGCTTGTGGAAGGCTACCCGAAACGTTGACCCA  
20 AGTTAAACAATTAAAGGCAATGCTACCAAATACTAG-3'

[0134] This sequence translates into the following amino acid sequence:

Seq-ID18:

25 MGKSKEISQDLRKKIVDLHKSGSSLGAISKRLKVPRSSVQTIVRKYKHHGT  
TQPSYRSGRRRVLSPRDERTLVRKVQINPRTTAKDLVKMLEETGTKVSISTVKRV  
LYRHNLKGRSARKKPLLQNRHKKARLRFATAHGDKDRTFWRNVLWSDETKIEL  
30 FGHNDHRYVWRKKGEACKPKNTIPTVKHGGGSIMLWGCFAAGGTGALHKIDGI  
MRKENYVDILKQHLKTSVRKLKLGRKWVFQMDNDPKHTSKVVAKWLKDNKV  
KVLEWPSQSPDLNPIENLWAELKKVRARRPTNLTQLHQLCQEEWAKIHPTYCG  
35 KLVEGYPKRLTQVKQFKGNATKY•

[0135] In order to generate a vertebrate cell expression vector for the *Sleeping Beauty* transposase enzyme, this ORF can be gene synthesized and cloned as a blunt ended DNA into the unique, blunt-cutting restriction enzyme site EcoRV 40 in the standard vertebrate cell expression vector pCDNA3.1-hygro(+) (catalogue # V870-20, Invitrogen, Carlsbad, CA, USA), by methods known in the art. The correct ligation of the *Sleeping Beauty* ORF, relative to the pCDNA3 promoter can be verified by diagnostic restriction enzyme digestion, and/or by DNA sequencing of the cloned *Sleeping Beauty* expression construct pCDNA3.1-hygro(+)-SB (Fig. 9).

[0136] The sequence of the *Sleeping Beauty* expression construct pCDNA3.1-hygro(+)-SB is provided in Seq-ID 19, 45 below:

Seq-ID19:

5'-

5 GACGGATCGGGAGATCTCCGATCCCCTATGGTCGACTCTCAGTACAATCTGC  
10 TCTGATGCCGCATAGTTAACGCCAGTATCTGCTCCCTGCTTGTGTGGAGGT  
15 CGCTGAGTAGTGCGCGAGCAAAATTAAAGCTACAACAAGGCAAGGCTTGACC  
20 GACAATTGCATGAAGAATCTGCTTAGGGTTAGGCCTTGCGCTGCTCGCGA  
25 TGTACGGGCCAGATATACCGCGTTGACATTGATTATTGACTAGTTATTAAAGT  
30 AATCAATTACGGGTCATTAGTTCATAGCCCATAATGGAGTTCCCGTACA  
35 TAACTTACGGTAAATGGCCCGCTGGCTGACCGCCAACGACCCCCGCCATT  
40 GACGTCAATAATGACGTATGTTCCATAGTAACGCCAATAGGGACTTCCATT  
45 GACGTCAATGGGTGGACTATTACGGTAAACTGCCACTTGGCAGTACATCA  
50 AGTGTATCATATGCCAAGTACGCCCTATTGACGTCAATGACGGTAAATGGC  
55 CCGCCTGGCATTATGCCAGTACATGACCTTATGGGACTTCCACTTGGCAG  
60 TACATCTACGTATTAGTCATCGCTATTACCATGGTATGCGGTTTGGCAGTA  
65 CATCAATGGCGTGGATAGCGGTTGACTCACGGGATTCCAAGTCTCCACC  
70 CCATTGACGTCAATGGAGTTGTTGGCACCAAATCAACGGACTTCCA  
75 AAATGTCGTAACAACCTCCGCCATTGACGCAAATGGCGGTAGGCGTGTAC  
80 GGTGGGAGGTCTATATAAGCAGAGCTCTGGCTAACTAGAGAACCCACTGC  
85 TTACTGGCTTATCGAAATTAAATACGACTCACTATAGGGAGACCCAAGCTGGCT  
90 AGCGTTAAACTTAAGCTTGGTACCGAGCTCGGATCCACTAGTCCAGTGTGGT  
95 GGAATTCTGCAGATATGGAAAATCAAAGAAATCAGCCAAGACCTCAGAA  
100 AAAAAATTGTAGACCTCCACAAGTCTGGTTCATCCTGGAGCAATTCCA  
105 CGCCTGAAAGTACCACTGTCATCTGTACAAACAATAGTACGCAAGTATAAAC  
110 ACCATGGGACCACGCAGCCGTACCGCTCAGGAAGGAGACGCCTCTGTC  
115 TCCTAGAGATGAACGTACTTGGTGCAGAAAGTGCACAAATCAATCCCAGAAC  
120 ACAGCAAAGGACCTGTGAAGATGCTGGAGGAAACAGGTACAAAGTATCTA  
125 TATCCACAGTAAAACGAGTCCTATATCGACATAACCTGAAAGGCCGCTCAGC  
130 AAGGAAGAAGCCACTGCTCCAAAACCGACATAAGAAAGCCAGACTACGGTT  
135 GCAACTGCACATGGGGACAAAGATCGTACTTTGGAGAAATGTCCTCTGGTC  
140 TGATGAAACAAAAATAGAACTGTTGCCATAATGACCATCGTTATGTTGGA  
145 GGAAGAAGGGGGAGGCTTGCAAGCCGAAGAACACCATCCCAACCGTGAAGC  
55

ACGGGGTGGCAGCATCATGTTGTGGGGTGCTTGCTGCAGGAGGGACTGG  
TGCACCTCACAAAATAGATGGCATCATGAGGAAGGAAAATTATGTGGATATA  
5 TTGAAGCAACATCTCAAGACATCAGTCAGGAAGTAAAGCTTGGTCGCAAAT  
GGGTCTTCAAATGGACAATGACCCCAAGCATACTTCAAAGTTGTGGCAA  
10 ATGGCTTAAGGACAACAAAGTCAAGGTATTGGAGTGGCCATCACAAAGCCCT  
GACCTCAATCCTATAGAAAATTGTGGCAGAACTGAAAAAGCGTGTGCGAG  
CAAGGAGGCCTACAAACCTGACTCAGTTACACCAGCTGTCAAGGAGGAATG  
15 GGCCAAAATTCAACCAACTTATTGTGGGAAGCTTGTGGAAGGCTACCCGAAA  
CGTTGACCCAAGTAAACAATTAAAGGCAATGCTACCAAATACTAGATCC  
AGCACAGTGGCGGCCGCTCGAGTCTAGAGGGCCGTTAAACCCGCTGATCA  
20 GCCTCGACTGTGCCTTAGTTGCCAGCCATCTGTTGTTGCCCTCCCCGTG  
CCTTCCTGACCTGGAAAGGTGCCACTCCACTGTCCTTCCTAATAAAATGA  
GGAAATTGCATCGCATTGTCTGAGTAGGTGTCATTCTATTCTGGGGGTGGGG  
25 TGGGGCAGGACAGCAAGGGGAGGATTGGGAAGACAATAGCAGGCATGCTG  
GGGATGCGGTGGGCTCTATGGCTCTGAGGCGGAAAGAACAGCTGGGCTC  
TAGGGGGTATCCCCACGCGCCCTGTAGCGGCGCATTAGCGCGCGGGTGTG  
30 GTGGTTACGCGCAGCGTACCGCTACACTGCCAGCGCCCTAGCGCCCGCTCC  
TTTCGCTTCTCCCTTCTCGCCACGTTGCCGGCTTCCCCGTCAAGCT  
CTAAATCGGGGCATCCCTTAGGGTCCGATTAGTGTCTTACGGCACCTCGA  
35 CCCAAAAAAACTGATTAGGGTATGGTCACGTAGTGGCCATGCCCTGAT  
AGACGGTTTCGCCCTTGACGTTGGAGTCCACGTTCTTAATAGTGGACTCT  
TGTTCCAAACTGGAACAACACTCAACCCATCTCGGTCTATTCTTTGATTAT  
40 AAGGGATTTGGGATTCGCCATTGGTAAAAATGAGCTGATTAAACAA  
AAATTAAACGCGAATTAAATTCTGTGGAATGTGTGTCAGTTAGGGTGTGGAAA  
GTCCCCAGGCTCCCCAGGCAGGCAGAAGTATGCAAAGCATGCATCTCAATT  
45 GTCAGCAACCAGGTGTGGAAAGTCCCCAGGCTCCCCAGCAGGCAGAAGTATG  
CAAAGCATGCATCTCAATTAGTCAGCAACCATAGTCCCAGGCTTAACCCGCC  
CATCCCGCCCTAACTCCGCCAGTCCGCCATTCTCCGCCATGGCTGAC  
50 TAATTTTTTATTATGCAGAGGCCAGGCCCTGCTGAGCTATTCC  
AGAAGTAGTGAGGAGGCTTTGGAGGCCTAGGCTTGTCAAAAGCTCCC  
GGGAGCTGTATATCCATTTCGGATCTGATCAGCACGTGATGAAAAAGCCTG  
55 AACTCACCGCGACGTCTGCGAGAAGTTCTGATCGAAAAGTCGACAGCGT

CTCCGACCTGATGCAGCTCTCGGAGGGCGAAGAACATCGTGTTCAGCTTCG  
ATGTAGGAGGGCGTGGATATGTCCTGCGGGTAAATAGCTGCGCCGATGGTT  
5 CTACAAAGATCGTTATGTTATCGGCACTTGCATCGGCCGCGCTCCGATTG  
CGGAAGTGCTTGACATTGGGAATTCAAGCAGAGCCTGACCTATTGCATCTCC  
CGCCGTGCACAGGGTGTACGTTGCAAGAACCTGCCTGAAACCGAACTGCCCG  
10 CTGTTCTGCAGCCGGTCGCGGAGGCCATGGATGCGATCGCTGCGGCCGATCTT  
AGCCAGACGAGCGGGTTCGGCCATTGGACCGCAAGGAATCGGTCAATACA  
15 CTACATGGCGTGATTTCATATGCGCGATTGCTGATCCCCATGTGTATCACTGG  
CAAACACTGTGATGGACGACACCGTCAGTGCCTCGTCGCGCAGGCTCTCGATG  
AGCTGATGCTTGGCCGAGGAATGCCCGAAGTCCGGCACCTCGTCACGC  
20 GGATTTCGGCTCCAACAATGCTTGACGGACAATGGCCGCATAACAGCGGT  
ATTGACTGGAGCGAGGCATGTTGGGGATTCCAATACGAGGTGCCAAC  
TCTTCTTCTGGAGGCCGTGGTGGCTTGTATGGAGCAGCAGCGCTACTTC  
25 GAGCGGAGGCATCCGGAGGCTGCAGGATGCCGCCGGCTCCGGCGTATATGC  
TCCGCATTGGTCTTGACCAACTCTATCAGAGCTTGGTGAAGGCAATTGAT  
GATGCAGCTGGCGCAGGGTCGATGCGACGCAATCGTCCGATCCGGAGCCG  
30 GGACTGTCGGCGTACACAAATGCCCGCAGAACGCGGGCGTCTGGACCGA  
TGGCTGTAGAAGTACTGCCGATAGTGGAAACCGACGCCAGCACTCGT  
CCGAGGGCAAAGGAATAGCACGTGCTACGAGATTGATTCCACCGCCGCCT  
35 TCTATGAAAGGTTGGCTCGGAATCGTTCCGGACGCCGGCTGGATGATC  
CTCCAGCGCGGGATCTCATGCTGGAGTTCTCGCCCACCCAACTGTTAT  
TGCAGCTTATAATGGTTACAAATAAGCAATAGCATCACAAATTTCACAAAT  
40 AAAGCATTTCACTGCATTCTAGTTGTGGTTGTCACAAACTCATCAATGTA  
TCTTATCATGCTGTATACCGTCGACCTCTAGCTAGAGCTTGGCGTAATCATG  
GTCATAGCTGTTCTGTGAAATTGTTATCCGCTCACAAATTCCACACAACA  
45 TACGAGCCGGAAGCATAAAGTGTAAAGCCTGGGTGCTAACATGAGTGAGCTA  
ACTCACATTAATTGCGTTGCGCTCACTGCCGCTTCCAGTCGGAAACCTGT  
CGTGCCAGCTGCATTAATGAATCGGCCAACGCGGGGAGAGGCGGTTGCG  
50 TATTGGCGCTTCCGCTTCCTCGCTCACTGACTCGCTGCGCTGGTCGTTGCG  
GCTGCCAGCGGTATCAGCTCACTCAAAGCGGTAAACGGTTATCCACA  
GAATCAGGGATAACGCAGGAAAGAACATGTGAGCAAAAGGCCAGCAAAAG  
55 GCCAGGAACCGTAAAAAGGCCGTTGCTGGCGTTTCCATAGGCTCCGCC

CCCTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTGGCGAAACCCG  
ACAGGACTATAAAGATACCAGGCCTTCCCCCTGGAAGCTCCCTCGCGCTC  
5 TCCTGTTCCGACCCTGCCGCTTACCGGATACCTGTCCGCCTTCTCCCTCGGG  
AAGCGTGGCGCTTCTCAATGCTCACGCTGTAGGTATCTCAGTCGGTAGG  
TCGTTCGCTCCAAGCTGGCTGTGCACGAACCCCCCGTTCAGCCGACCGC  
10 TGCGCCTTATCCGTAACATCGTCTGAGTCCAACCCGTAAGACACGACTT  
ATCGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGAGGTATGTA  
GGCGGTGCTACAGAGTTCTGAAGTGGTGGCTAACTACGGCTACACTAGAA  
15 GGACAGTATTGGTATCTGCGCTCTGCTGAAGCCAGTTACCTCGGAAAAAGA  
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55

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**Example 7:** Generation of a *Sleeping Beauty* transposable human Ig-kappa light chain expression library

[0137] In order to generate a diverse *Sleeping Beauty* transposable DNA library encoding human antibody light chain libraries, the VL region of *Sleeping Beauty* transposable vector pIRES-EGFP-sbT1T2-IgL of Example 5 needs to be replaced with a diverse VL gene repertoire. This can be done by gene synthesizing of human VL coding regions flanked by Clal and Eco47III restriction enzyme sites, and by allowing nucleotide variations in certain HCDR and LCDR positions, as already provided in Seq-ID-10 above. The Seq-ID10 sequence is flanked by Clal and Eco47III restriction enzymes allowing directed ligation into Clal-Eco47III linearized pIRES-EGFP-sbT1T2-IgL. This way a *Sleeping Beauty* transposable DNA library encoding diverse human antibody light chain can be generated.

[0138] This way, diverse transposable DNA libraries, encoding antibody heavy and light chains on separate vectors, in which the expression of the antibody chains are transcriptionally and therefore operably linked to a green fluorescent marker protein can be generated.

[0139] Having generated *PiggyBac* transposable human IgH chain expression vectors (Example 2) and human IgH chain library expression vectors (Example 3), as well as *Sleeping Beauty* transposable expression vectors for human IgL chains (Example 5) and human IgL chain libraries (Example 7), and in addition having provided the vertebrate expression vectors for the *PiggyBac* and *Sleeping Beauty* transposases (Examples 4 and 6, respectively), allows the realization of the invention by independent transposition of immunoglobulin heavy and light chain vectors into host cells using two separate transposition systems, if followed by standard cell biology and molecular biology methods, that include the culture of vertebrate or mammalian host cells in vitro, their transfection with the above-mentioned constructs by methods known in the art, such that stable transposition of the constructs occurs and expression of the polypeptides of interest from those constructs is effected, followed by standard cell screening and separation techniques for desired binding or function of the expressed proteins, and, lastly, followed by the identification of the coding sequences from transposed constructs contained in the selected cells by standard genomic or RT-PCR methods in combination with standard or next generation DNA sequencing.

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#### SEQUENCE LISTING

[0140]

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        65   70   75   80

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Ile Ser Ser Leu Gln Pro Glu Asp Val Ala Thr Tyr Tyr Cys Gln Arg

100

105

110

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Lys Arg Ser Val Ala Ala Pro Ser Val Phe Ile Phe Pro Pro Ser Asp  
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Phe Tyr Pro Arg Glu Ala Lys Val Gln Trp Lys Val Asp Asn Ala Leu  
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## EP 2 692 865 B1

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**Transponeringsmedieret identifikation af specifik binding eller funktionelle proteiner****Patentkrav**

- 5        1. En metode til at identificere et polypeptid, der har en ønsket bindingsspecificitet eller funktionalitet, indeholdende:
- (i) generering af en forskelligartet samling polynukleotider, fortrinsvis plasmidvektorer eller dobbeltstrengede DNA PCR-amplikoner, der koder polypeptider med forskellige bindingsspecificiteter eller funktionaliteter, hvori de nævnte 10 polynukleotider indeholder en sekvens, som koder et polypeptid, der er placeret mellem en første og en anden inverteret terminalrepeatsekvens, der genkendes af og fungerer med mindst et transposaseenzym;
- (ii) introduktion af den forskelligartede samling polynukleotider af (i) i værtsellerne;
- (iii) ekspression af mindst et transposaseenzym, der fungerer med de nævnte 15 inverterede terminalrepeatsekvenser i nævnte værtseller, så nævnte forskelligartede samling af polynukleotider er integreret i værtscellegenomet for at danne en værtscellepopulation, der eksprimerer nævnte forskelligartede samling af polynukleotider, der koder polypeptider med forskellige bindingsspecificiteter eller funktionaliteter;
- (iv) screening af nævnte værtseller for at identificere en værtscelle, der eksprimerer et 20 polypeptid, der har en ønsket bindingsspecificitet eller funktionalitet; og
- (v) isolering af polynukleotidsekvenser, der koder nævnte polypeptid, mod nævnte værtscelle.
2. En metode ifølge krav 1, hvori nævnte polynukleotider indeholder
- 25     a) en ligand-bindingssekvens af en receptor eller en målbindingssekvens af et bindingsmolekyle,
- b) en antigenbindingssekvens af et antistof,
- c) en sekvens, der koder en VH- eller VL-region af et antistof eller et antigenbindingsfragment deraf,
- 30     d) en sekvens, der koder en antistof-VH-region og en antistof-VL-region,
- e) en sekvens, der koder en fuldlang tung eller let immunoglobulinkæde eller et antigenbindingsfragment deraf og/eller
- f) en sekvens, der koder et Fv- eller et Fab-enkeltkædedomæne.
- 35     3. En metode ifølge enhver af de fornævnte krav, hvori generering af en forskelligartet samling polynukleotider indeholder udsættelse af V-region-gensekvenser for PCR under mutageniserende tilstande.

4. En metode ifølge enhver af de fornævnte krav, hvori trin (ii) indeholder introduktion af nævnte værtscellepolynukleotider indeholdende sekvenser, der koder
- a) VH- eller VL-immunoglobulinregioner eller antigenbindingsfragmenter deraf og hvori nævnte VH- og VL-regionssekvenser er kodet på separate vektorer og/eller
  - 5 b) fuldlange tunge eller lette immunoglobulinkæder eller antigenbindingsfragmenter deraf, hvori nævnte fuldlange tunge og lette kædesekvenser er på separate vektorer.
  - c) en vektor, der indeholder sekvenser, som koder VH- og VL-antistofkæder,
  - d) en vektor, der indeholder sekvenser, som koder en fuldlang tung immunoglobulinkæde og en fuldlang let immunoglobulinkæde.

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5. En metode ifølge enhver af de fornævnte krav, hvori nævnte ekspressionstrin (iii) indeholder introduktion i nævnte værtsceller af en ekspressionsvektor, som koder et transposaseenzym, der genkender og fungerer med nævnte inverterede terminalrepeatsekvenser, hvori nævnte transposaseenzym fortrinsvis eksprimeres kortvarigt i nævnte værtscelle.

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6. En metode ifølge enhver af de fornævnte krav, hvori nævnte screeningstrin (iv) indeholder magnetisk aktiveret cellesortering (MACS), fluorescensaktiveret cellesortering (FACS), udvaskning mod molekyler, der er immobiliseret på en fast flade, 20 selektion til binding til cellemembranassocierede molekyler, der er inkorporeret i en cellulær, naturlig eller kunstigt rekonstitueret lipidbilagsmembran eller high-throughput screening af individuelle cellekloner i multibrøndformat for en ønsket funktions- eller bindingsfænotype.

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7. En metode ifølge enhver af de fornævnte krav, hvori nævnte trin (v) for isolering af polynukleotidsekvenser, som koder polypeptidet, der har en ønsket bindingspecificitet eller funktionalitet, indeholder genom- eller RT-PCR-amplifikation eller dyb næstegenerationssekventering.

30

8. En metode ifølge enhver af de fornævnte krav, hvori
- a) nævnte inverterede terminalrepeatsekvenser er fra PiggyBac-transposonsystemet eller Sleeping Beauty-transposonsystemet og/eller
  - b) trin (iii), der indeholder introduktion i nævnte værtscelle af en vektor indeholdende en sekvens, der koder en funktionel PiggyBac-transposase eller Sleeping Beauty-transposase.

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9. En metode ifølge enhver af de fornævnte krav, hvori nævnte inverterede terminalrepeatsekvenser genkendes af og fungerer med mindst en transposase, der er valgt

fra gruppen bestående af: PiggyBac, Sleeping Beauty, Frog Prince, Himar1, Passport, Minos, hAT, Tol1, Tol2, Ac/Ds, PIF, Harbinger, Harbinger3-DR og Hsmar1.

10. Et bibliotek af polynukleotidmolekyler, der koder polypeptider med forskellige bindingsspecificiteter eller -funktionaliteter, som indeholder et flertal af polynukleotidmolekyler, fortrinsvis plasmider eller dobbeltstrengede DNA PCR-amplikoner, hvori nævnte polynukleotidmolekyler indeholder en sekvens, der koder et polypeptid med en bindingsspecificitet eller -funktionalitet, der er placeret mellem inverterede terminalrepeatsekvenser, som er genkendt af og fungerer med mindst et transposaseenzym.

11. Et bibliotek ifølge krav 10, hvori nævnte polynukleotider indeholder  
a) mindst en sekvens, der koder en antigenbindingssekvens af et antistof.  
b) en sekvens, der koder en VH- eller VL-region af et antistof eller et  
15 antigenbindingsfragment deraf.  
c) en sekvens, der koder en antistof-VH-region og en antistof-VL-region.  
d) en sekvens, der koder en fuldlang tung eller let immunoglobulinkæde eller et antigenbindingsfragment deraf.  
e) en sekvens, der koder et Fv- eller et Fab-enkeltkædedomæne.

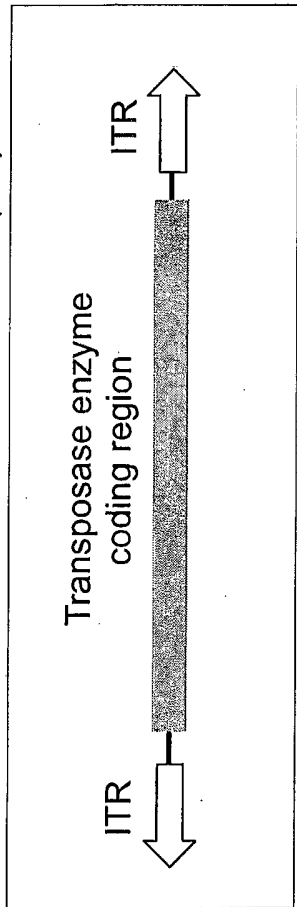
20  
12. Et bibliotek ifølge krav 10-11, hvori nævnte plasmider eller dobbeltstrengede DNA PCR-amplikoner yderligere indeholder en sekvens, der koder et transposaseenzym, som genkender og fungerer med de inverterede terminalrepeatsekvenser.

25  
13. En metode til at generere et bibliotek af transposable polynukleotider, der koder polypeptider med forskellige bindingsspecificiteter eller -funktionalitet, som indeholder generering af en forskelligartet samling polynukleotider indeholdende sekvenser, der koder polypeptider med forskellige bindingsspecificiteter eller -funktionaliteter, hvori nævnte polynukleotider indeholder en sekvens, som koder et polypeptid med en bindingsspecificitet eller -funktionalitet, der er placeret mellem inverterede terminalrepeatsekvenser, som er genkendt af og fungerer med mindst et transposaseenzym.

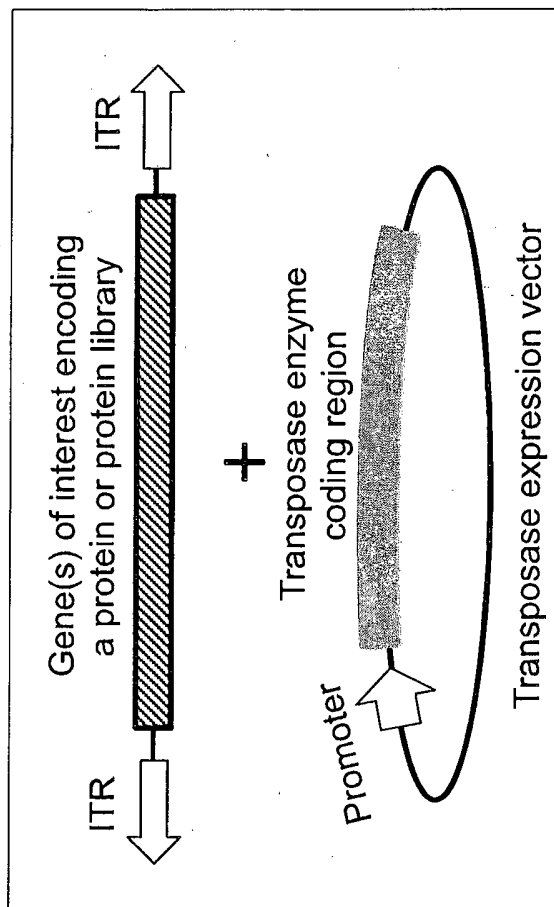
35  
14. En vektor indeholdende en sekvens, som koder en VH- eller VL-region af et antistof eller en antigenbindingsdel deraf, der er placeret mellem inverterede terminalrepeatsekvenser, som er genkendt af og fungerer med mindst et transposaseenzym.

- 15      En værtscelle indeholdende en vektor ifølge krav 14.
16.     En metode til at generere en population af værtsceller, der er i stand til at eksprimere polypeptider med forskellige bindingsspecificiteter eller –funktionaliteter  
5       indeholdende:
- (i)      generering af en forskelligartet samling polynukleotider indeholdende sekvenser, der koder polypeptider med forskellige bindingsspecificiteter eller –funktionaliteter, hvori nævnte polynukleotider indeholder en sekvens, som koder et polypeptid med en bindingsspecificitet eller -funktionalitet, der er placeret mellem 10 inverterede terminalrepeatsekvenser, som er genkendt af og fungerer med mindst et transposaseenzym; og
- (ii)     introduktion af nævnte forskelligartede samling polynukleotider i værtscellerne.
- 15      17.     Metoden eller vektoren ifølge de fornævnte krav 2-16, hvori vektoren, der indeholder VH-sekvensen, indeholder inverterede terminalrepeatsekvenser, der er genkendt af et andet transposaseenzym end de inverterede terminalrepeatsekvenser i vektoren indeholdende VL-sekvensen.
- 20      18.     En metode eller værtscelle ifølge de fornævnte krav 1-12, 15-17, hvori værtscellerne er hvirvelceller, helst celler fra pattedyr, allerhelst humane celler eller gnaverceller, så vidt muligt allerhelst lymfoïdceller stadig så vidt muligt allerhelst B-cell, i endnu højere grad allerhelst B-stamceller eller B-prækursorceller, specielt foretrukne Abelson-murin-leukæmivirus transformerede B-stamceller eller B-prækursorceller og 25 tidlige, EBV-transformerede humane proB- og preB-immunoglobulinnulceller.

Fig. 1:  
a.) autonomous transposable element (TE):



b.) bi-component transposon vector system



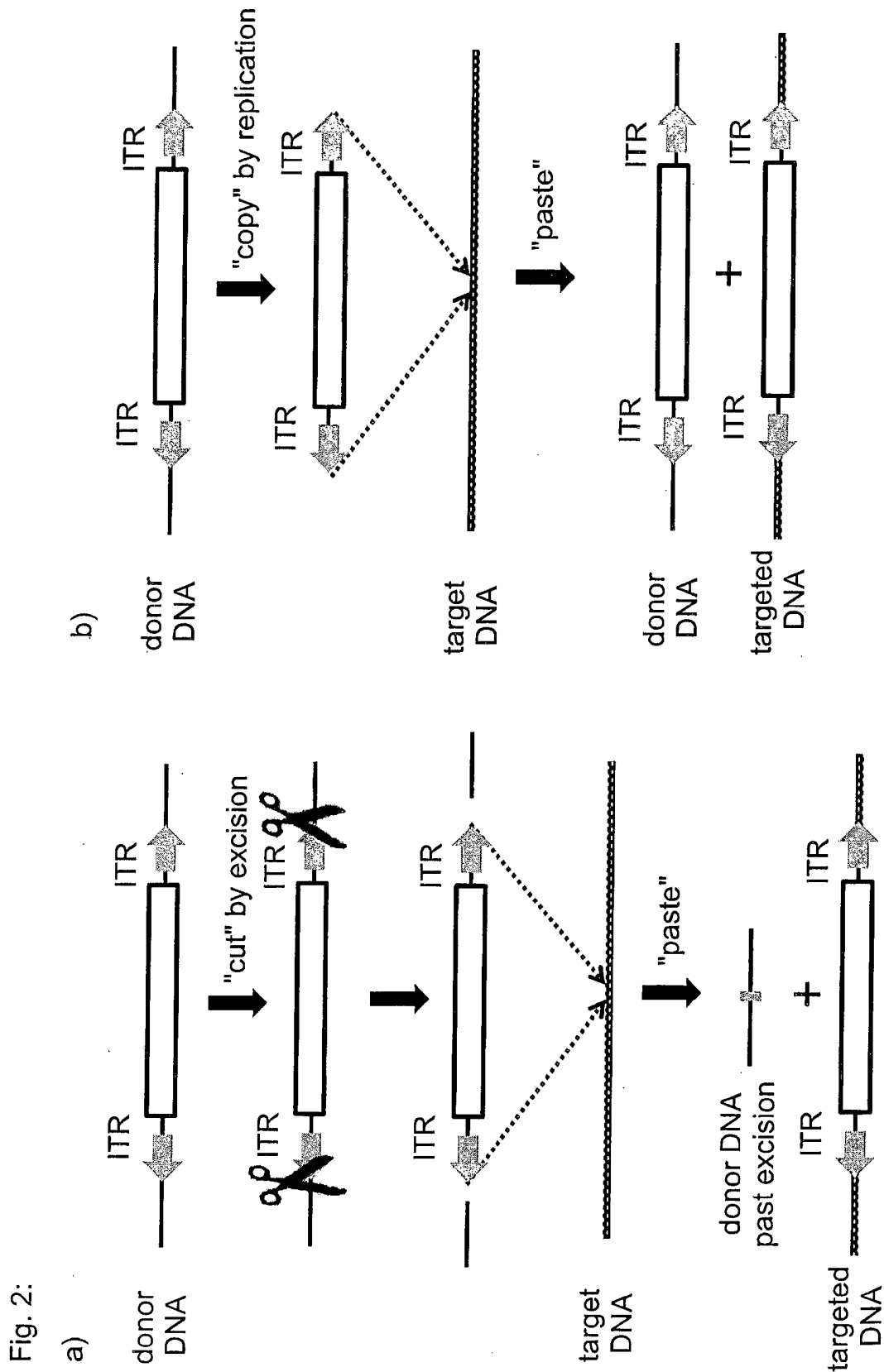


Fig. 3:

Transposon Family (Transposition)	Species or Organ	Host Cell Range
PiggyBac	<i>Trichoplusia ni</i>	Mouse, Human, Pig
Tcl-mariner (Sleeping Beauty)	<i>salmonid</i>	Zebrafish, Xenopus, Mouse, Human
(Frog Prince)	<i>R. pipiens</i>	Human, Hamster, Xenopus, Zebrafish
(Hmar1)	<i>H. irritans</i>	Human
(Passport)	<i>P. platessa</i>	Human, Monkey, Hamster, Turkey, Chicken, Pig
(Minos)	<i>D. hydei</i>	Human, Mouse
hAT (Tol1, Tol2)	<i>O. latipe</i>	Zebrafish, Xenopus, Mouse, Human, Chicken
AcDs	<i>Z. mays</i>	Zebrafish, Human
PIF, Harbinger, Harbinger3-DR	<i>D. rerio</i>	Zebrafish, Human
(Hsmar1)	<i>H. sapiens</i>	Human, Zebrafish

Fig. 4

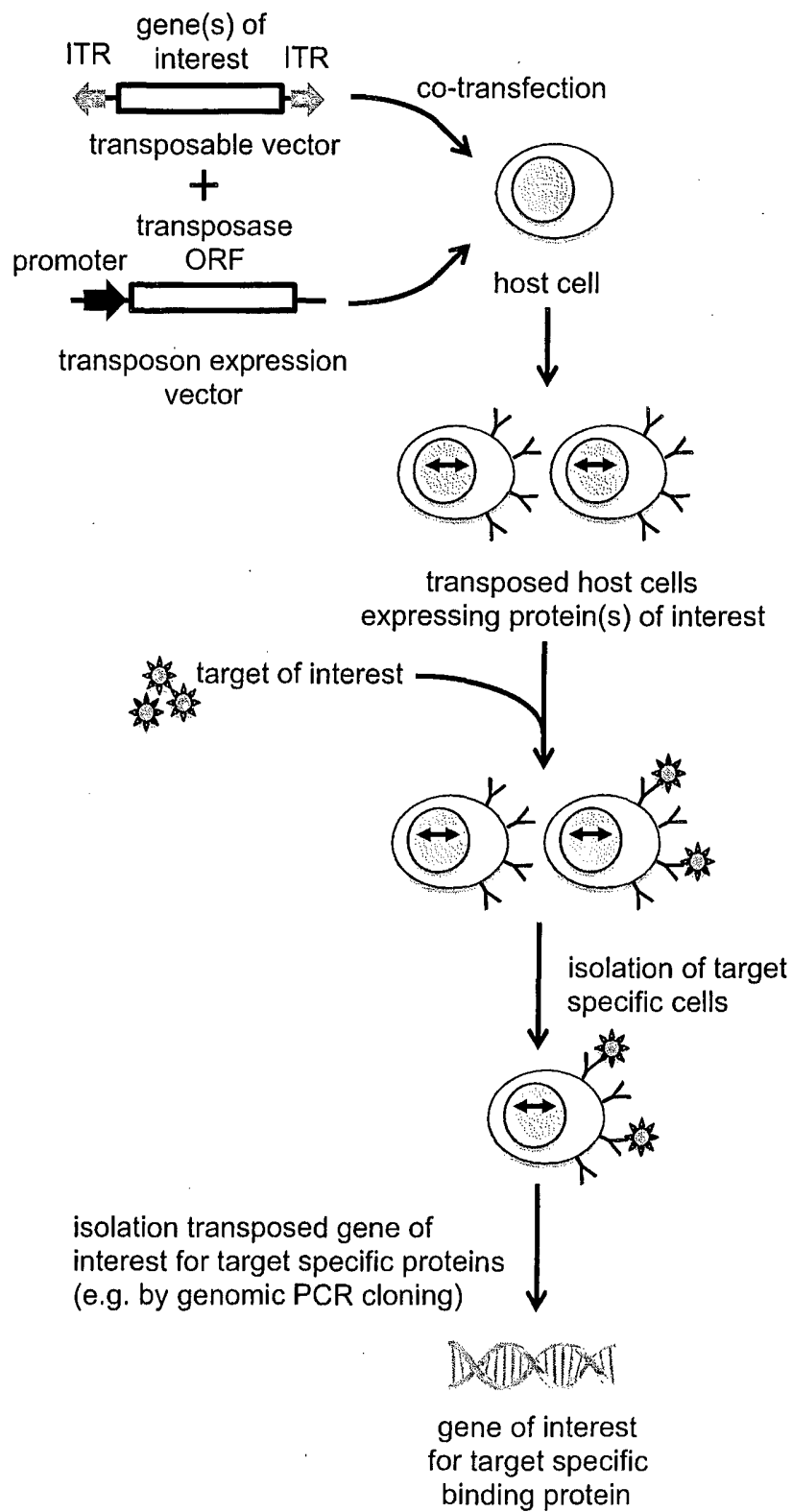


Fig. 5 a)

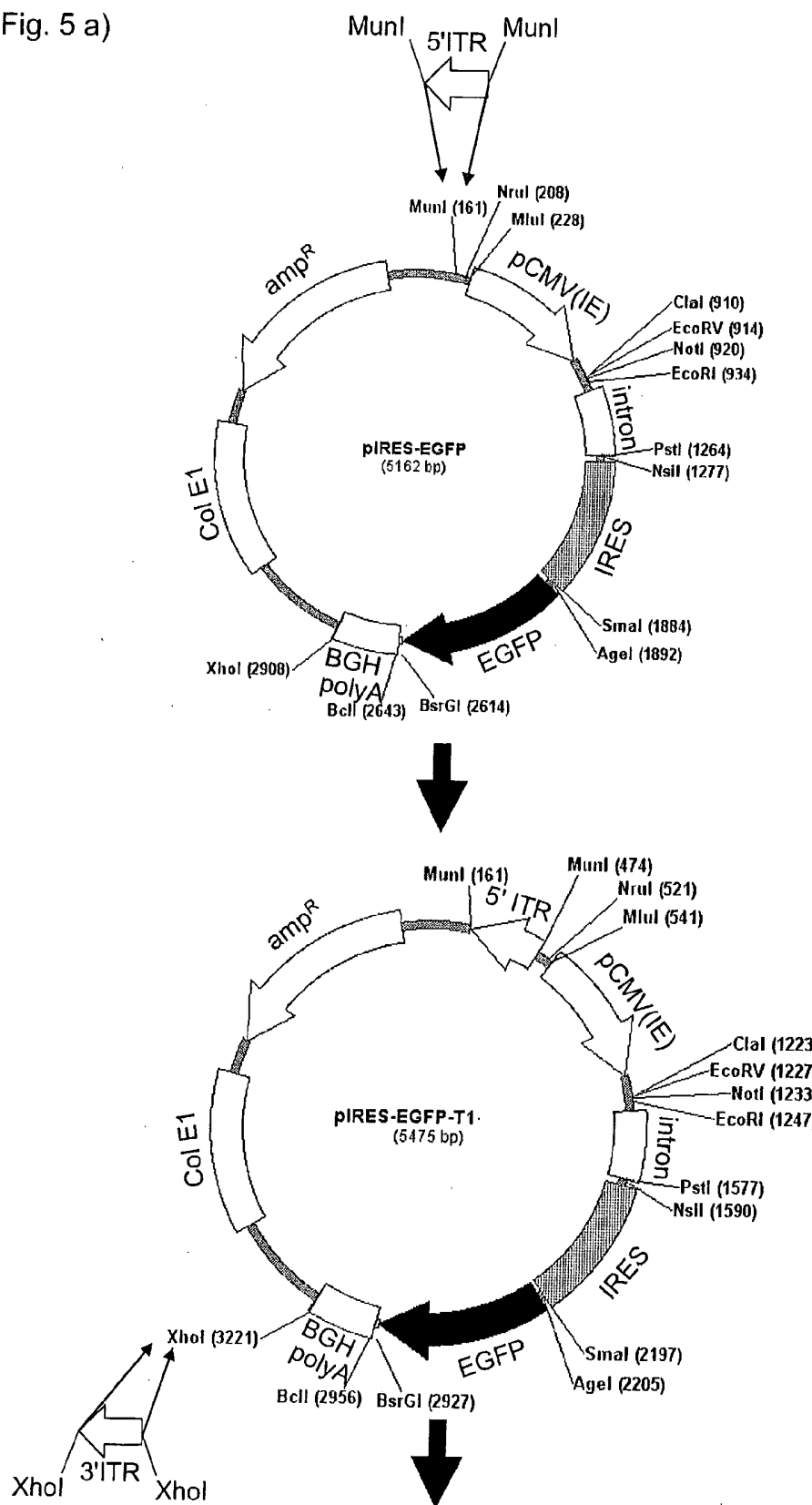


Fig. 5 b)

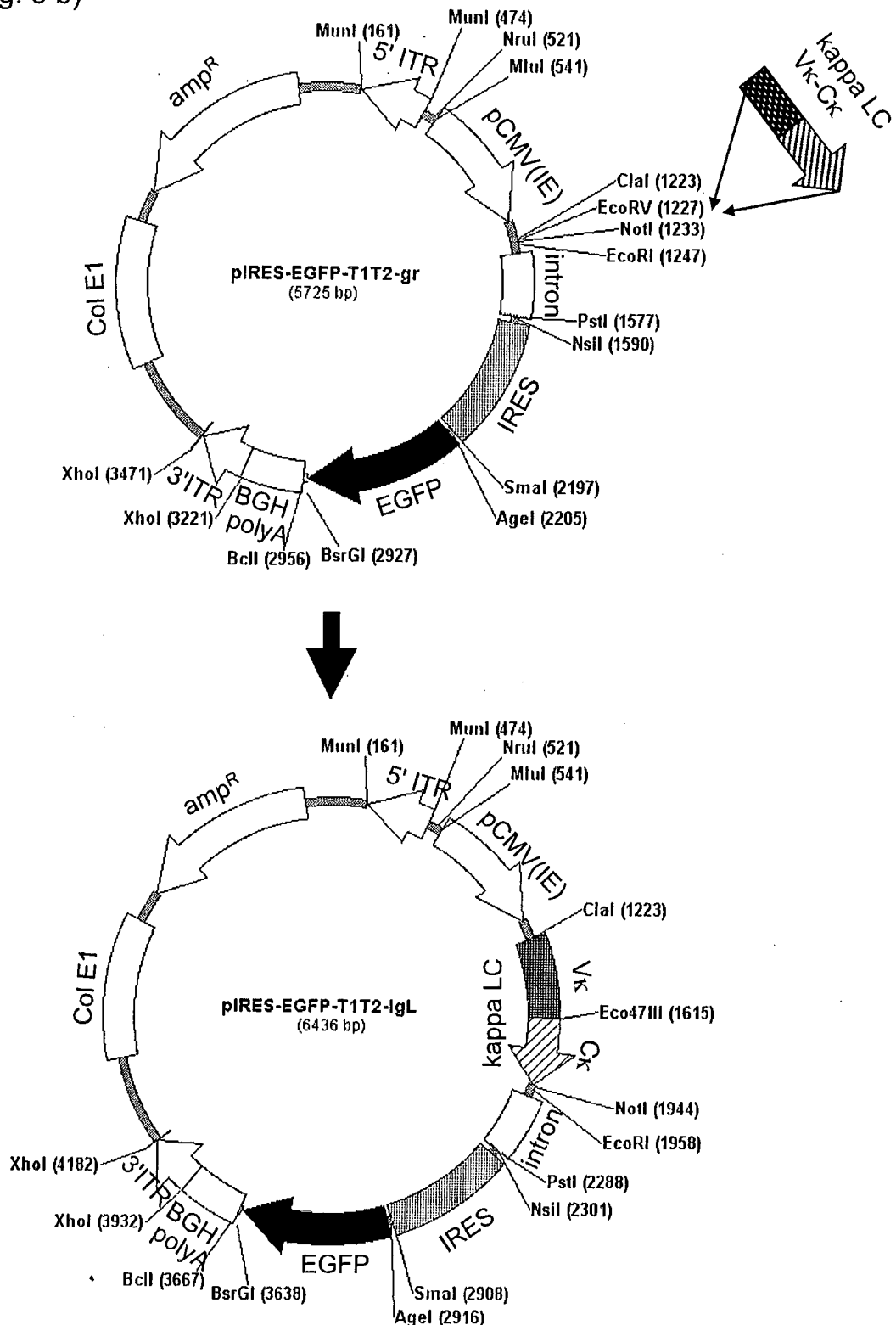


Fig. 6

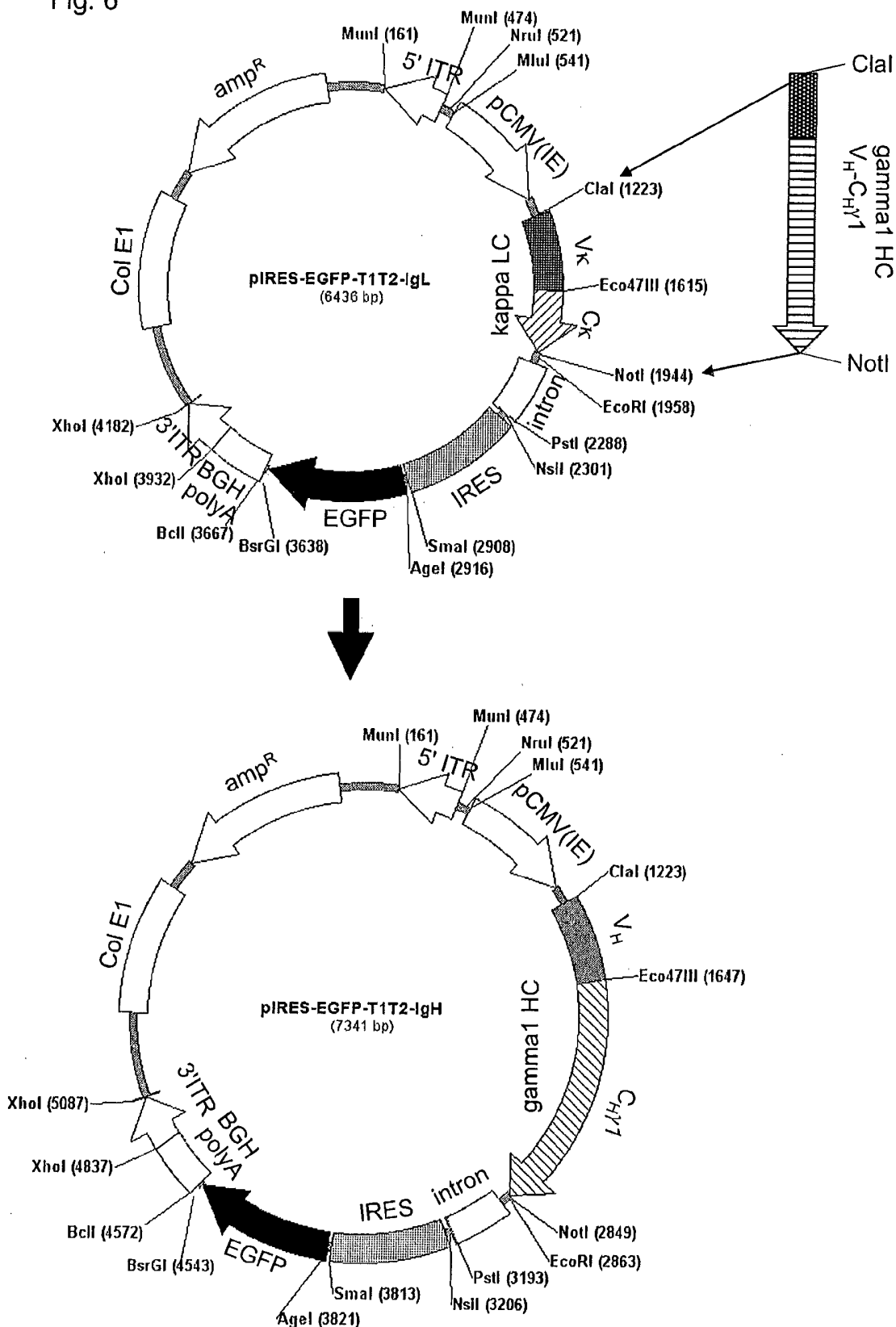


Fig. 7

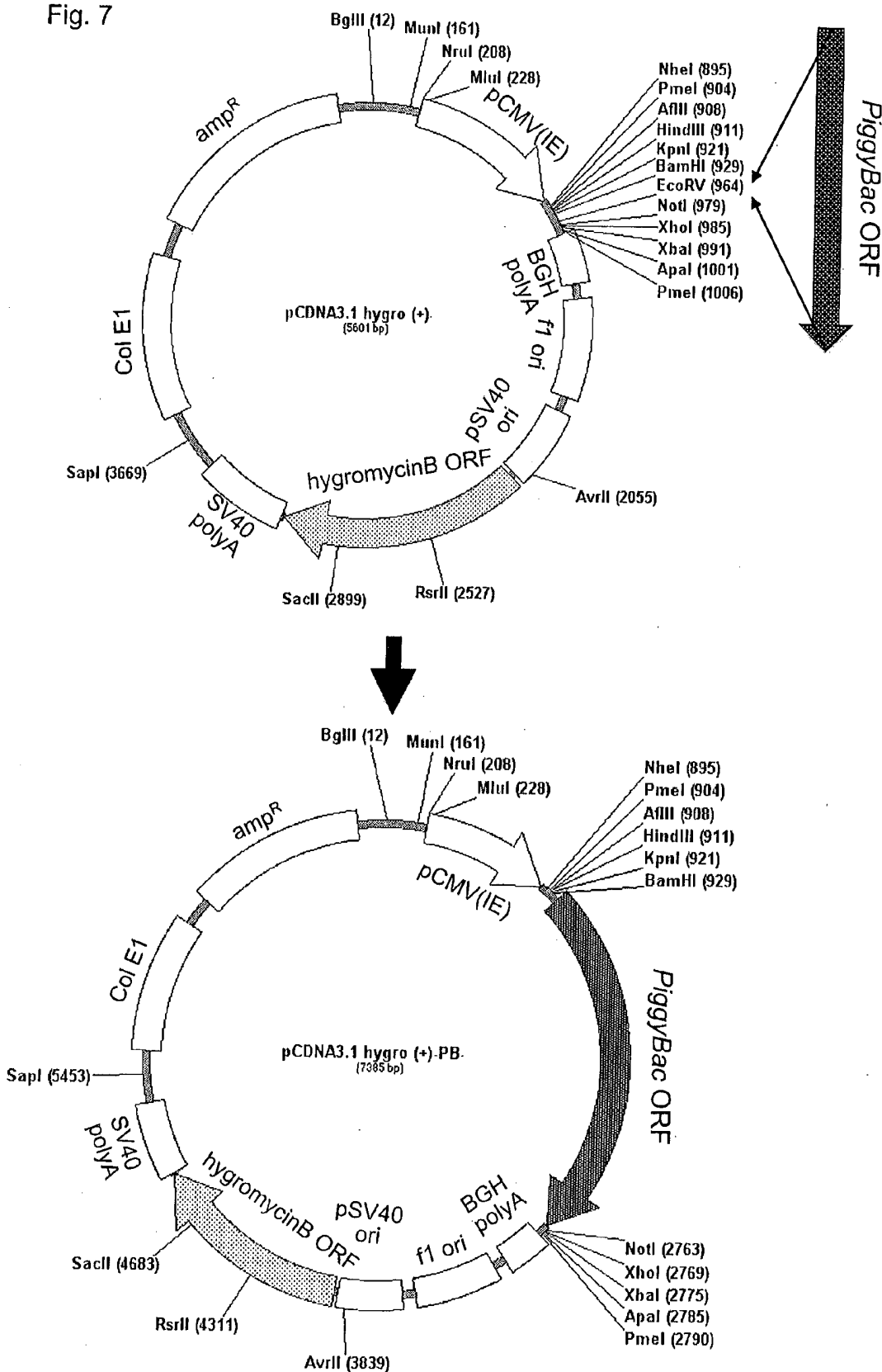


Fig. 8

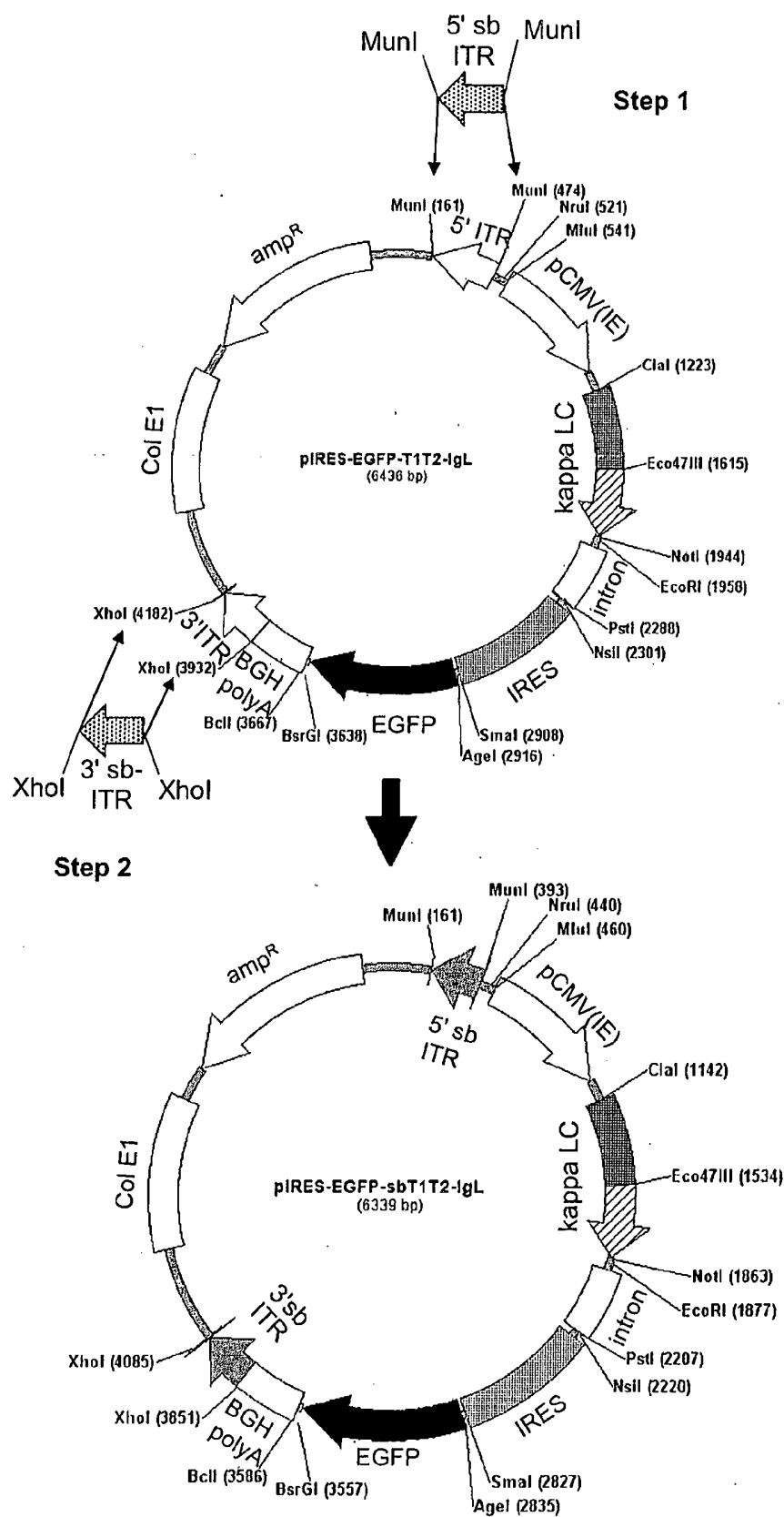


Fig. 9

