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**(71) Applicant:** MERUS N.V. [NL/NL]; Yalelaan 62, 3584 CM Utrecht (NL).

**(72) Inventors:** DE KRUIF, Cornelis Adriaan; c/o Yalelaan 62, 3584 CM Utrecht (NL). HENDRIKS, Linda Johanna Aleida; c/o Yalelaan 62, 3584 CM Utrecht (NL). LOGTENBERG, Ton; c/o Yalelaan 62, 3584 CM Utrecht (NL). VAN LOO, Pieter Fokko; c/o Yalelaan 62, 3584 CM Utrecht (NL).

**(74) Agent:** JANSEN, C.M.; P.O. Box 87930, 2508 DH Den Haag (NL).

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**(54) Title:** MULTIVALENT ANTIBODY

**(57) Abstract:** The invention relates to a multivalent antibody which comprises: a base antibody portion which comprises two binding domains; and at least one additional binding domain, wherein the base antibody portion is connected by a linker to the at least one additional binding domain, wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region, and wherein the linker comprises a hinge sequence or a sequence derived from a hinge sequence. The invention also relates to a multivalent antibody which comprises: a base antibody portion which comprises two binding domains; and at least one additional binding domain, wherein at least one additional binding domain comprises a CH1 region and is connected to the base antibody portion by said linker, linking a variable region of the base antibody portion and the CH1 region, and wherein the multivalent antibody binds to at least three different epitopes.

## **MULTIVALENT ANTIBODY**

### 5    **Field**

The invention relates to multivalent antibodies having three or more binding domains and to a method for making such multivalent antibodies. The invention further relates to constituent polypeptides of the multivalent antibodies and to linkers that can be used to connect 10 one or more binding domains of the multivalent antibody. The invention in addition relates to nucleic acids that encode such multivalent antibodies, linkers and to vectors comprising such nucleic acids, as well as to host cells that produce the multivalent antibodies. The invention also relates to multivalent antibodies which are capable of simultaneously binding three antigens or targets at once, including to target antigens present on cancer cells or tumor cell antigens and 15 targets that engage immune effector cells. Also the invention relates to a pharmaceutical composition which comprises the multivalent antibody and to the multivalent antibody for use in the treatment of the human or animal by therapy. Also, the invention relates to a method for the treatment of a human or animal using the antibody.

### 20    **Background**

Multivalent antibodies, such as bispecific antibodies, capable of binding two antigens or two epitopes are known in the art. Such multivalent binding proteins can be generated using various technologies, including cell fusion, chemical conjugation or recombinant DNA 25 techniques.

Antibodies typically are multimers comprised of four proteins, including two identical heavy chains and two identical light chains, wherein the heavy chain is comprised of a variable domain (VH), and three constant regions (CH1, CH2, CH3), and wherein the light chain is comprised of a variable light chain domain (VL) and a constant region (CL). Typically, the light 30 chain pairs with the heavy chain through the influence of many noncovalent interactions and also via disulphide bonds. The two heavy chains pair at the hinge region that connects CH1 to CH2 and/or through amino acid interactions in the interface between the two CH3 domains. The pairing of the VH with VL forms an antigen binding domain, and typically variability is found in three superficial-loop forming regions in the VH and VL domains, which are the 35 complementarity determining regions or CDRs.

Certain multivalent antibody formats are known in the art, such as antibodies having two different binding domains, such as in bispecific antibodies, that may bind two different antigens, or two different epitopes within the same antigen. Such a format may allow for the use of calibrated binding that will allow the multivalent antibody to be selectively targeted to cells or

targets that express two antigens or epitopes such as a tumor cell whilst not targeting healthy cells expressing one antigen, or to target such healthy cells expressing one antigen at lower expression levels. Similarly, having two different binding domains on a multivalent antibody, such as a bispecific antibody, may permit binding of different antigens, such that said

5 multivalent antibody could be used to target both an inhibitory and a stimulatory molecule on a single cell or on two interacting cells to result in enhanced potency of the multivalent antibody. A multivalent antibody could also be used to redirect cells, for example immunomodulatory cells, that could be redirected to a tumor.

The incorporation of more than two binding domains in a single antibody may allow for a 10 broader array of beneficial combinations of targets and efficacy. For example, a multivalent antibody having three or more binding domains may target the same antigen and epitope, permitting specificity for a given target and/or saturation of a target at a lower ratio of antibody to target. A multivalent antibody may contain two or more identical antigen binding domains to allow for high avidity binding to a target cell. This can be used to specifically target antigens 15 such as gangliosides that are over-expressed on tumor cells. These tumor associated antigens are present on normal cells but at much higher density on tumor cells. A multivalent antibody containing several lower affinity binding regions may allow for specific targeting to tumor cells while not reacting with healthy cells or doing so at a lower ratio and at the same time activating 20 or blocking additional receptors. Ultimately, three or more binding regions are useful in such applications.

While certain multivalent antibodies have been described in the art, there is a need in the art for new formats, and new linkers, that permit the efficient production of multivalent 25 antibodies, for which binding domains to an array of antigens can be readily made and converted into a multivalent antibody efficiently, stably, and that are capable of binding a wide array of antigens and epitopes. Engineering an antibody that contains more than two binding domains traditionally has been time-consuming, inefficient, and costly. Indeed, there are numerous impediments to efficiently producing high quality, low immunogenic multivalent 30 antibodies that can be generated to target a variety of antigens.

For example, existing multivalent formats containing three or four binding domains rely 35 on synthetic linkers such as Gly<sub>4</sub>Ser (G<sub>4</sub>S) repeats that contain sequence domains which firstly tend to restrict access to all binding domains in the molecule and secondly which may also be problematic for developability.

Existing multivalent antibody formats also rely on different heavy and light chains which are associated by disulphide bonds and amino acid interactions or can be joined by short linkers 35 in the case of single-chain fragment variables (scFvs). Yet the need to use differential variable chains (heavy and/or light) in each of the multiple variable domains used in a multispecific format of three or more binding specificities requires extensive engineering of such molecules to prevent mispairing of the heavy chains and light chains. Invariably this has an impact on complexity, stability, immunogenicity and production levels of these molecules.

Multivalent antibody formats may rely on the use of the same light chain for each binding domain, where one or more of the cognate variable regions paired with said light chain are forced to pair through chemical modifications, rather than the cognate chain being formed with and paired with a common light chain in response to antigenic exposure, and the processes 5 of co-evolution that occurs during B-cell development.

Multivalent antibody formats may also rely on the light chain from an existing monospecific antibody that binds one antigen, which is then used in a library to identify heavy chains capable of pairing said light chain, while also binding a distinct epitope or antigen.

Such pseudo-common light chains are not preferred for use in the present invention. 10 Preferred common light chains for the present invention are those that are capable of pairing with a diversity of cognate chains and are obtained from, derived from or based on common chains that pair with a rearranged cognate chain, which is encoded by DNA having undergone somatic recombination, and preferably somatic hypermutation in response to antigenic exposure.

15 A pseudo light chain approach limits the range of available cognate chains. Forcing the pairing of heavy and light chains, which were not formed together in a response to antigenic exposure results in loss of specificity and affinity limiting the utility. Further, it is typically rare for any given antibody to allow shuffling of VH and VL and conserve affinity and specificity.

20 Forcing light chain pairing with a heavy chain where the two have not co-evolved in the immune response while maintaining the capacity to bind an antigen is not trivial and may limit the flexibility of this approach.

25 Similarly, reusing a light chain of one antibody to identify heavy chains that are capable of binding said light chain and also bind a distinct antibody of interest, limits the range of available heavy chains, and becomes increasingly unlikely to identify suitable additional heavy chains the more antigens or epitopes sought to be bound. That is, one may use a light chain that pairs with a heavy chain to form a Fab that binds a given antigen to identify a subsequent heavy chain that pairs with said light chain, and is capable of binding a second antigen. However, use of that light chain a third time, to identify third heavy chain capable of pairing to 30 said light chain while binding a third antigen or epitope become increasingly rare, and rarer still the more heavy chains are sought to be identified capable of pairing with said light chain, while also binding distinct epitopes or antigens.

35 A preferred embodiment of the invention described herein employs a common chain, which pairs with a diversity of heavy chains in response to an antigen, and does not require reuse of an existing light chain from a monospecific antibody or forced pairing of a light chain to a cognate chain through chemical modification.

The successful construction of multivalent antibodies relies on the proper choice of protein linkers between the different domains as direct fusion of two domains can lead to compromised biological activity. The biophysical characteristics of the linker or linkers such as charge, rigidity or flexibility as well as the distance between the binding domains and the spatial

conformation between binding regions can impact epitope access and the ability of the multivalent protein to bind its targets. Thus there is a need for a multivalent format that in a modular fashion can employ a variety of linkers with different characteristics to permit construction of multivalent antibodies that allow simultaneous binding to different combinations 5 of epitopes that may be located on different molecules and/or on different cells.

Accordingly, there is a need in the art for the design of a set of linkers containing different characteristics of rigidity, flexibility, length that can be employed in a modular fashion depending on the combination of targets, while maintaining stability, low immunogenicity and ease of developability.

10 Accordingly, there is a need for new and useful formats for multivalent antibodies having three or more binding domains and linkers for the production of such antibodies which are widely applicable to the rapid and robust generation of a broad range of antibodies which comprise more than two binding domains.

15 **Summary**

The invention is based on new, modular, formats for a multivalent antibody comprising three or more binding domains which may be a multispecific antibody. In these formats, at least 20 one binding domain is connected to a base antibody portion, said base antibody portion comprising two binding domains. The additional binding domain may comprise a variable region, Fv domain, a Fab domain or a modified Fab domain or a functional fragment of any thereof. The base antibody portion may be, for example, a full length antibody or fragment thereof, but in each case comprises two binding domains.

The one or more additional binding domains are connected to the base antibody portion 25 via a linker(s), providing one or more binding moieties in addition to those of the base antibody portion.

A linker is used to connect the one or more additional binding domains to the base antibody portion. The linker comprises a peptide region, for example one or more hinge regions and/or one or more regions derived from a hinge region. The combination of the linker and a 30 constant region (e.g., CH1) to which it is connected may be critical in determining the properties of the multivalent antibody and allow correct functionality of the antibody and/or orientation of the one or more additional binding domains to the base antibody. Thus, if a linker sequence is based on a hinge of a given subtype it may be preferable that the constant region of the additional binding domain to which it is attached is of the same subtype.

35 The one or more additional binding domain(s) may comprise a variable region, Fv domain, a Fab domain or modified Fab domain.

Fab domains in particular constitute beneficial additional binding domains since they comprise protein domains having predictable behavior which is useful for the manufacture of multivalent molecules which are stable and can be readily manufactured.

Facilitating efficient production and developability of the multivalent antibody of the invention, said multivalent antibody may comprise a common variable region, which can be an immunoglobulin heavy chain variable region (VH) or a light chain variable region (VL), but is typically a common light chain (cLC) variable region.

5 The common variable region is typically paired with a cognate variable region that is encoded by a nucleic acid that has undergone somatic recombination and preferably affinity maturation or is based on or derived from a rearranged variable region that is encoded by a nucleic acid that has undergone the process of somatic recombination and preferably affinity maturation, and/or is based on or derived from known antibody generation techniques, such as  
10 phage display, immunization of animals, including transgenic animals with humanized immune systems and other techniques well known in the art.

The use of a common variable region that is essentially identical in each binding domain of the multivalent antibody of the invention facilitates the development and manufacture of such antibodies.

15 The choice of the common variable region for use in a multivalent antibody of the invention should thus be one which can be used widely with many different cognate heavy or light regions.

An identical, or substantially identical, common variable region, for example a cLC variable region, allows the use of complete, or substantially complete, Fab domains for all three  
20 or more binding regions without the need for extensive engineering as used in Crossmab technology or for linkers to prevent heavy and light chain mispairing, such as those used in scFv domains. Also, since essentially germline encoded and non-immunogenic common variable regions are known (see WO2009/157771), using it in each of the three or more Fab domains may provide for reduced immunogenicity.

25 An additional benefit of the format described herein for producing multivalent antibodies permits the use of transgenic animals, preferably transgenic rodents, that have within their genomes a common variable region, capable of pairing with a diversity of cognate variable regions (e.g., common light chain variable region pairing with a diversity of heavy chain variable regions) (see WO2009/157771), that permits the DNA encoding the cognate variable regions  
30 formed from exposure to different antigens to be introduced into a host cell with the DNA encoding the common variable region, which can each be expressed for the generation of multivalent antibodies.

For example, a transgenic mouse comprising in its germline, DNA encoding a common variable region and DNA encoding an unarranged immunoglobulin locus that can rearrange to  
35 form a cognate variable region and is capable of undergoing somatic recombination, can be exposed to one or more antigens, such that the rearranged variable regions produced based on exposure to the three or more antigens can then be used to generate a multivalent antibody of the present invention. Nucleic acid sequences encoding the common variable region, and the

three or more rearranged variable regions can be transformed into a host cell, to express a multivalent antibody of the present invention.

Thus, the formats described herein for producing multivalent antibodies make use of three or more binding domains that all may comprise a common variable region, preferably a common light chain variable region.

The formats of the present invention comprise one or more binding domains in addition to those of the base antibody portion. Such an additional binding domain may be a Fv domain, a Fab domain or a modified Fab domain preferably comprising a CH1 domain and a variable domain, which is connected to the base antibody portion via a linker. In a modified Fab domain, the CH1 domain is not paired with a CL. A suitable CH1 domain may be one which is engineered to remove one or more hydrophobic regions or may be one derived from a camelid animal or a shark. Alternatively, an additional binding domain may comprise a CL domain and a variable domain which is connected to the base antibody portion via a linker. The CL domain can either be a Cκappa or a Cλambda domain.

Typically, the additional binding domain(s) is/are connected to one or both binding domains of the base antibody portion at the N-terminal region of either a common variable portion or a rearranged variable region of the binding domain of the base antibody portion or both.

Alternatively, an additional binding domain may be connected to the base antibody portion via a linker connecting both a common chain and a rearranged variable domain of the binding domain of the base antibody portion to a CH1 and CL region of the additional binding domain. Where an additional binding domain lacks a constant region, novel linkers disclosed herein may connect both a common chain and a rearranged variable domain of the binding domain of the base antibody portion to the common chain and/or rearranged variable region of the additional binding domain.

Alternatively, an additional binding domain may be a  $V_H$  and  $V_L$  region, i.e. a Fv domain, which is connected by a single linker peptide to the base antibody portion. Typically, this type of additional binding domain(s) is/are attached to one or both binding domains of the base antibody portion at the N-terminal region of either the common variable portion or the rearranged variable region of the binding domain of the base antibody portion.

This format, when used with linkers of the invention disclosed herein, including different lengths, structures and degrees of rigidity, is surprisingly flexible. Thus, the invention provides a repertoire of linkers with different properties for use in the disclosed multivalent antibody formats that render it developmentally facile to combine three or more binding domains into a multivalent antibody.

By way of the linkers disclosed herein, the invention thus provides a modular approach in which selection of the appropriate linker together with selection of a corresponding set of binding domains, such as Fab domains, allows those binding domains to function together in a multivalent antibody for a variety of efficacy.

A multivalent antibody of the invention may be used in therapy, in particular, as a so-called "engager" antibody whereby the antibody is capable of forming a link between an immune effector cell and a tumor cell.

According to the invention, there is thus provided a multivalent antibody which  
5 comprises:

- a base antibody portion which comprises two binding domains; and
- at least one additional binding domain,

10 wherein the base antibody portion is connected by a linker to the at least one additional binding domain,

wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region, and

15 wherein the linker comprises a hinge sequence or a sequence derived from a hinge sequence.

The invention also provides a multivalent antibody which comprises:

20 - a base antibody portion which comprises two binding domains; and

- at least one additional binding domain,

wherein at least one additional binding domain comprises a CH1 region and is connected to the base antibody portion by said linker, linking a variable region of the base  
25 antibody portion and the CH1 region, and

wherein the multivalent antibody binds to at least three different epitopes.

Preferably, each binding domain of the base antibody portion and each of the at least one additional binding domains may all have a common variable region,

30 **Variety of the One or More Additional Binding Domain**

A preferred embodiment is a multivalent antibody, wherein one or more binding domains is a Fv domain comprising a heavy chain variable region (VH) and a light chain  
35 variable region (VL).

A further preferred embodiment is a multivalent antibody, wherein one or more binding domains is a Fab domain comprising a heavy chain variable region (VH) and a light chain variable region (VL), said heavy chain variable region of said Fab domain comprising a CH1 region (VH-CH1) and said light chain variable region of said Fab comprising a CL region (VL-

CL). Said Fab domain may contain a VL-CL that is either a Vkappa-Ckappa, Vlambda-Clambda, Vlambda-Ckappa or Vkappa-Clambda.

Another embodiment is a multivalent antibody, wherein the one or more additional binding domains is a modified Fab domain consisting of a VH-CH1 and VL. Alternatively, an 5 embodiment is a multivalent antibody, wherein the one or more additional binding domain is a modified Fab domain consisting of a VL-CL and a VH. In such modified Fab domains, a constant region, CH1 or CL, is present which is not paired with its cognate region and/or a variable region VH or VL, is present, which is not paired with its cognate region.

### **Common Chain**

10

A preferred embodiment is a multivalent antibody wherein the one or more additional binding domain comprises a Fab domain comprising a common rearranged variable region paired to a rearranged variable region which has undergone somatic rearrangement following exposure to an antigen or is encoded by nucleic acids obtained from, derived from, or based on 15 a sequence, which is the result of somatic rearrangement. Alternatively, the rearranged variable region could be one obtained from, derived from, or based on a synthetic repertoire where diversity is introduced into a repertoire using molecular biology techniques known in the art, including the use of synthetic phage display libraries. Preferably, said Fab domain comprises a common light chain variable region paired to a counterpart rearranged heavy chain variable 20 region. Preferably, said common light chain variable region is connected to a CL region and said rearranged heavy chain variable region is connected to a CH1 region. Preferably, said common light chain is paired to said heavy chain variable region via joining of the CL and CH1 regions. Alternatively, wherein the common chain is a heavy chain, the rearranged variable region is a light chain, and said chains may comprise a CH1 and CL domain respectively and 25 may be paired via joining of the CL and CH1 regions.

A preferred embodiment is a multivalent antibody wherein the three or more binding domains each comprise the same common chain, but wherein the three or more binding domains comprise different rearranged variable cognate chains, more preferably, wherein the said same common chain is a common light chain.

30

A preferred embodiment is a multivalent antibody, wherein the three or more binding domains comprise rearranged variable regions encoded by nucleic acids obtained from, derived from, or based on nucleic acid sequences of a transgenic animal comprising a common light chain and unarranged heavy chain variable region, which has been exposed to an antigen and has produced antibodies comprising a rearranged heavy chain variable region paired to a 35 common light chain. Alternatively, in an embodiment of a multivalent antibody, the three or more binding domains comprise rearranged variable regions encoded by nucleic acids obtained from, derived from, or based on nucleic acid sequences of a transgenic animal comprising a common heavy chain and unarranged light chain variable region, which has been exposed to

an antigen and has produced antibodies comprising a rearranged light chain variable region paired to a common heavy chain.

**Linker Composition**

5 A preferred embodiment is said multivalent antibody, wherein said linker is a naturally occurring sequence, or based on a naturally occurring sequence. More specifically, said linker is a hinge sequence or comprises a sequence based on a hinge sequence. More specifically said linker may comprise a hinge region based on an IgG1 hinge region, an IgG2 hinge region, an IgG3 hinge region or an IgG4 hinge region.

10 Alternatively, said linker comprises a peptide region comprising one or more of the following:

ESKYGPP (SEQ ID NO: 1)  
EPKSCDKTHT (SEQ ID NO: 2)  
15 GGGGSGGGGS (SEQ ID NO: 3)  
ERKSSVESPPSP (SEQ ID NO: 4)  
ERKCSVESPPSP (SEQ ID NO: 5)  
ELKTPLGDTTHT (SEQ ID NO: 6)  
ESKYGPPSPSSP (SEQ ID NO: 7)  
20 ERKSSVEAPPVAG (SEQ ID NO: 8)  
ERKCSVEAPPVAG (SEQ ID NO: 9)  
ESKYGPPAPEFLGG (SEQ ID NO: 10)  
EPKSCDKTHTSPPSP (SEQ ID NO: 11)  
EPKSCDGGSGGSGGGGS (SEQ ID NO: 12)  
25 GGGGSGGGGSAPPVAG (SEQ ID NO: 13)  
EPKSCDKTHTAPELLGG (SEQ ID NO: 14)  
ERKSSVESPPSPAPPVAG (SEQ ID NO: 15)  
ERKCSVESPPSPAPPVAG (SEQ ID NO: 16)  
ELKTPLGDTTHTAPEFLGG (SEQ ID NO: 17)  
30 ESKYGPPSPSSPAPEFLGG (SEQ ID NO 18)  
EPKSCDKTHTSPPSPAPELLGG (SEQ ID NO: 19)  
ERKSSVEAAAKEAAAKAPPVAG (SEQ ID NO: 20)  
ERKCSVEAAAKEAAAKAPPVAG (SEQ ID NO: 21)  
ESKYGPPEAAAKEAAAKAPEFLGG (SEQ ID NO: 22)  
35 EPKSCDKTHTEEAAAKEAAAKAPELLGG (SEQ ID NO: 23)  
ELKTPLGDTTHTEEAAAKEAAAKAPEFLGG (SEQ ID NO: 24)

or a sequence having at least about 85% sequence identity to any one thereto.

Preferably, a multivalent antibody of the invention comprises a linker that connects the base antibody portion to the one or more binding domains by comprising an amino acid sequence of any one of SEQ ID NOs: 1 to 24 or a polypeptide comprising an amino acid sequence having at least about 85% sequence identity to any of SEQ ID NOs: 1 to 24.

5 Table 1 illustrates how such linkers may be connected to a CH1 region.

A preferred multivalent antibody of the invention, comprises a linker that is rigid. More preferably, said multivalent antibody comprises a linker that comprises a helix-forming sequence.

10 A preferred multivalent antibody of the invention comprises a linker comprising a peptide sequence comprising a (EAAK)<sub>2</sub> motif.

A preferred multivalent antibody of the invention, comprises a linker that is flexible.

A preferred multivalent antibody of the invention, comprises a linker comprising three or more amino acid residues that correspond to a hinge region of a subtype of a constant region to which it is connected of said multivalent antibody.

15 A preferred multivalent antibody of the invention comprises a linker comprising a sequence of SEQ ID NOs: 1 to 24 that corresponds to a hinge region of a subtype of a constant region to which it is connected of said multivalent antibody.

#### Linker Location/Orientation

20 A preferred embodiment is a multivalent antibody, wherein the base antibody portion is connected to the one or more additional binding domains by a linker, wherein said linker joins an N-terminal end of a variable region of said base antibody portion to the C-terminal end of the one or more additional binding domains. Preferably, the base antibody portion comprises a Fab domain and the one or more additional binding domain comprises a Fab domain comprising a 25 CH1 domain and CL domain and the linker connects an N-terminal end of a variable region of the Fab of the base antibody portion to either or both of a C-terminal end of the CH1 domain and CL domain of the Fab domain of the one or more additional binding domain.

A preferred embodiment of the invention is a multivalent antibody comprising a common chain at each binding domain of the base antibody portion and each of the one or more 30 additional binding domains and a linker which connects a N-terminal end of a rearranged variable region of the base antibody portion to a C-terminal end of a rearranged variable region of the one or more additional binding domain. More preferably, the one or more additional binding domain comprises a Fab domain comprising a CH1 domain and CL domain and the linker connects an N-terminal end of a variable region of the Fab domain of the base antibody 35 portion to the CH1 domain or CL domain of the Fab domain of the one or more additional binding domain.

**Pairing of the regions comprising the additional binding domain**

Antibody assembly occurs through association of the light and heavy chains, namely the association (pairing) of VH with VL and CH1 with CL, which is based on interacting residues 5 in the interface between VH and VL, and between CH1 and CL. Typically, pairing is further stabilized, whereby a light chain is covalently connected to the heavy chain by a disulfide bond between a cysteine residue of the light chain in the CL, and a cysteine residue of the heavy chain at the CH1 or hinge, depending on the subtype.

Thus, in a multivalent antibody of the invention, an additional one or more binding 10 domain(s) is connected to the base antibody portion via a linker(s), wherein the one or more binding domain(s) comprises a Fv domain, a Fab domain or a modified Fab domain, and the counterpart immunoglobulin chains that comprise the binding domain (typically a heavy and light chain region) are paired together in a stable association.

For multivalent antibodies of the present invention that contain an additional binding 15 domain, wherein the binding domain is a Fv or Fab domain a cysteine residue may be present or engineered into the heavy and light chain domains, such that a disulfide bond forms to stabilize the pairing between the heavy and light chain of the additional binding domain. Where the multivalent antibody of the invention includes an additional binding domain comprising an IgG1 subclass, an upper hinge of IgG1 (EPKSC) of the heavy chain may be used that is 20 connected to and upstream (n-terminal side) of an artificial linker, such as (G4S)n, to provide a cysteine to covalently pair with the light chain of the additional binding domain. For other subclasses used in the additional binding domain, the skilled artisan will recognize the ability to engineer a cysteine residue in the linker employed to stabilize the pairing of light and heavy chain domains of the additional binding domain and to form a disulfide bridge between said light 25 chain and heavy chain used.

The wild-type IgG1 hinge region has the sequence: EPKSCDKTHTCPPCPAPELLGG (SEQ ID NO: 42). The underlined C residue is Cys220 which in the IgG1 heavy chain pairs with Cys214 of the light chain. Where a multivalent antibody of the invention comprises a linker based on such a hinge, preferably any Cys residues other than Cys220 are substituted with an 30 amino acid residue which cannot form a disulphide bond, for example Ser.

The wild type IgG2 hinge region has the sequence: ERKCCCCVECPPCPAPPVAG (SEQ ID NO: 46). The underlined C residue is Cys219 which in the IgG2-B heavy chain pairs with Cys214 of the light chain. In IgG2-A, Cys127 in the heavy chain pairs with Cys214. Where a multivalent antibody of the invention comprises a linker based on such a hinge, preferably any 35 Cys residues other than Cys215 are substituted with an amino acid residue which cannot form a disulphide bond, for example Ser.

The wild type IgG3 hinge region has the sequence:  
ELKTPPLGDTTHTCPRCPEPKSCDTPPPCPRCPEPKSCDTPPPCPRCPEPKSCDTPPPCPRCP

APEFLGG (SEQ ID NO: 50). In IgG3, Cys131 in the heavy chain pairs with Cys214 of the light chain.

The IgG4 hinge region has the sequence: ESKYGPPCPCPSCPAPEFLGG (SEQ ID NO: 54). In IgG4, Cys131 in the heavy chain pairs with Cys214 of the light chain. Where a 5 multivalent antibody of the invention comprises a linker based on such a hinge, preferably one or both of the Cys residues in the CPSPC region of the hinge are substituted with an amino acid residue which cannot form a disulphide bond, for example Ser.

As set out herein, to generate multivalent constructs, including trivalent constructs, based on an IgG structure capable of binding simultaneously three different epitopes, linkers are 10 employed based on IgG hinges from different subclasses to connect a binding domain of a base antibody portion to an additional binding domain that comprise a heavy chain constant region. To ensure stabilization of the covalent bond between a cysteine in the light chain and a cysteine in the heavy chain of the additional binding domain, the invention matches a linker comprising a hinge region or based on a hinge region of a particular subtype with the CH1 of the 15 additional binding domain that is from the same subtype.

Where the multivalent antibody comprises an additional binding domain comprised of pair regions (e.g., VH-CH1 paired with VL-CL, or VH paired with VL), stabilizing the interface between the regions may be accomplished in the present invention in a variety of ways. Where the one or more additional binding domain is a Fab domain comprised of a variable heavy chain 20 region and variable light chain region, a CH1 can be connected to the variable heavy chain region. The CH1 can be paired to a CL with a covalent bond, typically a disulphide bridge, that is connected to the variable light chain region. In addition heavy and light chains of a Fab domain are paired via non-covalent interactions. Alternatively, the linker that connects the one or more additional binding domains to the base antibody portion can also be used to pair the 25 variable heavy chain region to the variable light chain region of the additional binding domain by forming a peptide bond with either chain and a covalent linkage with the counterpart chain. This can be accomplished by designing a cysteine at or near the N-terminus of the linker and a cysteine at or near the C-termini of the variable heavy chain and/or variable light chain regions of the one or more additional binding domains, thereby forming a covalent bond between the 30 linker and the variable heavy chain and/or variable light chain regions of the one or more additional binding domains. Other means of pairing the domains that comprise the one or more additional binding domain, such as a Fab domain, are known to persons of ordinary skill in the art and described further in detail below.

A preferred embodiment is a multivalent antibody, comprising a base antibody portion 35 and one or more additional binding domains. Where the one or more additional binding domain is a Fv domain comprising a heavy chain variable (VH) region and light chain variable (VL) region, a linker of the invention connects the base antibody portion to said Fv, while pairing the heavy chain variable region and light chain variable region of the Fv.

Alternatively, the binding domain is a Fab domain comprising a variable heavy region comprising a CH1 region and a variable light region comprising a CL region. A linker of the invention connects the base antibody portion to the Fab domain at the CH1 region or CL region or both while pairing the CH1 and CL region of the Fab domain.

##### 5 **Pairing of the Base Antibody Portion**

Different techniques are known in the art to pair and cause heterodimerization of heavy chain constant regions (e.g., CH2 and CH3) of a base antibody portion. The use of, for example, DEKK mutations to cause heterodimerization of antibody heavy chains

10 (WO2013/157954 and De Nardis et al., *J. Biol. Chem.* (2017) 292(35) 14706-14717 incorporated herein by reference), further allows for efficient heterodimer formation, stable Fc region and ease of manufacture. This approach leaves the Fc region of the molecule functional and capable of engaging with immune receptors such as Fc receptors, complement and FcRn. Accordingly, certain multivalent antibody embodiments of the invention employ the DEKK  
15 modifications, or other Fc modifications known to skilled artisans, to preferentially heterodimerize the heavy chains of the base antibody portion.

Further, for certain embodiments of multivalent antibodies, it may be desirable not to engage the immune system's effector function (e.g. to limit antibody-dependent cellular cytotoxicity, antibody-mediated phagocytosis and/or cellular-dependent cytotoxicity), such as  
20 using a multivalent antibody to engage, stimulate and/or co-stimulate T-cells, in which case additional modifications may be employed to the Fc region to eliminate or mitigate effector function. Accordingly, certain multivalent antibody embodiments of the invention contain modifications to the heavy chain constant regions of the base antibody portion that eliminate or mitigate effector function(s).

25 An additional preferred embodiment of the invention is a multivalent antibody comprising a base antibody portion comprised of two heavy chains that lack CH2 or CH3 region, wherein said heavy chains are bound together at the hinge region.

The invention also provides a method for the preparation of a multivalent antibody, which method comprises providing a cell which comprises one or more nucleic acid sequences  
30 encoding polypeptides which are capable of assembly into a multivalent antibody of the invention. The cell may be cultivated under conditions to provide for expression of the base antibody portion, the at least one additional binding domain and the at least one linker and for their assembly into a multivalent antibody of the invention.

35 The invention also provides nucleic acids that encode for the constituent proteins of a multivalent antibody of the invention and the multivalent antibody they produce.

The invention also provides a vector comprising a nucleic acid sequence of the invention.

The invention also provides host cells that express said nucleic acids and produce said multivalent antibodies.

The invention also provides methods of generating said multivalent antibody, including through the use of transgenic animals comprising a common chain in its germline that produce common chain antibodies having diversity at a cognate chain, wherein said multivalent antibody comprises one or more binding domains having a rearranged variable region encoded by a

5 nucleic acid obtained from, derived from or based on one or more cognate chains of the common chain antibodies expressed by the transgenic animal exposed to an antigen.

The invention also provides a non-human transgenic animal comprising a common human light chain variable region capable of pairing with a diversity of human heavy chain variable regions, wherein the nucleic acids encoding the common light chain variable region and

10 human heavy chain variable regions are present at the non-human transgenic animal's endogenous variable region loci (light and heavy respectively or vice-versa) and/or are stably integrated elsewhere in the germline of the said transgenic animal (e.g., the Rosa locus), wherein contacting said transgenic animal with an antigen generates an array of rearranged human heavy chain variable regions that pair with said common light chain variable region,

15 wherein a nucleic acid encoding said rearranged human heavy chain variable regions is transformed into a host cell capable of producing a multivalent antibody of the invention, and the multivalent antibody comprises one or more binding domains comprising said rearranged human heavy chain variable region encoded by a nucleic acid obtained from, derived from or based on one or more rearranged human heavy chain variable regions produced by the

20 transgenic animal exposed to an antigen.

The invention also provides a pharmaceutical composition which comprises an antibody of the invention and a pharmaceutically acceptable carrier and/or diluent.

The invention also provides an antibody of the invention for use in the treatment of the human or animal body by therapy.

25 The invention also provides a method for the treatment of a human or animal suffering from a medical indication, which method comprises administering to the human or animal a therapeutically effective amount of an antibody of the invention.

#### **Brief description of the drawings**

30 For ease of reference, for Figures fifteen through twenty-eight, when describing trispecific molecules, the following format is used MFAxMFB:MFC or AntigenAxAntigenB:AntigenC, such that MFA or AntigenA followed by x constitutes the "short arm", while the x denotes the dimerization, followed by MFB or AntigenB describes the interior position of the long arm, followed by a ":" designating a linker followed by MFC or AntigenC

35 describes MFC or AntigenC at the distal domain of the long arm. Where the term "mock" is used in the context of a multivalent molecule, it refers to a binding domain of such molecule, which is capable of binding an antigen not present in the given assay in which it is tested. Typically, mock binding domains used herein bind tetanus toxin (TT), fibrinogen (Fibri) or thyroglobulin (Thyro).

Figure 1(a-u) sets out formats of multivalent antibodies of the invention, including different binding domain structures, linkers and base antibody portions.

Figure 2a sets out a schematic diagram of the VH1-CH1-linker-VH2 insert used for cloning the constructs in the vector MV1626. It is understood, though not shown, that the vector 5 also may encode CH3-CH2-CH1 region, which is connected to the VH2. Figure 2b shows a trispecific antibody, where VH1 binds a tetanus toxoid antigen, VH2 binds a fibrinogen antigen and VH3 binds a thyroglobulin antigen.

Figure 3 sets out a schematic diagram of the MV1626 vector.

Figure 4 sets out a schematic diagram of the MG1025C377 expression vector.

10 Figure 5 sets out the alignment of sequences of inserts used for cloning into MV1626. Note the alignment covers only CH1-linker for clarity purposes.

Figure 6 sets out a schematic diagram of the MV1057 vector.

Figure 7 sets out a schematic diagram of the MV1260 vector.

15 Figure 8 sets out the SDS-PAGE gels of IgGs in non-reducing (top) and reducing (bottom) conditions.

Figure 9 sets out screening data of 24 multivalent constructs.

Figure 10 sets out screening data of 18 multivalent constructs.

20 Figure 11A: Common light chain amino acid sequence. Figure 11B: Common light chain variable domain DNA sequence and translation (IGKV1-39/jk1). Figure 11C: Common light chain constant region DNA sequence and translation. Figure 11D: IGKV1-39/jk5 common light chain variable domain translation. Figure 11E: V-region IGKV1-39A; Figure 11F: CDR1, CDR2 and CDR3 of the common light chain.

Figure 12: Stability analysis of 18 multivalent IgG constructs and 4 control antibodies analyzed under 4 different conditions.

25 Figure 13: Bioinformatic modeling of 8 linkers.

Figure 14: Two engager trispecific formats, with a long arm interior immune cell binding domain and a short arm tumor cell antigen binding domain (14a) or long arm distal tumor cell antigen binding domain and a short arm tumor cell antigen binding domain (14b).

30 Figure 15: T cell activation in BxPC3 cells (median EGFR expression) by flow cytometry with the expression of CD25 and CD69 as a read out for both formats: EGFR on the short arm (EGFRxCD3:TT at Fig. 15a) and EGFR on the long arm (ThyroxCD3:EGFR at Fig. 15b). The bispecific antibody (EGFRxCD3) is used as a positive control.

35 Figure 16: T cell cytotoxicity in HCT116 cells (median EGFR expression) was determined by measuring ATP levels assessed by CellTiterGlo for both EGFRxCD3:TT (Fig. 16a) and ThyroxCD3:EGFR (Fig. 16b). The top charts ATP levels, measured by luminescence on an Envision Microplate reader results in Relative light unit (RLU) values, which were analyzed using GraphPad Prism, which the bottom chart correlates to percent killing based on the following equation, %Killing = (100- (RLU sample/RLU no IgG) x 100).

Figure 17: Effect of linkers on T-cell cytotoxicity in HCT116 cells (Fig. 17a).

Comparison of target cell lysis versus cytokine release for a range of linkers were demonstrated for IL-2 (Fig. 17b), IFN- $\gamma$  (Fig. 17c) and TNF- $\alpha$  (Fig. 17d).

Figure 18: Configuration of the CD3xPD-L1:EGFR trispecific T cell engager molecule  
5 with CD3 binding domain located on the short arm.

Figure 19: Configuration of the EGFRxCD3:PD-L1 trispecific T cell engager molecule  
with CD3 binding domain located on the internal region of the long arm.

Figure 20: T cell cytotoxicity activity data is provided against MDA-MB-231 cells  
comparing trispecific molecules combining a CD3 binding domain and two tumor cell antigen  
10 binding domains to trispecific controls with one tumor cell antigen binding domain, a mock  
domain and a CD3 binding domain, and the positive control described above.

Figure 21: T cell cytotoxicity activity data is provided against MDA-MB-231 cells  
comparing trispecific molecules combining a CD3 binding domain and two tumor cell antigen  
binding domains versus trispecific controls with one tumor cell antigen binding domain, a mock  
15 domain and a CD3 binding domain, where the trispecific molecules comprise tumor cell antigen  
binding domains comprising a range of affinities for targeting EGFR and PD-L1.

Figure 22: T cell cytotoxicity activity data is provided against HCT116 cells comparing  
trispecific molecules combining a CD3 binding domain and two tumor cell antigen binding  
domains versus trispecific controls with one tumor cell antigen binding domain, a mock domain  
20 and a CD3 binding domain, where the trispecific molecules comprise tumor cell antigen binding  
domains comprising a range of affinities for targeting EGFR and PD-L1.

Figure 23a: FACS data against MDA-MB-231 cells is shown as area under the curve  
(AUC) for a range of PD-L1 affinities and a range of EGFR affinities, demonstrating dual-antigen  
binding correlated with increasing affinity of the tumor antigen binding domains.

25 Figure 23b: T-cell cytotoxicity activity data is provided against BxPC3 cells  
demonstrating for certain trispecific molecules having the format of CD3xPD-L1:EGFR.  
Simultaneous dual antigen binding and immune cell engagement occurred with an additive  
effect on cytotoxicity over molecules binding a single antigen and CD3 (either CD3xEGFR:Mock  
or CD3xMock:PD-L1). Specific heavy chain sequences not shown.

30 Figure 24: Configuration of the EGFRxFibrinogen:CD3 trispecific T cell engager  
molecule with the CD3 binding domain located on the distal region of the long arm.

Figure 25: T-cell activation data is provided against HT29 cells, and demonstrating T-  
cell activation by a variety of EGFRxFibrinogen:CD3 trispecific T cell engager molecules using  
different CD3 binding domains as compared to the positive control EGFRxCD3 bispecific  
35 antibody used in Fig. 15.

Figure 26: Configuration of the EGFRxCD3:EGFR bispecific, trivalent molecule with the  
same EGFR binding domains (MF9891).

Figure 27: T cell activation activity in HCT116 cells was measured for a series of  
bispecific trivalent EGFRxCD3:EGFR molecules with the same EGFR binding domains

(MF9891) and different CD3 binding domains from different superclusters with a range of linkers.

Figure 28: T cell activation activity in MDA-MB-231 cells was measured for a series of bispecific trivalent EGFRxCD3:EGFR molecules with the same EGFR binding domains

5 (MF9891) and different CD3 binding domains from different superclusters with a range of linkers.

#### **Detailed description**

10 An "antibody" is a proteinaceous molecule belonging to the immunoglobulin class of proteins, containing one or more domains that bind an epitope on an antigen, where such domains are derived from or share sequence homology with the variable region of an antibody. Antibody binding has different qualities including specificity and affinity. The specificity determines which antigen or epitope thereof is specifically bound by the binding domain. The 15 affinity is a measure for the strength of binding to a particular antigen or epitope. It is convenient to note here that the 'specificity' of an antibody refers to its selectivity for a particular antigen, whereas 'affinity' refers to the strength of the interaction between the antibody's antigen binding site and the epitope it binds.

20 Thus, the "binding specificity" as used herein refers to the ability of an individual antibody binding site to react with an antigenic determinant. Typically, the binding site of the antibody of the invention is located in the Fab domains and is constructed from a hypervariable region of a heavy and/or light chains.

25 "Affinity" is the strength of the interaction between a single antigen-binding site and its antigen. A single antigen-binding site of an antibody of the invention for an antigen may be expressed in terms of the dissociation constant (KD). Typically, antibodies for therapeutic applications may have affinities of up to  $1 \times 10^{10}$  M or even higher.

30 An "antigen" is a molecule capable of inducing an immune response (to produce an antibody) in a host organism and/or being targeted by an antibody. At the molecular level, an antigen is characterized by its ability to be bound by the antigen-binding site of an antibody. Also mixtures of antigens can be regarded as an 'antigen', i.e. the skilled person would appreciate that sometimes a lysate of tumor cells, or viral particles may be indicated as 'antigen' whereas such tumor cell lysate or viral particle preparation exists of many antigenic determinants. An antigen comprises at least one, but often more, epitopes.

35 An "epitope" or "antigenic determinant" is a site on an antigen to which an immunoglobulin or antibody specifically binds. Epitopes can be formed from contiguous amino acids or noncontiguous amino acids juxtaposed by tertiary folding of a protein (so-called linear and conformational epitopes, respectively). Epitopes formed from contiguous, linear amino acids are typically retained on exposure to denaturing solvents, whereas epitopes formed by tertiary folding, conformation are typically lost on treatment with denaturing solvents. An epitope

may typically include 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15 amino acids in a unique spatial conformation.

The term "heavy chain" or "immunoglobulin heavy chain" includes an immunoglobulin heavy chain constant region sequence from any organism, and unless otherwise specified

5 includes a heavy chain variable domain. The term heavy chain variable domains include three heavy chain CDRs and four FR regions, unless otherwise specified. Fragments of heavy chains include CDRs, CDRs and FRs, and combinations thereof. A typical heavy chain has, following the variable domain (from N-terminal to C-terminal), a CH1 domain, a hinge, a CH2 domain, and a CH3 domain. A functional fragment of a heavy chain includes a fragment that is capable of  
10 specifically recognizing an antigen and that comprises at least one CDR.

The term "light chain" includes an immunoglobulin light chain variable domain, or  $V_L$ (or functional fragment thereof); and an immunoglobulin constant domain, or  $C_L$ (or functional fragment thereof) sequence from any organism. Unless otherwise specified, the term light chain may include a light chain selected from a human kappa, lambda, and a combination thereof.

15 Light chain variable ( $V_L$ ) domains typically include three light chain CDRs and four framework (FR) regions, unless otherwise specified. Generally, a full-length light chain includes, from N-terminus to C-terminus, a  $V_L$  domain that includes FR1-CDR1-FR2-CDR2-FR3-CDR3-FR4, and a light chain constant domain. Light chains that can be used with this invention include those, e.g., that do not selectively bind an epitope selectively bound by the heavy chains.

20 Suitable light chains for use in a multivalent antibody invention include a common light chain, such as those that can be identified by screening for the most commonly employed light chains in existing antibody libraries (wet libraries or in silico), where the light chains do not substantially interfere with the affinity and/or selectivity of the epitope-binding domains of the heavy chains, but are also suitable to pair with an array of heavy chains. For example, a  
25 suitable light chain includes one from a transgenic animal, such as a transgenic rodent, comprising the common light chain integrated into its genome and which can be used to generate large panels of common light chain antibodies having diversity at the heavy chain upon exposure to an antigen.

30 The term "common light chain" according to the invention refers to light chains which may be identical or have some amino acid sequence differences while the binding specificity of the antibody of the invention is not affected, i.e. the differences do not materially influence the formation of functional binding regions.

It is for instance possible within the scope of the definition of common chains as used herein, to prepare or find variable chains that are not identical but still functionally equivalent, 35 e.g., by introducing and testing conservative amino acid changes, changes of amino acids in regions that do not or only partly contribute to binding specificity when paired with a cognate chain, and the like. Such variants are thus also capable of binding different cognate chains and forming functional antigen binding domains. The term 'common light chain' as used herein thus refers to light chains which may be identical or have some amino acid sequence differences

while retaining the binding specificity of the resulting antibody after pairing with a heavy chain. A combination of a certain common light chain and such functionally equivalent variants is encompassed within the term "common light chain".

5 A "Fv domain" means a binding domain comprising a heavy chain variable region (VH) and a light chain variable region (VL).

A "Fab domain" means a binding domain comprising a variable region, typically a binding domain comprising a paired heavy chain variable region and light chain variable region. A Fab domain may comprise constant region domains, including a CH1 and a VH domain paired with a constant light domain (CL) and VL domain. Such pairing may take place, for 10 example, as covalent linkage via a disulfide bridge at the CH1 and CL domains.

A "modified Fab domain" means a binding domain comprising a CH1 and a VH domain, wherein the VH is paired with a VL domain and no CL domain is present. Alternatively, a modified Fab domain is a binding domain comprising a CL and a VL domain, wherein the VL is paired with a VH domain and no CH1 domain is present. In order that the CH1 or CL region can 15 be present in a non-paired form, it may be necessary to remove or reduce the lengths of regions of hydrophobicity. CH1 regions from species of animal that naturally express single-chain antibodies, for example from a camelid animal, such as a llama or a camel, or from a shark may be used. Other examples of a modified Fab domain include a constant region, CH1 or CL, which is not paired with its cognate region and/or a variable region VH or VL, is present, which 20 is not paired with its cognate region.

The term "immune effector cell" or 'effector cell' as used herein refers to a cell within the natural repertoire of cells in the mammalian immune system which can be activated to affect the viability of a target cell. Immune effector cells include cells of the lymphoid lineage such as natural killer (NK) cells, T cells including cytotoxic T cells, or B cells, but also cells of the myeloid 25 lineage can be regarded as immune effector cells, such as monocytes or macrophages, dendritic cells and neutrophilic granulocytes. Hence, said effector cell is preferably an NK cell, a T cell, a B cell, a monocyte, a macrophage, a dendritic cell or a neutrophilic granulocyte.

"Percent (%) identity" as referring to nucleic acid or amino acid sequences herein is defined as the percentage of residues in a candidate sequence that are identical with the 30 residues in a selected sequence, after aligning the sequences for optimal comparison purposes. In order to optimize the alignment between the two sequences gaps may be introduced in any of the two sequences that are compared. Such alignment can be carried out over the full length of the sequences being compared. Alternatively, the alignment may be carried out over a shorter length, for example over about 20, about 50, about 100 or more nucleic acids/based or amino 35 acids. The sequence identity is the percentage of identical matches between the two sequences over the reported aligned region.

A comparison of sequences and determination of percentage of sequence identity between two sequences can be accomplished using a mathematical algorithm. The skilled person will be aware of the fact that several different computer programs are available to align

two sequences and determine the identity between two sequences (Kruskal, J. B. (1983) An overview of sequence comparison In D. Sankoff and J. B. Kruskal, (ed.), Time warps, string edits and macromolecules: the theory and practice of sequence comparison, pp. 1 -44 Addison Wesley). The percent sequence identity between two amino acid sequences or nucleic acid

5 sequences may be determined using the Needleman and Wunsch algorithm for the alignment of two sequences. (Needleman, S. B. and Wunsch, C. D. (1970) J. Mol. Biol. 48, 443-453). The Needleman-Wunsch algorithm has been implemented in the computer program NEEDLE. For the purpose of this invention the NEEDLE program from the EMBOSS package is used to determine percent identity of amino acid and nucleic acid sequences (version 2.8.0 or higher, 10 EMBOSS: The European Molecular Biology Open Software Suite (2000) Rice, P. LongdenJ. and Bleasby, A. Trends in Genetics 16, (6) pp276— 277, <http://emboss.bioinformatics.nl/>). For protein sequences, EBLOSUM62 is used for the substitution matrix. For DNA sequences, DNAFULL is used. The parameters used are a gap-open penalty of 10 and a gap extension penalty of 0.5.

15 After alignment by the program NEEDLE as described above the percentage of sequence identity between a query sequence and a sequence of the invention is calculated as follows: Number of corresponding positions in the alignment showing an identical amino acid or identical nucleotide in both sequences divided by the total length of the alignment after subtraction of the total number of gaps in the alignment.

20 Herein, the term "connected" refers to domains which are joined to each other by way of peptide bonds at the primary amino acid sequence. For example, a heavy chain of a base antibody portion comprising VH-CH1-CH2-CH3 may be connected to a heavy chain of an additional binding domain VH-CH1 (or an additional binding domain to an additional binding domain) via a linker (connecting the heavy chain of the additional binding domain at the CH1 to 25 the VH region of the base antibody portion), which together constitutes one polypeptide chain. Similarly, a CH1 domain may be connected to a variable heavy region and a CL domain may be connected to a variable light region.

30 "Pairing" refers to interactions between the polypeptides constituting a multivalent antibody of the invention such that they may multimerize. For example, an additional binding domain may comprise a heavy chain region (VH-CH1) paired to a light chain region (VL-CL), where the CH1 and CL pair to form said binding domain. As described herein, pairing of antibody domains (e.g., heavy and light) occurs due to noncovalent interactions and also via disulphide bonds, and can be engineered through techniques disclosed herein and by methods known in the art.

35 Throughout the present specification and the accompanying claims, the words "comprise", "include" and "having" and variations such as "comprises", "comprising", "includes" and "including" are to be interpreted inclusively. That is, these words are intended to convey the possible inclusion of other elements or integers not specifically recited, where the context allows.

The articles "a" and "an" are used herein to refer to one or to more than one (i.e. to one or at least one) of the grammatical object of the article. By way of example, "an element" may mean one element or more than one element.

5                   The invention provides a multivalent antibody which comprises:

- a base antibody portion which comprises two binding domains; and
- at least one additional binding domain,

10                  wherein the base antibody portion is connected by a linker to the at least one additional binding domain,

                        wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region, and

15                  wherein the linker comprises a hinge sequence or a sequence derived from a hinge sequence.

20                  The invention also provides a multivalent antibody which comprises:

- a base antibody portion which comprises two binding domains; and
- at least one additional binding domain,

25                  wherein at least one additional binding domain comprises a CH1 region and is connected to the base antibody portion by said linker, linking a variable region of the base antibody portion and the CH1 region, and

                        wherein the multivalent antibody binds to at least three different epitopes.

30                  In such a multivalent antibody, each binding domain of the base antibody portion and each of the at least one additional binding domains may all have a common variable region,

                        The invention thus provides a multivalent antibody which is typically capable of binding to its target or targets via at least three binding domains, i.e. the antibody is a multivalent antibody. The multivalent antibody may optionally be a multispecific antibody. That is to say, an antibody of the invention may be capable of binding two or more different epitopes or two or more different antigens, for example two, three, four or more different epitopes or antigens.

Different Formats of the Multivalent Antibodies

It should be noted that other features and aspects of the invention are apparent from the detailed description, taken in conjunction with the accompanying drawings, which illustrate, 5 by way of example, the features in accordance with embodiments of the invention. The figures are exemplary and are not intended to nor do they limit the scope of the invention, which is defined by the claims and the full extent of the detailed disclosure, which describe and enable the inventions set out herein. A multivalent antibody of the invention may comprise a base antibody portion and an additional binding domain, preferably a Fab domain comprising a VH- 10 CH1 region paired to a VL-CL region. Said multivalent antibody comprises three VH regions, and three VL regions. Either of the VH or VL may be a common variable region (VHc or VLC) paired to a rearranged variable region of the cognate chain. For example, the three VL regions may be a common chain (VLC), and each VH region (VH1-VH3) may comprise a rearranged variable region, wherein said VH1, VH2 and VH3 regions may bind the same epitope or up to 15 three different epitopes. Wherein, the multivalent antibody comprises a common light chain (VLC) and three heavy chain variable regions (VH1-VH3), the additional Fab domain comprised of a VH3-CH1 paired with a VLC-CL may be connected to the base antibody via a linker positioned between a VH1 or VH2 region of the base antibody portion and CH1 of the additional Fab domain. See, for example, Figure 1a.

20 Alternatively, the additional Fab domain may be connected to the base antibody via a linker positioned between the common light chain region (VLC) of the base antibody and the CL region of the additional Fab domain. See, for example, Figure 1b. In another aspect of the invention, the three VH regions may be a common chain (VHc), and each VL region may comprise a rearranged variable region, wherein said three VL regions may bind the same or 25 differing epitopes (VL1-VL3). Wherein the multivalent antibody comprises a common heavy chain (VHc) and three light chain variable regions (VL1-VL3), the additional Fab domain may be connected to the base antibody via a linker positioned between a VL1 or VL2 region of the base antibody portion and CL of the additional Fab domain. See, for example, Figure 1c. Alternatively, the additional Fab domain may be connected to the base antibody via a linker 30 positioned between the common heavy chain region (VHc) of the base antibody and the CH1 region of the additional Fab domain. See, for example, Figure 1d.

35 Alternatively, the additional Fab domain may be connected to the base antibody via a linker positioned between both the heavy and light variable regions of the base antibody and the CH1 and CL regions of the additional Fab domain, irrespective of whether the common chain is heavy or light. See, for example, Figure 1e.

A multivalent antibody of the invention may comprise a base antibody portion and more than one additional binding domain, for example two Fab domains. Either the VH or VL regions of said multivalent antibody may be a common variable region (e.g., VHc or VLC) with the cognate chain comprising a rearranged variable region binding the same or different antigen or

epitope (e.g., VHc and VL1-VL4; or VH1-VH4 and VLc). The additional Fab domains may be connected to the base antibody portion via a linker positioned between the common chain of the base antibody portion (VHc or VLc) and the respective constant region of the common variable region of the additional Fab domains, or the rearranged variable domains (VH1 and VH2; or VL1 and VL2) of the base antibody and the respective constant region of the rearranged variable domain of the additional Fab domains. For example, Figure 1f depicts a multivalent antibody of the invention comprising a base antibody and two additional Fab domains, wherein the antibody comprises a common light chain (VLc), and four heavy chain variable regions (VH1-VH4), wherein a linker connects the base antibody to the additional Fab domains at rearranged heavy chain variable regions of the base antibody (VH2 and VH3) and the CH1 regions of the additional Fab domains. Alternatively, Figure 1g depicts a multivalent antibody of the invention, wherein the base antibody is connected to two additional Fab domains at a first rearranged heavy chain region (VH2) to the CH1 region of the first additional Fab domain, and a common light chain variable region (VLc) of the base antibody to the CL region of the second additional 15 Fab domain. Alternatively, Figure 1h depicts a multivalent antibody of the invention, wherein the base antibody is connected to two additional Fab domains via a linker connecting both common light chain regions (VLc) of the base antibody to the CL regions of the two additional Fab domains. Alternatively, Figure 1j depicts a multivalent antibody of the invention, wherein the base antibody is connected to a first additional Fab domain via a linker connecting both the 20 rearranged heavy chain variable region (VH3) and the common light chain region (VLc) of the base antibody to the first additional Fab domain at CH1 and CL respectively, and the second additional Fab domain is connected via a linker to the second rearranged heavy chain variable region (VH2) of the base antibody at the CH1 region of the second additional Fab domain. Alternatively, (Fig. 1i) the second additional Fab domain is connected via a linker to the common 25 light chain variable region (VLc) of the base antibody to the CL region of the second additional Fab domain. Alternatively, (Fig. 1k), the second additional Fab domain is connected via a linker to both the second rearranged heavy chain variable region (VH2) and the common light chain (VLc) of the base antibody portion at the CH1 and CL regions of the second additional Fab domain, respectively. The formats described herein and depicted at Figures 1f-1k also apply to 30 where the common chain is a heavy chain (VHc) and the multivalent antibody comprises four rearranged light chain variable regions (VL1-VL4) comprising up to four different binding specificities.

Further, two or more additional binding domains may be connected via linkers to only one binding domain of a base antibody portion, such that a first Fab domain is connected to a 35 second Fab domain via a linker, which is then connected to the base antibody portion. That is to say, a first linker is positioned between the base antibody portion and one of the additional Fab domains and a second linker is positioned between the two additional Fab domains. The two linkers may be the same or different.

In another aspect of the invention, the individual proteins that make up the multivalent antibody can mix heavy and light chains within the same protein. For example, a multivalent antibody may be comprised of a first protein comprising the additional Fab domain linked to the base antibody in order from N-terminus to C-terminus of VLc-CL-VH2-CH1-CH2-CH3, such that 5 a linker connects the VLc-CL region of the additional Fab domain to the base antibody portion at VH2-CL. A second protein comprising VH1-CH1, which pairs with the VLc-CL of the first protein. A third protein comprising in order from N-terminus to C-terminus VH3-CH1-CH2-CH3 such that the third and first proteins pair below their respective CH1 regions. And a fourth 10 protein comprises in order from N-terminus to C-terminus VLc-CL, which pairs with the VH2-CH1 of the first protein and VH3-CH1 of the third protein. See for example, Figure 1l.

While the format described and depicted in Figure 1l illustrates use of a common light chain and at least three rearranged heavy chain variable regions (VH1-VH3) comprising up to three different binding specificities, it should be understood that this format applies where the 15 common chain is a heavy chain (VHc) and the multivalent antibody comprises three or more rearranged light chain variable regions (VL1-VL3) comprising up to three different binding specificities.

Another aspect of the invention is a multivalent antibody comprising four proteins, where the common chain is a common light chain. The multivalent antibody is comprised of 20 four proteins in order of N-terminus to C-terminus comprising: a first protein of VH1-CH1-VLc-CL, wherein a linker connects the CH1 to VLc; a second protein of VLc-CL that pairs with the VH1-CH1 to form an additional Fab domain; a third protein comprising VH2-CH1-CH2-CH3, wherein the CH1 of the third protein pairs with the CL of the second protein; and a fourth protein comprising VH3-CH1-CH2-CH3, wherein the third and fourth proteins are paired below the CH1 region, and the second protein (VLc-CL) is paired with the fourth protein at the CH1 region of 25 the fourth protein. See, for example, Figure 1m.

A multivalent antibody of the invention may comprise a base antibody portion and an additional Fab domain, wherein either the VH or VL regions of the multivalent antibody may be a common variable region, and wherein the additional Fab domain may be connected to the base antibody portion via a linker positioned either at the VH (Figure 1n) or VL (Figure 1o) of the base 30 antibody, wherein said linker simultaneously connects the base antibody to the Fab domain and also pairs the cognate chains of the Fab domain. In such instances said Fab domain may optionally lack a CH1-CL domain, and use the linker to pair the variable domains of the Fab domain.

A multivalent antibody of the invention may comprise a base antibody portion and an 35 additional binding domain which comprises a paired VH and VL. Said additional binding domain which comprises a VH and VL may be paired via a cysteine bridge, formed between the VH and VL, such that it may not require the presence of a CH1 or CL region. See, for example, Figure 1p. Note, a cysteine bridge is depicted in figure 1p, although the person of ordinary skill in the

art understands that additional cysteine bridges are typically present at the CH1/CL interface (not shown in the figures).

A multivalent antibody of the invention may comprise a base antibody portion and an additional modified Fab domain. The modified Fab domain may comprise a modified CH1 such 5 that it does not need to pair with a CL. For example, the CH1 could be a camelid CH1 or based on a camelid CH1, or be modified to lack hydrophobic residues through techniques known in the art. Each VH or VL may be a common or rearranged variable region. The additional modified Fab domain may be connected to the base antibody portion via a linker positioned between the VH2 of the base antibody portion and CH1 of the modified Fab domain. The VH and VL of the 10 modified Fab domain may be paired via a cysteine bridge, or alternatively non-covalent interactions. See, for example, Figure 1q. Alternatively, the additional modified Fab domain may be connected to the base antibody portion via a linker positioned between the VL of the base antibody portion and CH1 of the modified Fab domain. The VH and VL of the modified Fab domain may be paired via a cysteine bridge. See, for example, Figure 1r.

15 A multivalent antibody of the invention may comprise a base antibody portion and an additional modified Fab domain, wherein the modified Fab domain may comprise a modified CL such that it does not need to pair with a CH1. For example, the CL could be engineered to remove hydrophobic regions. Each VH or VL of the modified Fab domain may be a common or rearranged variable region. The additional modified Fab domain may be connected to the base 20 antibody portion via a linker positioned between the VL of the base antibody portion and CL of the modified Fab domain. The VH and VL of the modified Fab domain may be paired via a cysteine bridge. See, for example, Figure 1s.

A multivalent antibody of the invention may comprise a base antibody portion and an additional modified Fab domain, wherein the modified Fab domain may comprise a modified CL 25 such that it does not need to pair with a CH1. For example, the CL could be engineered to remove hydrophobic regions. Each VH or VL of the modified Fab domain may be a common or rearranged variable region. The additional modified Fab domain may be connected to the base antibody portion via a linker positioned between the VH2 of the base antibody portion and CL of the modified Fab domain. The VH and VL of the modified Fab domain may be paired via a 30 cysteine bridge. See, for example, Figure 1t.

#### Base antibody portion of the invention

It should be noted that while figures 1a-1u depict a base antibody portion of the 35 multivalent antibody as including paired heavy chain constant regions comprising CH2 and CH3 regions, these regions are shown merely for illustrative purposes and the invention is not limited to these embodiments. Herein is described different formats for the base antibody portion and additional binding domain suitable for use in the antibodies disclosed.

The base antibody portion of the multivalent antibody of the invention may be a full length immunoglobulin, for example a full length IgG, IgA, IgE, IgD or IgM portion, but preferably IgG, and more preferably IgG1.

In any antibody of the invention, at least one of the additional binding domains, 5 preferably a Fab domain, may comprise a CH1 domain of an immunoglobulin subclass different from that of the CH1 domain(s) of the base antibody portion of the antibody and/or may have a light chain of a different class. For example, where the base portion of the antibody is a full length IgG1, at least one of the additional binding domains may comprise a CH1 domain of the subclass IgG2a, IgG2b, IgG3 or IgG4 and/or where the base portion of the antibody includes a 10 kappa light chain, at least one of the additional binding domain may include a lambda light chain.

The heavy chains of the base antibody may be designed to preferentially pair through techniques known to those of skill in the art, such as engineering the DEKK modifications in the 15 CH3 regions of the base antibody. See WO2013/157954 and De Nardis et al., *J. Biol. Chem.* (2017) 292(35) 14706-14717 incorporated herein by reference, demonstrating engineering in the CH3 region for driving heterodimerization of the heavy chains. Alternative approaches for driving heterodimerization which may be used in the invention include the knob-in-hole format (WO1998/050431) and use of charge engineering (Gunasekaran, *JBC* 2010, vol 285, pp 19637-19646).

20 The Fc region mediates effector functions of an antibody, such as complement-dependent cytotoxicity (CDC), antibody-dependent cellular cytotoxicity (ADCC) and antibody-dependent cell phagocytosis (ADCP). Depending on the multivalent antibody, it may be desired to either reduce or increase the effector function. Reduced effector function can be desired when an immune response is to be activated, enhanced or stimulated as in some of the 25 embodiments of the invention. Antibodies with reduced effector functions can be used to target cell-surface molecules of immune cells, among others. Increased effector function can be desired when an antibody is targeting harmful cells, thereby boosting the ability of immune effector cells or the complement cascade to eliminate or lyse such targets.

The effector function of the heavy chain Fc region can be mitigated or eliminated 30 through modifications known to those of ordinary skill in the art. Similarly, the effector function of the heavy chain Fc region can be enhanced through modifications known to those of ordinary skill in the art. For example, ADCC may be enhanced via the genetic modification of the CH2 domain. See, for example, Strohl, *Curr. Opin. Biotechnol.* 2009 (6) 685-91.

A multivalent antibody of the invention can in one embodiment be afucosylated. A 35 multivalent antibody of the invention preferably comprises a reduced amount of fucosylation of the N-linked carbohydrate structure in the Fc region, when compared to the same antibody produced in a normal CHO cell.

An aspect of the multivalent antibody of the invention also includes a base antibody, which lacks a CH2 or CH3 region, wherein said heavy chains of the base antibody portion may be joined by a cysteine covalent bridge below the CH1 region.

Aspects of the invention, including variations to the base antibody portion of the invention are illustrated at Figure 1u.

Herein is described a repertoire of linkers that can be used to connect the base portion of the antibody of the invention with one or more binding domains. One or more binding domains, such as a variable region, an Fv domain, a Fab domain or a modified Fab domain, may be connected to the base antibody portion of an antibody of the invention.

The antibody of the invention comprises a base antibody portion and, attached thereto, via a linker or linkers, one or more binding domains.

Multivalent antibodies comprising a full length IgG base antibody portion are preferred because such structures typically have beneficial properties such as a favorable half-life, predictable biophysical behavior and lower immunogenicity. Antibodies of the invention are typically suitable for therapeutic use and therefore are comprised of human sequences for the use of human therapeutics. Alternatively, said antibodies have sequences of the species for which the therapeutic is being used or based on consensus sequences within that given species, using techniques well known to those of ordinary skill in the art.

Where the base antibody portion of an antibody of the invention is a full length IgG, the full length IgG may comprise mutations that provide desired characteristics. Such mutations are typically not deletions of substantial portions of any of the regions. However, full-length IgG portions wherein one or several amino acid residues are inserted, deleted or substituted, without essentially altering the binding characteristics of the resulting IgG portion, are embraced within the term "full length IgG". For instance, such IgG portions can have one or more insertions, deletions or substitutions of between 1 and 10 amino acid residues, preferably in non-CDR regions, wherein the inserted, deleted or substituted amino acids are not essential for the binding specificity of the IgG.

IgG1 may be favored based on its long circulatory half-life in man. Also, in order to mitigate immunogenicity in humans, it is preferred that the base antibody portion of an antibody according to the invention is a human antibody.

The base portion of the antibody of the invention may be a full length immunoglobulin which is defined as comprising an essentially complete antibody. Such an essentially complete antibody may not necessarily have all the functions of an intact antibody.

A full length base portion of an antibody as described herein comprises two heavy and two light chains. Each chain contains constant (C) and variable (V) regions, which can be broken down into domains designated CH1, CH2, CH3, V<sub>H</sub> for the heavy chain, and C<sub>L</sub>, V<sub>L</sub> for the light chain. The antibody can interact with molecules and cells of the immune system through the constant domains, typically through the Fc portion.

The constant region of an antibody of the present invention, including a bispecific or multispecific antibody, is preferably a human constant region. The constant region may contain one or more, preferably not more than 10, preferably not more than 5 amino-acid differences with the constant region of a naturally occurring human antibody. Various variable domains of 5 antibodies produced herein are derived from a human antibody variable domain library. As such these variable domains are human. The unique CDR regions may be derived from humans, be synthetic or derived from another organism. An antibody or bispecific antibody of the invention is preferably a human or humanized antibody. Suitable heavy chain constant regions are non-limitingly exemplified in Table 21.

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An antibody of the invention typically has an intact Fc region that maintains half-life and stability of the multispecific antibody. The Fc may also allow interaction with immune effector molecules such as Fc receptors, complement and FcRn. As understood by persons of skill in the art, techniques are available to design an Fc region to prevent or mitigate interactions with 15 Fc receptors or to enhance interactions with Fc receptor.

The base antibody portion and one or more additional binding domains, for example, Fab domains are connected via one or more linkers. The at least one additional Fab domain may be of a given isotype or subclass, eg, IgG1, 2a, 2d, 3 or 4; at least one additional Fab may be of a different subclass to that of the Fab domains of the full length IgG portion or may carry a 20 light chain of a different class (kappa or lambda).

#### Linkers for use in the Multivalent Antibody Format

An antibody of the invention comprises one or more linkers which connect the one or 25 more additional binding domains to the base antibody portion. The linker together with the binding domain to which the linker is connected determines, at least in part, the functionality of the multivalent antibody.

In an antibody of the invention, the peptide region of a linker may comprise a hinge sequence or comprise a sequence based on a hinge sequence. Thus, the amino acid 30 sequence of a suitable peptide region may comprise a naturally-occurring sequence or comprise a sequence based on a naturally-occurring sequence. The use of such sequences may help developability of multivalent antibodies of the invention and help to ensure low immunogenicity.

A hinge region is a flexible amino acid stretch in the central part of the heavy chains of 35 the IgG and IgA immunoglobulin classes (i.e., that portion which connects the Fab to the Fc), which pairs these two heavy chains by disulfide bonds. It is rich in cysteine and proline amino acids, and bears little resemblance to any other immunoglobulin region.

Accordingly, a suitable linker to connect the one or more additional binding domains to the base antibody portion for use in a multivalent antibody of the invention may be derived from

an IgG or IgA hinge sequence. The linker region may be based on an IgG1 hinge region, an IgG2 hinge region, an IgG3 hinge region or an IgG4 hinge region.

Typically, the type of the hinge region used is matched with the type of the constant region, for example the CH1, of the additional Fab domain to which the linker is connected.

5 That is to say, if a linker is based on a sequence or sequences from a IgG1 hinge region, the CH1 of the additional Fab domain to which it is connected is a CH1 from a IgG1.

A linker of an antibody may be based on an upper, middle or lower hinge region, or a subset of such a region.

The IgG1 hinge region has the sequence: EPKSCDKTHTCPCPAELLGG (SEQ ID NO: 42).

The upper hinge region is defined as: EPKSCDKTHT (SEQ ID NO: 43)

The middle hinge region is defined as: CPPCP (SEQ ID NO: 44)

The lower hinge region is defined as: APELLGG (SEQ ID NO: 45)

Thus, in an antibody of the invention, the linker may comprise one or more of these sequences and/or a sequence based on one or more of these sequences.

The IgG2 hinge region has the sequence: ERKCVECPPCPAPPVAG (SEQ ID NO: 46).

The upper hinge region is defined as: ERKCCVE (SEQ ID NO: 47)

The middle hinge region is defined as: CPPCP (SEQ ID NO: 48)

20 The lower hinge region is defined as: APPVAG(SEQ ID NO: 49)

Thus, in an antibody of the invention, the linker may comprise one or more of these sequences and/or a sequence based on one or more of these sequences.

The IgG3 hinge region has the sequence:

The upper hinge region is defined as: ELKTPPLGDTTHT (SEQ ID NO: 51)

The middle hinge region is defined as:

CPRCPEPKSCDTPPPCPRCPEPKSCDTPPPCPRCPEPKSCDTPPPCPRC (SEQ ID NO: 52)

The lower hinge region is defined as: APEFLGG (SEQ ID NO: 53)

30 The IgG4 hinge region has the sequence: ESKYGPPCPSCPAPAEFLGG (SEQ ID NO:  
54).

The upper hinge region is defined as: ESKYGPP (SEQ ID NO: 55)

The middle hinge region is defined as: CPSCP (SEQ ID NO: 56)

The lower hinge region is defined as: APEFLGG (SEQ ID NO: 57).

35 The middle region with consensus sequence CXXC connects both IgG heavy chains in the context of a wildtype IgG and is rigid. These disulfide bridges are not required for the current application and, therefore, where a linker comprises a middle hinge sequence, preferably, one or both Cys residues in the CXXC concensus are substituted, for example with a Ser residue. Thus, in a preferred embodiment CxxC may be SxxS.

A linker suitable for use in a multivalent antibody of the invention may be one based on a middle hinge sequence, for example a sequence which comprises a middle hinge sequence, but which does not comprise a lower and/or an upper hinge sequence. A linker suitable for use in a multivalent antibody of the invention may be one based on an upper hinge sequence, for 5 example a sequence which comprises an upper hinge sequence, but which does not comprise a lower and/or a middle hinge sequence. A linker suitable for use in a multivalent antibody of the invention may be one which does not comprise a middle hinge sequence, for example a sequence which comprises a combination of lower and upper hinge sequences.

Accordingly, the invention provides a linker comprising an amino acid sequence of one 10 of SEQ ID NOs: 3 to 5, 7 to 11 or 13 to 24.

Thus, in an antibody of the invention, the linker may comprise one or more of these sequences and/or a sequence based on one or more of these sequences. A peptide region may consist essentially of a middle region sequence or be based on such a sequence or consist essentially of an upper and a lower region sequence or be based on such sequences.

15 A linker suitable for use in an antibody of the invention may be defined with reference to a sequence comprising the amino acid sequence of any linker sequence as set out herein in which from 0 to 5 amino acid insertions, deletions, substitutions or additions (or a combination thereof) is made. In some embodiments, the linker comprises an amino acid sequence comprising from 0 to 4, preferably from 0 to 3, preferably from 0 to 2, preferably from 0 to 1 and 20 preferably 0 amino acid insertions, deletions, substitutions or additions (or a combination thereof) with respect to a linker sequence as set out herein.

25 A suitable linker may be from about 7 to about 29 amino acids in length, for example from about 10 to about 20 amino acids in length. However, a suitable linker may be a short linker, for example from about 7 to about 10 amino acids in length or may be a long linker, for example from about 20 to about 29 amino acids in length.

The linker may comprise an Ig hinge region or comprise a sequence based on an IgG hinge region connected to a CH1 region of the same subclass as the linker and may comprise cysteines for covalent linkage of the common light chain.

30 A linker suitable for use in an antibody of the invention may be based on an IgG1 hinge region, an IgG2 hinge region, an IgG3 hinge region or an IgG4 hinge region.

If a  $(G_4S)_n$  sequence is to be used, preferably it is used in combination with a hinge sequence from an isotype other than IgG or a subclass other than IgG1 and includes a CH1 region.

35 In an antibody of the invention, the linker may be rigid or flexible may comprise a charged sequence, may be straight or bent.

A rigid sequence for the purposes of this invention is sequence having a Karplus and Schulz flexibility Prediction of about 1.015 or less. A partially flexible sequence is one having a Karplus and Schulz flexibility Prediction of from about 1.015 to about 1.04. A flexible sequence for the purposes of this invention is sequence having a Karplus and Schulz flexibility Prediction

of at least about 1.015 (Karplus PA, Schulz GE. Prediction of Chain Flexibility in Proteins - A tool for the Selection of Peptide Antigens. *Naturwissenschaften* 1985; 72:212-3; <http://tools.immuneepitope.org/bcell/>). The flexibility prediction is calculated over consecutive windows of 7 residues along the sequence (1 residue step) yielding the predicted "flexibility" 5 index per window. The overall flexibility over the linker sequence is given as the average over the whole sequence.

Removal or Substitution of Cys residues in an IgG hinge region may make a linker based on that hinge more flexible including through replacement of the Cys residue with a serine (Ser). Alternatively, a linker may be a rigid linker in view of the presence of a helix-10 forming sequence. Accordingly, a middle hinge region, for example the conserved CPPCP motif, may be replaced by a helix-forming sequence, for example (EAAAK)<sub>2</sub>, which will result in a short rigid helix in the linker. Therefore, in an antibody of the invention, the linker may comprise a helix-forming sequence, for example comprising the amino acid sequence (EAAAK)<sub>2</sub>. The use of such a sequence may help to add rigidity.

15 A linker of the invention may, preferably comprise an amino acid sequence as set out in any one of SEQ ID NOs: 3 to 5, 7 to 11 or 13 to 24 or an amino acid sequence having at least about 90% sequence identity to any one thereto, preferably at least about 95% sequence identity to any one thereto, more preferably at least 97% sequence identity to any one thereto, more preferably at least about 98% sequence identity to any one thereto, more preferably at 20 least about 99% sequence identity to any one thereto.

For example, a linker suitable for use in a multivalent antibody of the invention may be defined with reference to a sequence comprising the amino acid sequence of any one of SEQ ID NOs: 1 to 24 in which from 0 to 5 amino acid insertions, deletions, substitutions or additions (or a combination thereof) is made. In some embodiments, the linker comprises an amino acid 25 sequence having from 0 to 4, preferably from 0 to 3, preferably from 0 to 2, preferably from 0 to 1 and preferably 0 amino acid insertions, deletions, substitutions or additions (or a combination thereof) with respect to a sequence set out in SEQ ID NOs: 3 to 5, 7 to 11 or 13 to 24.

A linker suitable for use in a multivalent antibody of the invention may be defined with 30 reference to a sequence comprising the amino acid sequence of any one of SEQ ID NOs: 1 to 24 or an amino acid sequence having at least about 85% sequence identity to any one thereto, such as at least about 90% sequence identity to any one thereto, for example at least about 95% sequence identity to any one thereto, such as at least about 98% sequence identity to any one thereto, for example at least about 99% sequence identity to any one thereto.

Table 1 illustrates how a linker sequence may be connected to CH1 and VH2 regions.

**Table 1: The underlined sequence is the linker sequence; the flanking sequences are the CH1 region of the additional Fab's CH1 region and the VH2 region of a following heavy chain VH region**

#	Name	Linker sequences (underlined) containing CH1 region and VH sequence preceding and following the linker sequence respectively. In the control IgG1 sequence the CH2 region is present (underlined)
1	IgG1 H	NVNHKPSNTKVDKR <u>VEPKSCDKTHTSPPS</u> PAELLGGEVQLVESGG GVVQPG (SEQ ID NO: 58)
2	IgG1 MH	NVNHKPSNTKVDKR <u>VEPKSCDKTHTSPPS</u> PEVQLVESGGGVQPG (SEQ ID NO: 59)
3	IgG1 UH	NVNHKPSNTKVDKR <u>VEPKSCDKTHT</u> EVQLVESGGGVQPG (SEQ ID NO: 60)
4	IgG1 G4S	NVNHKPSNTKVDKR <u>VEPKSCDGGGGSGGG</u> SEVQLVESGGGVQ PG (SEQ ID NO: 61)
5	IgG1 R	NVNHKPSNTKVDKR <u>VEPKSCDKTHT</u> EA AAKEAAAKA <u>PELLGGEVQL</u> VESGGGVQPG (SEQ ID NO: 62)
6	IgG1 UL	NVNHKPSNTKVDKR <u>VEPKSCDKTHT</u> APE <u>LLGGEVQL</u> VESGGGVQPG (SEQ ID NO: 63)
7	IgG2A H	NVDHKPSNTKVDKT <u>VERKSSVESPPSAPPVAGE</u> VQLVESGGGV QPG (SEQ ID NO: 64)
8	IgG2A MH	NVDHKPSNTKVDKT <u>VERKSSVESPPSPEVQL</u> VESGGGVQPG (SEQ ID NO: 65)
9	IgG2A UL	NVDHKPSNTKVDKT <u>VERKSSVEAPPVAGE</u> VQLVESGGGVQPG (SEQ ID NO: 66)
10	IgG2B H	NVDHKPSNTKVDKT <u>VERKCSVESPPSAPPVAGE</u> VQLVESGGGV QPG (SEQ ID NO: 67)
11	IgG2B MH	NVDHKPSNTKVDKT <u>VERKCSVESPPSPEVQL</u> VESGGGVQPG (SEQ ID NO: 68)
12	IgG2B UL	NVDHKPSNTKVDKT <u>VERKCSVEAPPVAGE</u> VQLVESGGGVQPG (SEQ ID NO: 69)
13	IgG2A G4SL	NVDHKPSNTKVDKT <u>VGCGGSGGGGSAPPVAGE</u> VQLVESGGGVQ PG (SEQ ID NO: 70)
14	IgG2A G4SS	NVDHKPSNTKVDKT <u>VGCGGSGGG</u> SEVQLVESGGGVQPG (SEQ ID NO: 71)
15	IgG2A R	NVDHKPSNTKVDKT <u>VERKSSVEAAKEAAAKAPPVAGE</u> VQLVESGG GVQPG (SEQ ID NO: 72)
16	IgG2B R	NVDHKPSNTKVDKT <u>VERKCSVEAAKEAAAKAPPVAGE</u> VQLVESGG GVQPG (SEQ ID NO: 73)
17	IgG3 ULH	NVNHKPSNTKVDKR <u>VELKTPLGDTT</u> TAPE <u>FLGGEVQL</u> VESGGGVQPG (SEQ ID NO: 74)

18	IgG3 UH	NVNHKPSNTKVDKRVE <u>LKTPLGDTTH</u> EVQLVESGGGVQPG (SEQ ID NO: 75)
19	IgG3 R	NVNHKPSNTKVDKRVE <u>LKTPLGDTTH</u> TEAAAKEAAAKAPE <u>FLGGEV</u> QLVESGGGVQPG (SEQ ID NO: 76)
20	IgG4 H	NVDHKPSNTKVDKRVE <u>SKYGPSPSSPAPEFLGGEV</u> QLVESGGGVQPG (SEQ ID NO: 77)
21	IgG4 MH	NVDHKPSNTKVDKRVE <u>SKYGPSPSSPEV</u> QLVESGGGVQPG (SEQ ID NO: 78)
22	IgG4 UL	NVDHKPSNTKVDKRVE <u>SKYGP</u> PAPE <u>FLGGEV</u> QLVESGGGVQPG (SEQ ID NO: 79)
23	IgG4 UH	NVDHKPSNTKVDKRVE <u>SKYGPPEV</u> QLVESGGGVQPG (SEQ ID NO: 80)
24	IgG4 R	NVDHKPSNTKVDKRVE <u>SKYGPPEAAAKEAAAKAPEFLGGEV</u> QLVESGGGVQPG (SEQ ID NO: 81)
25	IgG1 hinge	NVNHKPSNTKVDKRVE <u>PKSCDKTHCP</u> PCPAPE <u>LLGGPSVFLFPPK</u> PKDTLM (SEQ ID NO: 82)

Note, the VH2 sequence, following the linker (underscored above) may vary, depending on the specific variable region used. In other embodiments, the sequence following the linker may be a light chain variable region, including a common light chain.

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#### Use of linkers to pair regions of the additional binding domain

The linkers used herein may connect the base antibody portion to the at least one additional binding domain. In addition, where the at least one additional binding domain is a Fab domain or is comprised of pairing of a heavy chain variable region and a light chain variable region, the linker may pair the heavy and light chains via covalent linkage, typically via a disulphide bridge. The disulphide bridge may form between a cysteine residue in the linker and a variable region of the additional binding domain(s). Such pairing caused by the linker may apply to an additional binding domain, comprising a Fab domain comprising a common light chain and a counterpart rearranged heavy chain variable region or comprising a common heavy chain and a counterpart rearranged light chain variable region.

#### Multivalency and multispecificity

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Where the two binding domains of the base antibody of a multivalent protein of the invention bind different antigens, said first and second antigens may be two different molecules or moieties that are located on one cell or on different cell types. Antibodies comprising two binding domains that mediate cytotoxicity by recruiting and activating endogenous immune cells are an emerging class of antibody therapeutics. This can be achieved by combining antigen binding specificities for target cells (i.e., tumor cells) and effector cells (i.e., T cells, NK cells, and

macrophages) in one molecule (see, for example, WO2014/051433). An antibody of the invention comprises at least three binding domains. The base antibody portion will typically comprise two different binding domains (although the two binding domains may have the same sequence or bind the same epitope). A multivalent antibody comprising three or more binding domains may target one, two, three or more tumor associated antigens, permitting a specific targeting of deleterious cells over healthy cells. For example, one binding domain or two binding domains of the multivalent antibody may bind an antigen on an aberrant (tumor) cell, whereas a second or third binding domain of the multivalent antibody may bind an antigen on an immune effector cell that can cause directed killing of the tumor cell expressing the one or more tumor associated antigens. Alternatively, two binding domains of the multivalent antibody may bind specifically to two different epitopes on an identical antigen or different antigens expressed on tumor cells while the affinities of these arms are attenuated to mitigate binding to cells expressing only one antigen or where only one binding domain of the multivalent antibody is engaged. Or three binding domains of the multivalent antibody of the invention may bind to three different antigens or to identical antigens, but at different epitopes of immune effector cells.

Similarly, a multivalent antibody comprising three or more binding domains may bind a functional target such as a ligand or enzyme, triggering a biological response or blocking the function of the target, resulting in inhibitory or agonistic cellular activity. At least one binding domain of a multivalent antibody of the invention is connected via a linker to a binding domain of the base antibody portion. Where the binding domain of the base antibody portion is a Fab domain, this may take the form, for example, of VH-CH1-linker-VH-CH1, wherein the linker connects the heavy chain of the base antibody portion to the at least one additional binding domain, preferably a Fab domain.

Alternatively, this may take the form, for example, of VL-CL-linker-VL-CL, wherein the linker connects the light chain of the base antibody portion to the at least one additional binding domain, preferably a Fab domain.

An additional binding domain, such as a Fab domain, may be connected to each of the binding domains of the base antibody portion, each via a separate linker. The two or more linkers connecting the additional binding domains to the base antibody portion or additional binding domains may be the same or different. Further, the linkers may allow pairing of the cognate chains of the binding domain.

If an antibody of the invention comprises more than one linker, those linkers may be the same or different or a combination thereof. An example of the latter situation is where a multispecific antibody comprises three linkers, two of which are the same and a third which is different (from the other two).

Further a binding domain connected via a linker to a binding domain of a base antibody portion, may itself be attached to a binding domain connected via a linker described herein, wherein the base antibody portion may be extended in a modular fashion by connecting through

a linker to an additional binding domain, and connecting that binding domain to a second additional binding domain through a linker and so on.

In this way an antibody of the invention may be capable of binding three or more epitopes. Thus, a multispecific antibody of the invention may be capable of specifically binding 5 to three or more epitopes.

An antibody of the invention may be capable of binding two, three or more antigens. A multispecific antibody of the invention may thus be capable of specifically binding to two, three or more antigens.

An antibody of the invention may comprise two or more binding domains, such as two 10 or more Fab domains, which are capable of binding to different epitopes on one antigen

Accordingly, an antibody of the invention comprises at least three binding domains, such as two or more Fab domains which are different.

Another aspect of the invention comprises a multivalent antibody comprising at least three Fab domains and therefore is capable of binding to three epitopes which are typically all 15 different from each other.

An antibody of the invention may be multivalent. An antibody of the invention may also be multispecific. Multivalent indicates that the antibody has at least three binding domains and therefore has at least three antigen-binding sites. Multispecific indicates that the antibody is capable of binding at least two different epitopes, for example two different antigens or two 20 epitopes on the same antigen. Trispecific indicates that the antibody is capable of binding three different epitopes. Quadspecific indicates that the antibody is capable of binding four different epitopes and so on.

An antibody of the invention may bind target epitopes which are located on the same molecule. This may allow for more efficient counteraction of the (biological) function of said 25 target molecule as compared to a situation wherein only one epitope is targeted. For example, an antibody of the invention may simultaneously bind to 2 or 3 or more epitopes present on an antigen cell, e.g., growth factor receptors or soluble molecules critical for tumors cells to proliferate, thereby effectively blocking several independent signaling pathways leading to uncontrolled proliferation.

30 Any combination of at least two antibodies of the invention may simultaneously bind to 2, 3, 4 or more epitopes present on a target molecule, such as a growth factor receptor or soluble molecule.

The target moiety may be a soluble molecule or may be a membrane-bound moiety or may be a moiety present on a cell-surface that internalizes upon binding.

35 The target epitopes may be located on different moieties, for example on two (i.e. two or more target epitopes on a first moiety and one or more target epitopes on a second moiety) or three different moieties (i.e. at least one target epitope on each of three moieties). In this case, each of the different target moieties may either be a soluble moiety or a membrane-bound moiety or a moiety present on a cell-surface that internalizes upon binding. In one embodiment,

the different target moieties are soluble moieties. Alternatively, at least one target moiety is a soluble moiety whereas and at least one target moiety is a membrane bound moiety. In yet another alternative, all target moieties are membrane bound moieties. In one embodiment, the different target moieties are expressed on the same cell, whereas in other embodiments the different target moieties are expressed on different cells.

As a non-limiting example, any antibody of the invention or any combination of an antibody of the invention and an additional antibody may be suitable for simultaneously blocking multiple membrane-bound receptors, neutralizing multiple soluble molecules such as cytokines or growth factors for tumor cells or for neutralizing different viral serotypes or viral strains.

10 In an antibody of the invention, at least one target epitope may be located on a tumor cell. Alternatively, or additionally, at least a target epitope may be located on the surface of an effector cell. This is for instance suitable for recruitment of T cells or NK cells for tumor cell killing. For instance, an antibody of the invention may be capable of recruiting immune effector cells, preferably human immune effector cells, by specifically binding to a target molecule

15 located on immune effector cells. In a further embodiment, said immune effector cell is activated upon binding of the antibody of the invention to the target molecule. Recruitment of effector mechanisms may for instance encompass the redirection of immune modulated cytotoxicity by administering an Ig-like molecule produced by a method according to the invention that is capable of binding to a cytotoxic trigger molecule such as the T cell receptor or an Fc gamma

20 receptor, thereby activating downstream immune effector pathways or immune effector cells.

#### Immune cell engagers

25 A multivalent multimer, such as an antibody of the invention may be an immune effector cell engager antibody. That is to say, a multivalent antibody of the invention may be one which comprises at least one binding domain which binds specifically to an antigen on an immune effector cell, such as a T cell, and also comprises at least one binding domain which binds specifically to an antigen on an aberrant cell, such as a cancer or tumor cell.

30 A multivalent multimer of the invention, such as a trispecific antibody, may be one having three binding domains bringing three cells together in an engager complex, including a tumor cell, and two immune effector cells.

A multivalent multimer of the invention, such as a trispecific antibody, may further be one having three binding domains targeting two cells and a soluble molecule.

35 For the embodiments set out here, the Fc may be a wild-type Fc, may be enhanced for ADCC or binding of Cq1 based on means known to persons of skill in the art, or may be abrogated for such activity based on means known to persons of skill in the art.

The components of such immune cell engaging multivalent antibodies can be arranged with respect to each other in a variety of configurations. Exemplary configurations are depicted in Figures. 1a-1u. In particular embodiments, the present invention is directed to an immune

cell engaging multivalent antibody wherein a third binding domain is linked at the C-terminus of the Fab heavy chain to the N-terminus of a first or second binding domain of the base antibody.

In one embodiment, an immune cell engaging multivalent antibody comprises three binding domains, i.e. a base antibody portion and one additional binding domain, so that the 5 said multivalent antibody is trispecific,

One of the binding domains of the base antibody portion may be bind an antigen on an immune effector cell. Alternatively, the additional binding domain may bind an antigen of an immune effector cell. That is to say, the binding domain for an antigen on an immune effector cell may be at position 1, 2 or 3, wherein these positions correspond to the VH1, VH2 and VH3 10 indicated in Figure 1a. Alternatively, where a common heavy chain is used, the binding domain for an antigen on an immune effector cell may be at position 1, 2 or 3, wherein these positions correspond to the VL1, VL2 and VL3, for example, as indicated in Figure 1c.

In an immune effector cell engager antibody of the invention, at least one of the binding domains may specifically bind to an antigen on an aberrant cell. Typically, at least two binding 15 domains bind to an antigen on an aberrant cell, typically at least two binding domains bind to at least two different antigens or epitopes on a an antigen on an aberrant cell. In an immune effector cell engager antibody of the invention, two or more binding domains may bind the same target, including antigen and epitope, and another binding domain engaging an immune effector cell.

20 In a preferred embodiment of the invention, a multivalent antibody of the invention specifically binds to an antigen on an immune effector cell and also binds specifically to two different antigens on an aberrant cell, such as a tumor cell.

In one embodiment of the invention, the T cell engaging multivalent antibody is capable of simultaneous binding to a target cell antigen, particularly a tumor cell antigen, and a surface 25 antigen of a human T cell. In one embodiment, the T cell engaging multivalent antibody is capable of simultaneously binding to a target cell antigen, particularly a tumor cell antigen, and human CD3. In one embodiment, the T cell engaging multivalent antibody is capable of crosslinking a T cell and a target cell by simultaneous binding to a target cell antigen and CD3. In another embodiment, the simultaneous binding results in lysis of the target cell, particularly a 30 tumor cell. In one embodiment, the simultaneous binding results in activation of the T cell. In other embodiments, the simultaneous binding results in a cellular response of a T lymphocyte, particularly a cytotoxic T lymphocyte, selected from the group of: proliferation, differentiation, cytokine secretion, cytotoxic effector molecule release, cytotoxic activity, and expression of activation markers. In one embodiment, binding of the T cell engaging polypeptide to CD3 35 without simultaneous binding to the target cell antigen does not result in T cell activation, where, for example the remaining binding domains do not bind a tumor cell antigen.

In one embodiment, the T cell engaging multivalent antibody is capable of re-directing cytotoxic activity of a T cell to a target cell. In one embodiment, the re-direction is independent of MHC-mediated peptide antigen presentation by the target cell and and/or specificity of the T

cell. In one embodiment, the T cell is a cytotoxic T cell. In another embodiment, the T cell is a CD4<sup>+</sup> or a CD8<sup>+</sup> T cell. In another embodiment the T cell is a CD8<sup>+</sup> T cell.

The T cell engaging multivalent antibody of the invention comprises at least one antigen binding domain capable of binding to a surface antigen of a human T cell. In one embodiment, 5 the binding domain binds CD3 (also referred to herein as an "CD3 antigen binding domain").

The term "CD3" (cluster of differentiation 3) refers a protein complex, which is composed of a CD3 $\gamma$  chain (SwissProt P09693), a CD3 $\delta$  chain (SwissProt P04234), CD3 $\epsilon$  chains (SwissProt P07766), and a CD3 zeta chain homodimer (SwissProt P20963). CD3 $\epsilon$  is known under various aliases some of which are: "CD3e Molecule, Epsilon (CD3-TCR

10 Complex)"; "CD3e Antigen, Epsilon Polypeptide (TIT3 Complex)"; T-Cell Surface Antigen T3/Leu-4 Epsilon Chain; T3E; T-Cell Antigen Receptor Complex, Epsilon Subunit Of T3; CD3e Antigen; CD3-Epsilon 3; IMD18; TCRE. Ids for CD3E Gene are HGNC: 1674; Entrez Gene: 916; Ensembl: ENSG00000198851; OMIM: 186830 and UniProtKB: P07766. These chains associate with the T-cell receptor (TCR) and the  $\zeta$ -chain to form a TCR complex that can upon 15 mitogenic signaling generates an activation signal in T lymphocytes. CD3 is expressed on T cells and NK T cells. Where reference is made to CD3 herein, the reference is to human CD3, unless specifically stated otherwise.

In a particular embodiment, the T cell engaging polypeptide comprises not more than one binding domain capable of specific binding to CD3. In one embodiment the T cell engaging 20 polypeptide provides monovalent binding to CD3. In one embodiment, the T cell engaging polypeptide comprises one member of a supercluster of CD3 binding binding domains. A 'super-cluster' is used hereinto to refer to variable regions having amino acid changes that are tolerated, for example, with respect to heavy chain variable regions, including a VH or VL and/or CDR therein of the present invention without losing binding specificity to the particular antigen.

25 More specifically, a 'super-cluster' is a group of clones sharing the same VH V-gene usage and having at least 70% sequence identity in HCDR3 and the same HCDR3 length. The clones in a supercluster are expected to bind the same antigen potentially with different affinities and/or different location on the epitope.

The CD3 binding domain may range in affinity, epitope and other characteristics. 30 Specific variable domains that can bind an extracellular part of CD3 are variable domains that comprise the amino acid sequence of the VH of MF8057, MF8058, MF8078 and variable regions of this supercluster, MF8397 and variable regions of this supercluster, MF8508 and variable regions of this supercluster, and MF9249 and MF9267 and variable regions of this supercluster.

35 The CD3 antigen binding domain comprises at least one heavy chain complementarity determining region (CDR) selected from the group consisting VH of MF8057, MF8058, MF8078 and variable regions of this supercluster, MF8397 and variable regions of this supercluster, MF8508 and variable regions of this supercluster, and MF9249 and MF9267 and variable regions of this supercluster (SEQ ID NO: 97, SEQ ID NO:106, SEQ ID NO:115, SEQ ID

NO:124, SEQ ID NO:133, SEQ ID NO:142, SEQ ID NO:98, SEQ ID NO:107, SEQ ID NO:116, SEQ ID NO:125, SEQ ID NO:134 SEQ ID NO:143 SEQ ID NO:152SEQ ID NO: 99, SEQ ID NO:108, SEQ ID NO:117, SEQ ID NO:126, SEQ ID NO:135, SEQ ID NO:144 and/or SEQ ID NO:153) and at least one light chain CDR selected from the group of SEQ ID NO: 254, SEQ ID 5 NO: 255, SEQ ID NO: 256.

In one embodiment the CD3 antigen binding domain comprises the heavy chain CDR1 of SEQ ID NO: 97, SEQ ID NO:106, SEQ ID NO:115, SEQ ID NO:124, SEQ ID NO:133, or SEQ ID NO:142, the heavy chain CDR2 of SEQ ID NO:98, SEQ ID NO:107, SEQ ID NO:116, SEQ ID NO:125, SEQ ID NO:134 SEQ ID NO:143 or SEQ ID NO:152, the heavy chain CDR3 of SEQ ID 10 NO: 99, SEQ ID NO:108, SEQ ID NO:117, SEQ ID NO:126, SEQ ID NO:135, SEQ ID NO:144 or SEQ ID NO:153, the light chain CDR1 of SEQ ID NO: 254, the light chain CDR2 of SEQ ID NO: 255, and the light chain CDR3 of SEQ ID NO: 256.

In one embodiment the CD3 antigen binding domain comprises a heavy chain variable region sequence that is at least about 95%, 96%, 97%, 98%, 99% or 100% identical to an 15 amino acid sequence selected from the group of: SEQ ID NO: 100, SEQ ID NO:109, SEQ ID NO:118, SEQ ID NO:127, SEQ ID NO:135, SEQ ID NO:145 and SEQ ID NO:154, and a light chain variable region sequence that is at least about 95%, 96%, 97%, 98%, 99% or 100% identical to an amino acid sequence selected from the group of: SEQ ID NO: 37 and SEQ ID NO: 40.

20 In one embodiment the CD3 antigen binding domain comprises the heavy chain variable region of SEQ ID NO: 100, SEQ ID NO:109, SEQ ID NO:118, SEQ ID NO:127, SEQ ID NO:135, SEQ ID NO:145 or SEQ ID NO:154 and the light chain variable region comprising the amino acid sequence of SEQ ID NO: 37 or SEQ ID NO:40.

The positions of the binding domains in the multivalent antibody can be defined. When 25 the binding or variable domains of the base antibody are termed binding domains 1 and 2, the additional one or more binding domains can be referred to as binding domains 3, 4 etc. A binding domain is also referred to as BD. BD3 may be linked to BD1 or BD2. When BD3 is linked to one of BD1 or 2, BD4, when present, is linked to the other of BD1 and 2.

30 The T cell engaging multivalent antibody of the invention comprises at least two antigen binding domains capable of binding to a target cell antigen (also referred to herein as an "target cell antigen binding domain" or "second" or "third" antigen binding domain). In certain embodiments, the T cell engaging multivalent antibody comprises two antigen binding domains capable of binding to a target cell antigen. In one embodiment, each of these antigen binding 35 domains specifically binds to the same antigenic determinant. In another embodiment, the target cell antigen binding domains are identical. In one embodiment, the T cell engager polypeptide comprises not more than two target cell antigen binding domains capable of binding to a target cell antigen.

In one embodiment the multivalent antibody comprises an BD1 which is a CD3 binding domain, an BD2 which is a binding domain that binds a first target cell antigen, further referred TA1 and an BD3 which is a binding domain that binds a second target cell antigen, further referred TA2. In this embodiment BD3 can be linked to BD1 or BD2. In one embodiment TA2 5 binding BD3 is linked to CD3 binding BD1. In another embodiment the TA2 binding BD3 is linked to TA1 binding BD2.

In one embodiment the multivalent antibody comprises a TA1 binding BD1, a TA2 binding BD2, and a CD3 binding BD3. In this embodiment BD3 can be linked to BD1 or BD2. In one embodiment the CD3 binding BD3 is linked to TA1 binding BD1. In another embodiment 10 CD3 binding BD3 is linked to TA2 binding BD2.

In one embodiment the invention provides a multivalent antibody wherein the base antibody comprises binding domains 1 and 2 (BD1 and 2) and wherein the additional binding domain 3 (BD3) is linked to binding domain 1 (BD1) and wherein an optional additional binding 15 domain 4 (BD4) is linked to binding domain 2 (BD2). In one embodiment binding domain 1 is a CD3 binding domain and binding domains 2 and 3 bind to different target cell antigens. In another embodiment binding domain 2 is a CD3 binding domain and binding domains 1 and 3 bind to different target cell antigens. In a further embodiment binding domain 3 is a CD3 binding domain and binding domains 1 and 2 bind to different target cell antigens.

20 In a preferred embodiment the multivalent antibody comprises a binding domain 4 that binds yet a further different target cell antigen.

The invention further provides a multivalent antibody as described herein wherein the base antibody comprises binding domains 1 and 2 and wherein the additional binding domain 3 is 25 linked to binding domain 1 and wherein an optional additional binding domain 4 is linked to binding domain 2. In one embodiment binding domain 1 is a CD3 binding domain and binding domains 2 and 3 bind to different target cell antigens. In another embodiment binding domain 2 is a CD3 binding domain and binding domains 1 and 3 bind to different target cell antigens. In a further embodiment binding domain 3 is a CD3 binding domain and binding domains 1 and 2 30 bind to different target cell antigens.

When comprising a binding domain 4 the domain preferably binds yet a further different target cell antigen.

35 In one embodiment a first of said target cell antigen binding domain binds PD-L1, EGFR, CD137, CLEC12A, fibrinogen, or thyroglobulin. In one embodiment a first and a second of said target cell antigen binding domains bind antigens selected from PD-L1, EGFR, CD137, CLEC12A fibrinogen, and thyroglobulin. The first and second target cell binding domain preferably bind different antigens.

A CD3 binding domain preferably comprises a heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid sequence of the CDR1, CDR2 and CDR3 of MF8057, or of MF8058, or of MF8078, or of MF8397, or of MF8508, or of MF9249 or of

5 MF9267. The CD3 binding domain preferably comprises a heavy chain variable region with the amino acid sequence of the VH of MF8057, of MF8058, of MF8078, of MF8397, of MF8508, of MF9249 or of MF9267 with 0-10, preferably 0-5 amino acid insertions, deletions, substitutions, additions or a combination thereof at one or more positions other than the CDRs.

10 A target cell antigen binding domain can be a PD-L1 binding domain. If present the PD-L1 binding domain preferably comprises a heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid sequence of the CDR1, CDR2 and CDR3 of MF5377, or of MF5444, or of MF5380. The PD-L1 binding domain preferably comprises a heavy chain variable region with the amino acid sequence of the VH of MF5377, of MF5444, or of MF5380 with 0-10, 15 preferably 0-5 amino acid insertions, deletions, substitutions, additions or a combination thereof at one or more positions other than the CDRs.

A target cell antigen binding domain can be an EGFR binding domain. If present the EGFR binding domain preferably comprises a heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid sequence of the CDR1, CDR2 and CDR3 of MF8233, or of

20 MF9891, or of MF9886, or of MF9873, or of MF9988. The EGFR binding domain preferably comprises a heavy chain variable region with the amino acid sequence of the VH of MF8233, of MF9891, of MF9886, of MF9873, or of MF9988 with 0-10, preferably 0-5 amino acid insertions, deletions, substitutions, additions or a combination thereof at one or more positions other than

25 the CDRs.

A target cell antigen binding domain can be a CLEC12A binding domain. If present the CLEC12A binding domain preferably comprises a heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid sequence of the CDR1, CDR2 and CDR3 of

30 MF4327. The CLEC12A binding domain preferably comprises a heavy chain variable region with the amino acid sequence of the VH of MF4327 with 0-10, preferably 0-5 amino acid insertions, deletions, substitutions, additions or a combination thereof at one or more positions other than the CDRs.

In one embodiment the multivalent antibody comprises a CD3 binding domain, an EGFR binding domain and a PD-L1 binding domain.

The binding domains with the indicated heavy chain variable regions comprise a light chain variable region. The light chain variable region preferably comprises a CDR1, CDR2, and CDR3 region comprising the amino acid sequence CDR1 - QSISSY, CDR2 – AAS, CDR3 –

QQSYSTP, i.e. the CDRs of IGKV1-39 (according to IMGT). The amino acid variations, insertions, deletions, substitutions, additions or combination thereof are preferably not in the CDR3 region of the light chain variable region, preferably not in the CDR1 or CDR2 region of the light chain variable region. In a preferred embodiment the light chain variable region does 5 not comprise a deletion, addition or insertion with respect to the sequence indicated. In this embodiment the light chain variable region can have 0-5 amino acid substitutions with respect to the indicated amino acid sequence. An amino acid substitution is preferably a conservative amino acid substitution. The CDR1, CDR2 and CDR3 of a light chain of an antibody of the invention preferably comprises respectively the amino acid sequence CDR1 - QSISSY, CDR2 - 10 AAS, CDR3 - QQSYSTP, i.e. the CDRs of IGKV1-39 (according to IMGT).as described elsewhere herein. The light chains of the binding domains with the indicated heavy chain variable regions preferably all comprise the same light chain. Preferably a common light chain as defined elsewhere herein.

15 Amino acid insertions, deletions, substitutions, additions or combination thereof are preferably not in the CDR3 region of the heavy chain variable region, preferably not in the CDR1 and/or CDR2 region of the heavy chain variable region. In a preferred embodiment the heavy chain variable region does not comprise a deletion, addition or insertion with respect to the sequence indicated. In one embodiment the heavy chain variable region can have 0-10, 20 preferably 0-5 amino acid substitutions with respect to the indicated amino acid sequence. In a preferred embodiment the heavy chain variable region comprises 0-9, 0-8, 0-7, 0-6, 0-5, 0-4, preferably 0-3, preferably 0-2, preferably 0-1 and preferably 0 amino acid insertions, deletions, substitutions, additions with respect to the indicated amino acid sequence, or a combination thereof at positions other than the CDRs. A combination of an insertion, addition, deletion or 25 substitution is a combination as claimed if aligned sequences do not differ at more than 10, preferably no more than 5 positions. A gap in one of the aligned sequences counts for as many amino acids as skipped in the other sequence. An amino acid substitution, if any, is preferably a conservative amino acid substitution.

30 In one embodiment, the target cell antigen binding domain is a Fab molecule. In one embodiment, the target cell antigen binding domain is a Fab molecule that binds to a specific antigenic determinant and is able to direct the T cell engaging multivalent antibody to a target site, for example to a specific type of tumor cell that bears the antigenic determinant. In certain embodiments the target cell antigen binding specifically binds Programmed Cell Death 35 1 protein (PD-L1), preferably human PD-L1 (SEQ ID NO: 257).

PD-L1 is a type 1 transmembrane protein that plays a role in suppressing an immune response during particular events such as pregnancy, tissue allografts, autoimmune disease and other disease states such as hepatitis. The binding of PDL1 to PD-1 or B7.1 (CD80) transmits an inhibitory signal which reduces the proliferation of the PD-1 expressing T cells. PD-

1 is thought to be able to control the accumulation of foreign antigen specific T cells through apoptosis. PD-L1 is expressed by a variety of cancer cells and the expression thereof is thought to be at least in part responsible for a dampening of an immune response against the cancer cell. PD-L1 is a member of the B7-family of protein and is known under a variety of other names 5 such as CD274 Molecule; CD274 Antigen; B7 Homolog 1; PDCD1 Ligand 1; PDCD1LG1; PDCD1L1; B7H1; PDL1; Programmed Cell Death 1 Ligand 1; Programmed Death Ligand 1; B7-H1; and B7-H. External Ids for CD274 are HGNC: 17635; Entrez Gene: 29126; Ensembl: ENSG00000120217; OMIM: 605402; UniProtKB: Q9NZQ7.

The PD-L1 binding domain may range in affinity, epitope and other characteristics. 10 Specific variable domains that can bind an extracellular part of PD-L1 are variable domains that comprise the amino acid sequence of the VH of MF5377, MF5444 or MF5380 .

The PD-L1 antigen binding domain comprises at least one heavy chain CDR selected from the group consisting of VH of SEQ ID NO: 160, SEQ ID NO:169, SEQ ID NO:178, SEQ ID NO: 161, SEQ ID NO:170, SEQ ID NO:179 SEQ ID NO: 162, SEQ ID NO:171 and SEQ ID 15 NO:180 and at least one light chain CDR selected from the group of SEQ ID NO: 254, SEQ ID NO: 255, and SEQ ID NO: 256.

In one embodiment the PD-L1 antigen binding domain comprises the heavy chain CDR1 of SEQ ID NO: 160, SEQ ID NO:169 or SEQ ID NO:178, the heavy chain CDR2 of SEQ ID NO: 161, SEQ ID NO:170 or SEQ ID NO:179, the heavy chain CDR3 of SEQ ID NO: 162, 20 SEQ ID NO:171 or SEQ ID NO:180, the light chain CDR1 of SEQ ID NO: 254, the light chain CDR2 of SEQ ID NO: 255, and the light chain CDR3 of SEQ ID NO: 256.

In one embodiment the PD-L1 antigen binding domain comprises a heavy chain variable region sequence that is at least about 95%, 96%, 97%, 98%, 99% or 100% identical to an amino acid sequence selected from the group of: SEQ ID NO: 163, SEQ ID NO:172 and 25 SEQ ID NO:181, and a light chain variable region sequence that is at least about 95%, 96%, 97%, 98%, 99% or 100% identical to an amino acid sequence selected from the group of: SEQ ID NO: 37 and SEQ ID NO: 40.

In one embodiment the PD-L1 antigen binding domain comprises the heavy chain variable region of SEQ ID NO: 163, SEQ ID NO:172 or SEQ ID NO:181 and the light chain variable 30 region comprising the amino acid sequence of SEQ ID NO: 37 or SEQ ID NO:40.

In certain embodiments, the PD-L1 antigen binding domain comprises the heavy and light chain variable regions of the PD-L1 antibodies comprising the amino acid sequences disclosed for MPDL3280A, RG7446, see US 2010/0203056 A1; MEDI-4736, see WO 2011/066389; MSB-0010718C, see WO 2013/079174; STI-1014 see WO2013/181634; CX-072, 35 see WO2016/149201; KN035, see Zhang et al., Cell Discov. 7:3 (March 2017); LY3300054, see, e.g., WO 2017/034916; and CK-301, see Gorelik et al., AACR:Abstract 4606 (Apr 2016)), and 12A4 or MDX-1105, see, e.g., WO 2013/173223.

In certain embodiments, the PD-L1 antigen binding domain binds the same epitope as the heavy and light chain variable regions of the PD-L1 antibodies MPDL3280A, RG7446, see

US 2010/0203056 A1; MEDI-4736, see WO 2011/066389; MSB-0010718C, see WO 2013/079174; STI-1014 see WO2013/181634; CX-072, see WO2016/149201; KN035, see Zhang et al., *Cell Discov.* 7:3 (March 2017); LY3300054, see, e.g., WO 2017/034916; and CK-301, see Gorelik et al., AACR:Abstract 4606 (Apr 2016)), and 12A4 or MDX-1105, see, e.g.,

5 WO 2013/173223.

In certain embodiments, the PD-L1 antigen binding domain competes for binding to PD-L1 with the heavy and light chain variable regions of the PD-L1 antibodies MPDL3280A, RG7446, see US 2010/0203056 A1; MEDI-4736, see WO 2011/066389; MSB-0010718C, see WO 2013/079174; STI-1014 see WO2013/181634; CX-072, see WO2016/149201; KN035, see

10 Zhang et al., *Cell Discov.* 7:3 (March 2017); LY3300054, see, e.g., WO 2017/034916; and CK-301, see Gorelik et al., AACR:Abstract 4606 (Apr 2016)), and 12A4 or MDX-1105, see, e.g., WO 2013/173223.

In certain embodiments the target cell antigen binding specifically binds human epidermal growth factor receptor (EGFR) (SEQ ID NO: 258). 'ErbB1' or 'EGFR' is a member of

15 a family of four receptor tyrosine kinases (RTKs), named Her- or cErbB-1, -2, -3 and -4. The EGFR has an extracellular domain (ECD) that is composed of four sub-domains, two of which are involved in ligand binding and one of which is involved in homo-dimerisation and heterodimerisation. The reference numbers used in this section refer to the numbering of the references in the list headed "References cited in the specification". EGFR integrates

20 extracellular signals from a variety of ligands to yield diverse intracellular responses. The major signal transduction pathway activated by EGFR is composed of the Ras-mitogen-activated protein kinase (MAPK) mitogenic signalling cascade. Activation of this pathway is initiated by the recruitment of Grb2 to tyrosine phosphorylated EGFR. This leads to activation of Ras through the Grb2-bound Ras-guanine nucleotide exchange factor Son of Sevenless (SOS). In

25 addition, the PI3-kinase-Akt signal transduction pathway is also activated by EGFR, although this activation is much stronger in case there is co-expression of Her3. The EGFR is implicated in several human epithelial malignancies, notably cancers of the breast, bladder, non-small cell lung cancer lung, colon, ovarian head and neck and brain. Activating mutations in the gene have been found, as well as over-expression of the receptor and of its ligands, giving rise to

30 autocrine activation loops. This RTK has therefore been extensively used as target for cancer therapy. Both small-molecule inhibitors targeting the RTK and monoclonal antibodies (mAbs) directed to the extracellular ligand-binding domains have been developed and have shown hitherto several clinical successes, albeit mostly for a select group of patients. A database accession number for the human EGFR protein and the gene encoding it is (GenBank

35 NM\_005228.3). The accession number is primarily given to provide a further method of identification of EGFR protein as a target, the actual sequence of the EGFR protein bound by an antibody may vary, for instance because of a mutation in the encoding gene such as those occurring in some cancers or the like. Where reference herein is made to EGFR, the reference refers to human EGFR unless otherwise stated. The antigen-binding site that binds EGFR,

binds EGFR and a variety of variants thereof such as those expressed on some EGFR positive tumors.

The EGFR binding domain may range in affinity, epitope and other characteristics. Specific variable domains that can bind an extracellular part of EGFR are variable domains that 5 comprise the amino acid sequence of the VH of MF8233, MF9891, MF9886, MF9873, MF9988.

The EGFR antigen binding domain comprises at least one heavy chain CDR selected from the group consisting of SEQ ID NO: 187, SEQ ID NO:196, SEQ ID NO:205, SEQ ID NO:214, SEQ ID NO:223, SEQ ID NO: 188, SEQ ID NO:197, SEQ ID NO:206, SEQ ID NO:215, 10 SEQ ID NO:224 SEQ ID NO: 189, SEQ ID NO:198, SEQ ID NO:207, SEQ ID NO:216 and SEQ ID NO:225 and at least one light chain CDR selected from the group of SEQ ID NO: 254, SEQ ID NO: 255, SEQ ID NO: 256.

In one embodiment the EGFR antigen binding domain comprises the heavy chain CDR1 of SEQ ID NO: 187, SEQ ID NO:196, SEQ ID NO:205, SEQ ID NO:214, or SEQ ID 15 NO:223, the heavy chain CDR2 of SEQ ID NO: 188, SEQ ID NO:197, SEQ ID NO:206, SEQ ID NO:215, or SEQ ID NO:224, the heavy chain CDR3 of SEQ ID NO: 189, SEQ ID NO:198, SEQ ID NO:207, SEQ ID NO:216 or SEQ ID NO:225, the light chain CDR1 of SEQ ID NO: 254, the light chain CDR2 of SEQ ID NO: 255, and the light chain CDR3 of SEQ ID NO: 256.

In one embodiment the EGFR antigen binding domain comprises a heavy chain 20 variable region sequence that is at least about 95%, 96%, 97%, 98%, 99% or 100% identical to an amino acid sequence selected from the group of: SEQ ID NO: 190, SEQ ID NO:199, SEQ ID NO:208, SEQ ID NO:217 and SEQ ID NO:226, and a light chain variable region sequence that is at least about 95%, 96%, 97%, 98%, 99% or 100% identical to an amino acid sequence selected from the group of: SEQ ID NO: 37 and SEQ ID NO: 40.

25 In one embodiment the EGFR antigen binding domain comprises the heavy chain variable region of SEQ ID NO: 190, SEQ ID NO:199, SEQ ID NO:208, SEQ ID NO:217 or SEQ ID NO:226 and the light chain variable region comprising the amino acid sequence of SEQ ID NO: 37 or SEQ ID NO:40.

In certain embodiments, the EGFR antigen binding domain comprises the heavy and 30 light chain variable regions of the EGFR antibodies cetuximab (C225, Erbitux®, Lilly) or panitumumab (Vectibix, Amgen).

In certain embodiments, the EGFR antigen binding domain binds the same epitope as the heavy and light chain variable regions of the EGFR antibodies cetuximab (C225, Erbitux®, Lilly) or panitumumab (Vectibix, Amgen).

35 In certain embodiments, the EGFR antigen binding domain competes for binding to EGFR with the heavy and light chain variable regions of the EGFR antibodies cetuximab (C225, Erbitux®, Lilly) or panitumumab (Vectibix, Amgen).

Common variable region

The multivalent antibody of the invention preferably uses a common chain at each of the three or more binding domains. As described, the base antibody portion of the multivalent antibody invention preferably has a first heavy chain variable region/light chain variable region (VH/VL) combination that binds one antigen and a second VH/VL combination that binds a second antigen. Each additional binding domain connected to the base antibody portion may also comprise an additional VH/VL combination that binds a further epitope on an antigen.

A base antibody portion of the invention preferably comprises two heavy chains (one or both comprising one or more additional CH1 and VH domain) and a light chain which pairs with each CH1 and VH domain. Preferably the two heavy chains have compatible heterodimerization domains, and preferably the light chain is a common light chain. Alternatively, the base antibody portion of the multivalent antibody of the invention comprises two light chains (one or both comprising one or more additional CL and VL domain) and a heavy chain variable region which pairs with each CL and VL domain, and the heavy chain variable region comprises a common heavy chain variable region.

Where the embodiment of the invention includes a multivalent antibody comprising a common light chain, where said light chain is expressed within a host cells that includes DNA encoding two or more heavy chain variable regions, said light chain is capable of pairing with each available heavy chains (or CH1-VH1 regions), thereby forming at least three functional antigen binding domains.

A functional antigen binding domain is capable of specifically binding to an epitope on an antigen. Preferably, a common light chain used in a multivalent antibody of the invention is capable of pairing with all heavy chains (or CH1-VH1 regions) produced with a method according to the invention, thereby forming functional antigen binding domains, so that mispairing of unmatched heavy and light chains is avoided or produced at a significantly lower ratio than the multivalent antibody.

It is a preferred aspect of the present invention that a multivalent antibody of the invention has a common light chain (variable region) that can combine with an array of heavy chain variable regions to form an antibody with functional antigen binding domains (WO2004/009618, WO2009/157771).

A common light chain (variable region) for use in the multivalent antibody of the invention is preferably a human light chain (variable region). A common light chain (variable region) preferably has a germline sequence. A preferred germline sequence is a light chain variable region that is frequently used in the human repertoire and has good thermodynamic stability, yield and solubility. A preferred germline light chain is O12. A common light chain is preferably the rearranged germline human kappa light chain IgV $\kappa$ 1-39\*01/IGJ $\kappa$ 1\*01 (Figure 11A; SEQ ID NO: 35). The common light chain variable region is preferably the variable region of the rearranged germline human kappa light chain IgV $\kappa$ 1-39\*01/IGJ $\kappa$ 1\*01 (Figure 11A; SEQ ID NO:

35). A common light chain preferably comprises a light chain variable region as depicted in Figure 11B or 8D (SEQ ID NOs: 37 or 40 respectively) with 0-5 amino acid insertions, deletions, substitutions, additions or a combination thereof. The common light preferably further comprises a light chain constant region, preferably a kappa light chain constant region. A nucleic acid that 5 encodes the common light chain can be codon optimized for the cell system used to express the common light chain protein. The encoding nucleic acid can deviate from a germ-line nucleic acid sequence.

The common light chain (variable region) for use in the multivalent antibodies of the invention can be a lambda light chain and this is therefore also provided in the context of the 10 invention, however a kappa light chain is preferred. The common light chain of the invention may comprise a constant region of a kappa or a lambda light chain. It is preferably a constant region of a kappa light chain, preferably wherein said common light chain is a germline light chain, preferably a rearranged germline human kappa light chain comprising the IgV<sub>K</sub>l-39 gene segment, for example the rearranged germline human kappa light chain IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>l\*01 15 (Figure 11). The terms rearranged germline human kappa light chain IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>l\*01, IGKV1-39/IGJ<sub>K</sub>l, huV<sub>K</sub>l-39 light chain or in short huV<sub>K</sub>l-39, or simply 1-39 are used 20 interchangeably throughout the application. Those of skill in the art will recognize that "common" also refers to functional equivalents of the light chain of which the amino acid sequence is not identical. Many variants of said light chain exist wherein mutations (deletions, substitutions, additions) are present that do not materially influence the formation of functional binding regions.

IgV<sub>K</sub>l-39 is short for Immunoglobulin Variable Kappa 1-39 Gene. The gene is also known as Immunoglobulin Kappa Variable 1-39; IGKV139; IGKV1-39; O12a or O12. External 25 IDs for the gene are HGNC: 5740; Entrez Gene: 28930; Ensembl: ENSG00000242371. A preferred amino acid sequence for IgV<sub>K</sub>l-39 is given in Figure 11. This lists the sequence of the V-region. The V-region can be combined with one of five J-regions. Figure 11 describes two preferred sequences for IgV<sub>K</sub>l-39 in combination with a J-region. The joined sequences are indicated as IGKV1-39/jk1 and IGKV1-39/jk5; alternative names are IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>l\*01 or IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>5\*01 (nomenclature according to the IMGT database worldwide web at 30 imgt.org).

A common light chain variable region is preferably linked to a kappa light chain constant region. In a preferred embodiment the light chain variable region used in the multivalent antibody of the invention comprises the kappa light chain IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>l\*01 or IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>5\*01. In a preferred embodiment the common light chain in the multivalent antibody 35 is IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>l\*01.

A cell that produces a common light chain can produce for instance rearranged germline human kappa light chain IgV<sub>K</sub>l-39\*01/IGJ<sub>K</sub>l\*01 and a light chain comprising the variable region of the mentioned light chain fused to a lambda constant region. Where herein

reference is made to a germ-line sequence it is preferred that the variable region is a germ-line sequence.

A preferred common light chain for use in a multivalent antibody of the invention is one comprising the sequence set out in SEQ ID NO: 29.

5 The common chain for use in the multivalent antibodies of the invention can also be a heavy chain and this is therefore also provided in the context of the invention. Common heavy chains have been used in the art to make bispecific antibodies, and can be used here in making a multivalent antibody comprising three or more binding domain, two or more of said binding domains comprise a common heavy chain known in the art. For example, the use of antibody 10 libraries in which the heavy chain variable domain is the same for all the library members and thus the diversity is based on the light chain variable domain. Such libraries are described, for example, PCT/US2010/035619, and PCT/US2010/057780, each of which is hereby incorporated by reference in its entirety. These and other techniques to generating binding 15 domains having common heavy chains can be generated by the skilled artisan, and can be employed in the present invention to produce multivalent antibodies having novel formats disclosed herein.

#### Production of a multivalent antibody

20 A multivalent antibody of the invention may be produced by co-transfection of individual cells with one or more genetic constructs which together encode the three or more proteins that form a multimer comprising the multivalent antibody such as those described above, including in Figures 1a-u. For example, a host cell may be co-transfected with nucleic acid encoding three or more heavy chain variable regions and a common light chain variable region to produce a multivalent antibody. Alternatively, a multivalent antibody of the invention may be produced by 25 co-transfection of individual cells with one or more genetic constructs which together encode the three or more light chain variable regions and a common heavy chain.

30 Several methods have been published to favor the production antibodies which are heterodimers. In the present invention it is preferred that the cell favors the production of the heterodimers over the production of the respective homodimers. This is typically achieved by modifying the constant region of the heavy chains such that they favor heterodimerization (i.e. dimerization with one heavy chain combining with the second heavy chain) over homodimerization. In a preferred embodiment the antibody of the invention comprises two different immunoglobulin heavy chains with compatible heterodimerization domains.

35 The compatible heterodimerization domains are preferably compatible immunoglobulin heavy chain CH3 heterodimerization domains. When wildtype CH3 domains are used, co-expression of two different heavy chains (A and B) and a common light chain will result in three different antibody species, AA, AB and BB. AA and BB are designations for the two homodimer antibodies and AB is a designation for the heterodimer antibody. To increase the percentage of the desired heterodimer product (AB) CH3 engineering can be employed, or in other words, one

can use heavy chains with compatible hetero-dimerization domains, as defined hereunder. The art describes various ways in which such hetero-dimerization of heavy chains can be achieved.

The term 'compatible hetero-dimerization domains' as used herein refers to protein domains that are engineered such that engineered domain A' will preferentially form 5 heterodimers with engineered domain B' and vice versa, homo-dimerization between A'-A' and B'-B' is diminished.

In US13/866,747 (now issued as US 9,248,181), US14/081,848 (now issued as US 9,358,286), WO2013/157953 and WO2013/157954, methods and means are disclosed for 10 producing multivalent antibodies using compatible heterodimerization domains. These means and methods can also be favorably employed in the present invention. Specifically, an antibody 15 of the invention preferably comprises mutations to produce essentially only bispecific full length IgG molecules. Preferred mutations are the amino acid substitutions L351K and T366K (EU numbering) in the first CH3 domain or at positions corresponding thereto (the 'KK-variant' heavy chain) and the amino acid substitutions L351D and L368E in the second domain or at positions 20 corresponding thereto (the 'DE-variant' heavy chain), or vice versa. It was previously demonstrated in our US 9,248,181 and US 9,358,286 patents as well as the WO2013/157954 PCT application that the DE-variant and KK-variant preferentially pair to form heterodimers (so-called 'DEKK' bispecific molecules). Homodimerization of DE-variant heavy chains (DEDE homodimers) or KK-variant heavy chains (KKKK homodimers) hardly occurs due to repulsion between the charged residues in the CH3-CH3 interface between identical heavy chains.

In a preferred host cell of the present invention, capable of expressing proteins that multimerize to form a multivalent antibody, the host cell is transformed with a nucleic acid that encodes three proteins. In order from N-terminus to C-terminus, the encoded proteins include a 25 first protein comprising VH1-CH1-VH2-CH1-CH2-CH3, wherein a linker connects VH2 and CH1 on the first protein, a second encoded protein comprising VLc-CL, a third encoded protein comprising VH3-CH1-CH2-CH3, wherein the CH1 of the first and third encoded protein pairs with the CL of the second encoded protein, and the encoded CH3 region of the first and third proteins encode amino acid L351K and T366K (EU numbering) in the first CH3 protein or at 30 positions corresponding thereto and the amino acids L351D and L368E in the third protein or a corresponding positions thereto respectively, or vice versa. Alternatively, said first and third proteins comprise other compatible hetero-dimerization domains that cause the efficient pairing of the CH3 domains of each of these proteins.

Said nucleic acids encoding said three proteins may be on one or more vectors, to generate a multivalent antibody of the invention. Similarly, host cells can be generated 35 encoding more than three proteins for each of the multivalent antibodies described above, including those in Figures 1a-1u.

Said nucleic acids encoding said three proteins may further be stably integrated into the host cell's genome, preferably at chromosomal regions known for high expression and an absence or reduction of gene silencing.

According to the invention, there is thus provided a method for the preparation of a multivalent antibody, which method comprises:

5 providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into a multivalent antibody according to the invention; and

cultivating said host cell under conditions to provide for expression of the polypeptides and for their assembly into a multivalent antibody.

10

A host cell of the present invention may be capable of producing the multivalent antibody at a purity of at least about 50%, at least about 60%, least about 70%, at least about 80%, at least about 90%, at least about 95%, at least about 98% of the multivalent antibody of the invention on the basis of total expressed immunoglobulin.

15

A host cell of the invention may be capable of producing the multivalent antibody, wherein at least about 50%, at least about 60%, least about 70%, at least about 80%, at least about 90%, at least about 95%, at least about 98% of the multivalent antibody produced comprises a variable rearranged region paired with a cognate common chain for all binding sites.

20

A host cell of the invention may be capable of producing the multivalent antibody, wherein at least about 50%, at least about 60%, least about 70%, at least about 80%, at least about 90%, at least about 95%, at least about 98% of the common chain expressed is paired to the multivalent antibody and is not free, unassociated protein.

25

Suitable cells for antibody production are a hybridoma cell, a Chinese hamster ovary (CHO) cell, an NS0 cell or a PER-C6 cell. In a particularly preferred embodiment said cell is a CHO cell. Cells for the production of an antibody as disclosed herein are also referred to as host cells.

30

Various institutions and companies have developed cell lines for the large scale production of antibodies, for instance for clinical use. Non-limiting examples of such cell lines are CHO cells, NS0 cells or PER.C6 cells. At least some of these cells are also used for other purposes such as the production of proteins. Cell lines developed for industrial scale production of proteins and antibodies are herein further referred to as industrial cell lines. In a preferred embodiment the invention provides an industrial cell line that produces and an antibody of the invention.

35

The invention in one embodiment provides a cell (host cell) comprising an antibody according to the invention and/or a nucleic acid according to the invention. Said cell is preferably an animal cell, more preferably a mammal cell, more preferably a primate cell, most preferably a human cell. For the purposes of the invention a suitable cell, suitable host cell is

any cell capable of comprising and preferably of producing an antibody according to the invention and/or a nucleic acid according to the invention.

The invention further provides a cell comprising an antibody according to the invention. Preferably said cell (typically an in vitro, isolated or recombinant cell) produces said antibody. In 5 a preferred embodiment said cell is a hybridoma cell, a Chinese hamster ovary (CHO) cell, an NS0 cell or a PER.C6 cell. In a particularly preferred embodiment said cell is a CHO cell. Further provided is a cell culture comprising a cell according to the invention. Various 10 institutions and companies have developed cell lines for the large scale production of antibodies, for instance for clinical use. Non-limiting examples of such cell lines are CHO cells, NS0 cells or PER.C6 cells. These cells are also used for other purposes such as the production 15 of proteins. Cell lines developed for industrial scale production of proteins and antibodies are herein further referred to as industrial cell lines. Thus in a preferred embodiment the invention provides the use of a cell line developed for the large scale production of antibody for the production of a multivalent antibody of the invention. The invention further provides a cell for producing a multivalent antibody comprising one or more nucleic acid molecules that alone or together code for a multivalent antibody as claimed.

The invention also provides a method for producing two or more antibodies by the same cell wherein at least one of said antibodies is a multivalent antibody as described herein. This 20 embodiment is now exemplified by the previously described DE/KK heterodimerization system. The invention is, however, not limited to a particular method for enabling heterodimerization of heavy chains. As previously described, the DE-variant and KK-variant preferentially pair to form heterodimers (so-called 'DEKK' bi/multivalent molecules). Homodimerization of DE-variant heavy chains (DEDE homodimers) or KK-variant heavy chains (KKKK homodimers) hardly 25 occurs due to repulsion between the charged residues in the CH3-CH3 interface between identical heavy chains. Introducing a further heavy chain that has either the DE- or the KK- variant heavy chain, allows the production of a further DEKK bi/multivalent molecule. A newly introduced DE-heavy chain (DE2) can associate with the existing KK heavy chain. The cell thus produces two bi/multivalent antibodies a DE1KK and a DE2KK bivalent antibody. If a new KK 30 heavy chain (KK2) is introduced instead of the new DE heavy chain, the bivalent antibodies with the combinations DEKK1 and DEKK2 are produced. The levels at which the different antibodies can be produced by the cell is typically best adjusted by adjusting the relative expression of the DE1/2 and KK1/2 chains with respect to each other. The light chain is typically produced sufficiently to reduce the level of single heavy chains and the level at which one chain is 35 produced is typically sufficient to allow efficient pairing with the complementary chains. In the DE1/2; KK example the DE1 and DE2 heavy chains are preferably produced to a level that together matches the level of the KK heavy chain. The level of the respective antibodies can be adjusted by adjusting the levels at which the DE1 and DE2 are produced relative to each other. For the KK1/2 DE variant the situation is of course similar but now for the KK1 and

KK2 chains in relation to each other. Depending on the number of binding domains or variable domains associated with each of the heavy chains this method allows the production of a variety of different bi/multivalent antibodies. Several non-limiting examples are now here described. In this example heavy chain DE1 has one heavy chain variable region that together with light chain 5 common to all binding domains for a binding domain or variable domain that binds antigen V, heavy chain DE2 has two heavy chain variable regions that together with the common light chain form two binding domains or variable domains that bind antigens W and X. Heavy chain KK has one heavy chain variable region that together with the common light chain forms a binding domain or variable domain that binds antigen Y. Producing these heavy and light chains 10 in a cell will produce an antibody DE1KK and an antibody DE2KK wherein antibody DE1KK is a bivalent antibody that binds antigens V and Y. Antibody DE2KK is a multivalent antibody that binds the antigens W, X and Y. If in the above example DE1 also has two heavy chain variable regions that together with the common light chain form two binding domains or variable domains 15 two multivalent antibodies of the invention are produced. The KK heavy chain can also be provided with an additional heavy chain variable region thereby adding yet further binding domains of the same or different antigen binding specificity. Combination of two or more different heterodimerization domains such as the DE/KK described above and the known in hole domains can add further diversity in the oligoclonal antibody production. For instance adding two heavy chains, one with the knob and the other with the complementary hole allows the 20 production of an independent bi/multivalent antibody comprising of a knob heavy chain and a hole heavy chain. Depending on the number of heavy chain variable regions associated with each heavy chain and depending on whether they are the same or different, a further monospecific antibody, or a further bi or multivalent antibody is produced.

25 According to the invention, there is provided a composition comprising two or more antibodies, at least one of which may be a multivalent antibody of the invention. Such a composition of the invention may comprise two or more multivalent antibodies of the invention. Such a composition may comprise three, four, five or more antibodies, at least one of which 30 may be a multivalent antibody of the invention. Such a composition may comprise three, four, five or more antibodies, all of which may be a multivalent antibody of the invention. In such a composition, one or more of the antibodies present in the composition may have one heavy chain in common.

35 A host cell of the invention may express or may be capable of expressing two or more antibodies, at least one of which may be a multivalent antibody of the invention. A host cell of the invention may express or may be capable of expressing two or more multivalent antibodies of the invention. Such host cells may express or may be capable of expressing three, four, five or more antibodies, at least one of which may be a multivalent antibody of the invention. Such host cells may express or may be capable of expressing three, four, five or more antibodies, all of which may be a multivalent antibody of the invention.

According to the invention, there is thus provided a method for the preparation of a composition comprising two or more antibodies, which method comprises:

providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into two or more antibodies, at least one of which is a multivalent antibody according to the invention; and

10 cultivating said host cell under conditions to provide for expression of the polypeptides and for their assembly into the two or more antibodies, at least one of which is a multivalent antibody according to the invention.

The invention also provides a method for producing two or more antibodies by the same cell wherein at least one of said antibodies is a multivalent antibody as described herein.

The invention provides a method for producing a composition comprising two or more antibodies of which at least one is a multivalent antibody as claimed the method comprising providing a cell with

- nucleic acid that encodes a first heavy chain with a heavy chain variable region that together with a common light chain forms a binding domain or variable domain that binds to a first antigen;

- nucleic acid that encodes a second heavy chain with a heavy chain variable region that together with said common light chain forms a variable domain that binds to a second antigen and a heavy chain variable region that together with said common light chain forms a variable domain that binds a third antigen;

- a nucleic acid that encodes a third heavy chain with a heavy chain variable region that together with said common light chain forms a variable domain that binds to a fourth antigen; and

- a nucleic acid that encodes a polypeptide comprising said common light chain;

25 wherein two or more of said nucleic acids may be physically linked or not and wherein each of said nucleic acids further comprises an expression control sequence to allow expression of the encoded heavy and light chains in said cell and wherein the method further comprises culturing said cell to allow expression of said heavy and light chains and, optionally collecting said two or more antibodies. In one embodiment said first and second heavy chains have a compatible

30 heterodimerization domain preferably a DE/KK heterodimerization domain. In a preferred embodiment said third heavy chain comprises one of the parts of the compatible heterodimerization domain as a result of which two antibodies are produced. In one embodiment the method further comprises providing a collection of cells with said nucleic acid and selecting from said collection a cell with a desired ratio of expression of the respective

35 heavy and light chains. In a preferred embodiment said two or more antibodies are two or more multispecific antibodies. In a preferred embodiment the cells produce essentially equimolar amounts of the two or more antibodies. In some embodiments the cells produce more of one antibody than of another of said two or more antibodies.

Non-human animals

5       Synthesis and expression of multivalent binding proteins has been problematic, in part due to issues associated with identifying a suitable light chain that can associate and express with two or more different heavy chains, and in part due to isolation issues. Further, the art has lacked an array of linkers that permit a diverse array of antibody valence, flexibility with stability and low immunogenicity.

10      The methods and compositions described herein allow for making suitable multivalent binding proteins having binding domains obtained from, derived from, or based on suitable methods. Suitable methods may include phage display methods (including modification of germline sequences generated in phage display systems), and other *in vitro* methods known in the art. A particularly useful method is having a genetically modified non-human animal make, through natural processes of somatic recombination, and affinity maturation, a suitable heavy chain variable domain that can associate and express with a common light chain.

15      In one embodiment, the variable domains used in a multivalent antibody of the invention are obtained from, derived from or based on heavy and light chain variable regions of a non-human transgenic animal that comprises in its germline an unarranged heavy chain variable locus and expresses a single rearranged human light chain variable domain, e.g., a common light chain mammal, such as a rodent. Such a non-human, transgenic animal upon exposure to 20 an antigen will express a diversity of heavy chain variable regions paired with a common light chain, which can then be used to develop nucleic acid sequences encoding heavy chain variable regions obtained from, derived from or based on those from said transgenic animal that are able to be efficiently transformed into host cells for the production of multivalent antibodies.

25      In particular, the human variable region sequences from suitable B cells of an immunized common light chain animal that are genetically engineered to express human light chain variable domains derived from no more than one, or no more than two, human VL gene segments may be used as a source of potential  $V_H$  domains for a multivalent antibody of the invention. The B cells from said animals that are immunized with one or more antigens of interest, which are, in various embodiments, antigens to which the multivalent antibody will bind. 30      Cells, tissues, or serum, splenic or lymph materials of the said animals are screened to obtain heavy chain variable domains (or B cells that express them) that exhibit desired characteristics with respect to the antigens of interest, e.g., high affinity, low affinity, blocking ability, activation, internalization or other characteristics. Because virtually all of the heavy chain variable domains that are generated in response to an antigenic stimulation in said transgenic animal are made in 35 conjunction with the expresses of a human immunoglobulin light chain derived from no more than one, or no more than two,  $V_L$  gene segments, the heavy chain variable regions are capable of expressing and associating with common light chain domains that are expressed in the transgenic animal.

In one aspect, an epitope-binding protein as described herein is provided, wherein human V<sub>L</sub> and V<sub>H</sub> sequences are encoded by nucleic acid based on nucleic acid obtained from the B-cell of a transgenic mouse described herein, and/or a transgenic animal as disclosed in WO2009/157771, incorporated herein by reference, that has been immunized with an antigen 5 comprising an epitope of interest.

Nucleic acid sequences, polypeptides, vectors and cells

The invention further provides: nucleic acid sequences encoding polypeptides or linkers 10 that may be used in the assembly of a multivalent antibody of the invention; vectors comprising such nucleic acid sequences; a cell which is capable of producing a multivalent antibody of the invention; and a method for the preparation of such a multivalent antibody using such a cell.

Multivalent antibodies according to the invention are typically produced by cells that express nucleic acid sequences encoding the polypeptides that together assemble to form an antibody of the invention.

15 Accordingly, the invention provides a linker which comprises an amino acid sequence as set out in any one of SEQ ID NOs 1 to 3 or 5 to 24 or a polypeptide having at least about 85% sequence identity to any one thereto at least about 85% sequence identity to any one thereto, such as at least about 90% sequence identity to any one thereto, for example at least about 95% sequence identity to any one thereto, such as at least about 98% sequence identity 20 to any one thereto, for example at least about 99% sequence identity to any one thereto.

The invention further provides a polypeptide comprising:a VH1-CH1-hinge-based linker-VH2-CH1.

In certain embodiments VH1 and VH2 bind the same epitope. In certain embodiment 25 the VH1 and VH2 bind the same antigen, but different epitopes. And in certain embodiments, VH1 and VH2 bind separate epitopes and antigens.

Also provided by the invention is a nucleic acid sequence encoding such a linker or polypeptide and a vector comprising such a nucleic acid sequence.

30 The nucleic acid sequences employed to make the described polypeptides may be placed in any suitable expression vector and, in appropriate circumstances, two or more vectors in a single host cell.

Generally, nucleic acid sequences encoding variable domains are cloned with the appropriate linkers and/or constant regions and the sequences are placed in operable linkage with a promoter in a suitable expression construct in a suitable cell line for expression.

35 Accordingly, the invention also provides a method for the preparation of an antibody, which method comprises:

providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into a multivalent antibody of the invention; and

cultivating said host cell under conditions to provide for expression of the polypeptides and for their assembly into a multivalent antibody.

Expression of a multivalent antibody

5

Expression of antibodies in recombinant host cells has been described in the art. The nucleic acid molecules encoding the light and heavy chains of an antibody of the invention may be present as extrachromosomal copies and/or stably integrated into the chromosome of the host cell. The latter is preferred in which case a loci may be targeted that is known for lack of 10 gene silencing.

To obtain expression of nucleic acid sequences encoding the polypeptides which assemble as an antibody of the invention, it is well known to those skilled in the art that sequences capable of driving such expression can be functionally linked to the nucleic acid sequences encoding the polypeptides. Functionally linked is meant to describe that the nucleic 15 acid sequences encoding the polypeptides or precursors thereof are linked to the sequences capable of driving expression such that these sequences can drive expression of the polypeptides or precursors thereof. Useful expression vectors are available in the art, e.g. the pcDNA vector series of Invitrogen. Where the sequence encoding the polypeptide of interest is properly inserted with reference to sequences governing the transcription and translation of the 20 encoded polypeptide, the resulting expression cassette is useful to produce the polypeptide of interest, referred to as expression. Sequences driving expression may include promoters, enhancers and the like, and combinations thereof. These should be capable of functioning in the host cell, thereby driving expression of the nucleic acid sequences that are functionally linked to them. Promoters can be constitutive or regulated, and can be obtained from various sources, 25 including viruses, prokaryotic, or eukaryotic sources, or artificially designed.

Expression of nucleic acid sequences of the invention may be from the natural promoter or a derivative thereof or from an entirely heterologous promoter. Some well-known and much used promoters for expression in eukaryotic cells comprise promoters derived from viruses, such as adenovirus, e.g. the E1A promoter, promoters derived from cytomegalovirus (CMV), such as the CMV immediate early (IE) promoter, promoters derived from Simian Virus 40 (SV40), and the like. Suitable promoters can also be derived from eukaryotic cells, such as methallothionein (MT) promoters, elongation factor 1a (EF- 1a) promoter, actin promoter, an immunoglobulin promoter, heat shock promoters, and the like. Any promoter or enhancer/promoter capable of driving expression of a nucleic acid sequence of the invention in a host cell is suitable in the invention. In one embodiment the sequence capable of driving expression comprises a region from a CMV promoter, preferably the region comprising nucleotides -735 to +95 of the CMV immediate early gene enhancer/promoter. The skilled person will be aware that the expression sequences used in the invention may suitably be combined with elements that can stabilize or enhance expression, such as insulators, matrix

attachment regions, STAR elements and the like. This may enhance the stability and/or levels of expression.

Any cell suitable for expressing a recombinant nucleic acid sequence may be used to generate an antibody of the invention. Preferably said cell is adapted for suspension growth.

5 A multivalent antibody of the invention may be expressed in host cells, typically by culturing a suitable cell of the invention and harvesting said antibody from said culture. Preferably said cell is cultured in a serum free medium. An antibody of the invention may be recovered from the cells or, preferably, from the cell culture medium by methods that are generally known to the person skilled in the art.

10 Further provided is an antibody obtainable by a method for producing an antibody according to the invention. The antibody is preferably purified from the medium of the culture.

After recovery, an antibody may be purified from the culture by using methods known in the art. Such methods may include precipitation, centrifugation, filtration, size -exclusion chromatography, affinity chromatography, cation- and/or anion-exchange chromatography,

15 hydrophobic interaction, chromatography, and the like. Affinity chromatography, including based on the linker sequence as a means of separating the multivalent antibody of the invention may be used.

#### Pharmaceutical compositions and methods of use

20 Also provided by the invention is a pharmaceutical composition which comprises an antibody of the invention and a pharmaceutically acceptable carrier and/or diluent.

Accordingly, the invention provides a multispecific antibody as described herein for use in the treatment of the human or animal body by therapy.

25 Further provided by the invention is a method for the treatment of a human or animal suffering from a medical condition, which method comprises administering to the human or animal a therapeutically effective amount of an antibody as described herein.

30 The amount of antibody according to the invention to be administered to a patient is typically in the therapeutic window, meaning that a sufficient quantity is used for obtaining a therapeutic effect, while the amount does not exceed a threshold value leading to an unacceptable extent of side-effects. The lower the amount of antibody needed for obtaining a desired therapeutic effect, the larger the therapeutic window will typically be. An antibody according to the invention exerting sufficient therapeutic effects at low dosage is, therefore, preferred.

35 A reference herein to a patent document or other matter which is given as prior art is not to be taken as an admission that that document or matter was known or that the information it contains was part of the common general knowledge at the priority date of any of the claims.

The disclosure of each reference set forth herein is incorporated herein by reference in its entirety.

For the purpose of clarity and a concise description features are described herein as part of the same or separate embodiments, however, it will be appreciated that the scope of the invention may include embodiments having combinations of all or some of the features described.

5

The following Examples illustrate the invention. For ease of reference, for Examples eight to fifteen, when describing trispecific molecules, the following format is used MFAxMFB:MFC or AntigenAxAntigenB:AntigenC, such that MFA or AntigenA followed by x constitutes the “short arm”, while the x denotes the dimerization, followed by MFB or AntigenB 10 describes the interior position of the long arm, followed by a “:” designating a linker followed by MFC or AntigenC describes MFC or AntigenC at the distal domain of the long arm.

## EXAMPLES

15 **Example 1: Cloning of the variable domains and linker for the generation of a vector capable of expressing a multispecific antibody**

24 linker constructs were cloned in pools according to their size as detailed in Table 2 into the MV1626 vector (see Figure 3), containing the KK residues (L351K, T366K) in the CH3 20 region for the generation of IgG heavy chain heterodimers (WO2013/157954 and WO2013/157953). The constructs were cloned into vector MV1626 using restriction enzyme *Sfi* and *Xho*. All constructs contain sequentially the VH gene of MF1337, a CH1 domain, the linker sequences of which the translations are listed in Table 2, and the VH gene of MF1122. As an example, the DNA sequence of construct MF1337xIgG4 UHxMF1122 is provided below in Table 25 3. Schematically the construct is depicted in Figure 2a. The constructs are based on both the CH1 and linker sequence of the IgG isotype indicated in the name of the constructs. A translation of all 24 CH1 regions in combination with the linker sequences is provided in Figure 5. A translation of all three VH genes and the common light chain gene are provided below in Table 4.

30

Table 2: The sequences of the 24 different linkers/ constructs and naming as used; note there are also differences in CH1 (Figure 5). The linker sequence is indicated below. The respective CH1 sequences together with the linker are indicated in figure 5. The “linker name” refers to the sequence of the indicated together with the CH1 domain.

	Linker name	Sequence	Linker size (aa)
<b>Pool1</b>			
1	IgG4 UH	ESKYGPP (SEQ ID NO: 1)	7
2	IgG1 UH	EPKSCDKTHT (SEQ ID NO: 2)	10
3	IgG2A G4SS	GGGGSGGGGS (SEQ ID NO: 3)	10
4	IgG2A MH	ERKSSVESPPSP (SEQ ID NO: 4)	12
5	IgG2B MH	ERKCSVESPPSP (SEQ ID NO: 5)	12

<b>6</b>	IgG3 UH	ELKTPLGDTTHT (SEQ ID NO: 6)	12
<b>7</b>	IgG4 MH	ESKYGPSPSPSP (SEQ ID NO: 7)	12
<b>8</b>	IgG2A UL	ERKSSVEAPPVAG (SEQ ID NO: 8)	13
<b>9</b>	IgG2B UL	ERKCSVEAPPVAG (SEQ ID NO: 9)	13
<b>10</b>	IgG4 UL	ESKYGPAPPFLGG (SEQ ID NO: 10)	14
<b>11</b>	IgG1 MH	EPKSCDKTHTSPPSP (SEQ ID NO: 11)	15
<b>12</b>	IgG1 G4S	EPKSCDGGGGSGGGGS (SEQ ID NO: 12)	16
<b>13</b>	IgG2 G4SL	GGGGSGGGGSAPPVAG (SEQ ID NO: 13)	16
<b>Pool2</b>			
<b>1</b>	IgG1 UL	EPKSCDKTHTAPELLGG (SEQ ID NO: 14)	17
<b>2</b>	IgG2A H	ERKSSVESPPSPAPPVAG (SEQ ID NO: 15)	18
<b>3</b>	IgG2B H	ERKCSVESPPSPAPPVAG (SEQ ID NO: 16)	18
<b>4</b>	IgG3 ULH	ELKTPLGDTTHTAPEFLGG (SEQ ID NO: 17)	19
<b>5</b>	IgG4 H	ESKYGPSPSPPAPEFLGG (SEQ ID NO: 18)	19
<b>6</b>	IgG1 H	EPKSCDKTHTSPPSPAPELLGG (SEQ ID NO: 19)	22
<b>7</b>	IgG2A R	ERKSSVEAAAKEAAAKAPPVAG (SEQ ID NO: 20)	23
<b>8</b>	IgG2B R	ERKCSVEAAAKEAAAKAPPVAG (SEQ ID NO: 21)	23
<b>9</b>	IgG4 R	ESKYGPPEAAAKEAAAKAPEFLGG (SEQ ID NO: 22)	24
<b>10</b>	IgG1 R	EPKSCDKTHTEAIAAKEAAAKAPELLGG (SEQ ID NO: 23)	27
<b>11</b>	IgG3 R	ELKTPLGDTTHTEAIAAKEAAAKAPEFLGG (SEQ ID NO: 24)	29

**Table 3: DNA sequence of construct MF1337xIgG4 UHxMF1122**

DNA sequence	SEQ ID NO
ggcccagccggccatggccgaggtgcagctggagactgggctgaggtgaagaagccggggcctca gtgaaggctcctgcaaggctctgactacatctcaccaaataatgacatcaactgggtgcgccaggccc ctggacaaggctgaatggatggatggatgagcgctaacactggaaacacaggctatgcacagaattt ccaggcagagtaccatgaccggacacgtccataaacacagcctacatggagctgagcagcctgaca tctggtacacggccgttattctgtcgaggagtgtctttcaagacagagacggccctactatc acttcgctctggacgtctgggccaagggaccacggtcaccgtccatgtccatgtccaccaaggccccag cgtgtccccctggcccccgtcagccggagcaccaggcgagagcaccggccctggctgcgtggtaag gactactcccgagccgtgaccgtgagctggaaacagcggccctgaccagcggcgtcaccatcc ccgcccgtgtcagaggcagcggctgtacagcctgagcagcgtggtagcggccagcagcagcctgg caccatggccatggccatggccatggccatggccatggccatggccatggccatggccatggccatgg aagtacggcccccggaggtgcagctggagactctggggaggcgtggtagcggccatggccatggccatgg gactctctgtcagccctggattcacctcagtagctatggcatgcactggcccgccaggctccagg caaggggctggagtggtggcagttatcatatgatggaaagtaataactatgcagactccgtgaag ggccgattcaccatcccgagacaattccagaacacacgttatctgcaaatgaacacgctgagagctg aggacacggccgtgtattactgtcaagagccctttcacgaccatgcccattggactattggggccaagg taccctgtcaccgtctcgagtt gaggtgcagctggagactgggctgaggtgaagaagccggggcctca gtgaaggctcctgcaaggctctgactacatctcaccaaataatgacatcaactgggtgcgccaggccc ctggacaaggctgaatggatggatggatgagcgctaacactggaaacacaggctatgcacagaattt ccaggcagagtaccatgaccggacacgtccataaacacagcctacatggagctgagcagcctgaca	SEQ ID NO: 25
	SEQ ID NO: 284

tctggtgacacggccgttattctgtgcaggagtgcgtttcaagacagagacggccctactatc acttcgcctggacgtctggggccaaggaccacggtcaccgtctccagtgcaccaagggccccag cgtttccccctggcccccctgcagccggagcaccagcgagagcaccggccctggctgctggtaag gactactccccgagccgtgaccgtgagctggaacagcggccctgaccagcggcgtgcacaccc ccggccgtgcagagcagcggccgtacagcctgagcagcgtggtgcacggccagcagcggctgg caccagacctacacctgcaacgtggaccacaagcccagcaacaccaagggtggacaagcggggag aagtacggccccccgaggtcagctggagtcggggaggcgtggccagcctggaggtccctga gactctctgtgcagcctctggattcacccctgtagctatggcatgcactggcccgccaggctccagg caagggctggagtggtggcagttatcatatgatgaaataactatgcagactccgtgaag ggccgattcaccatctccagagacaattccaagaacacgcgttatctgcaaatacggactattgggccaagg tacccttgtcaccgtctcgagt	
EVQLVETGAEVKKPGASVKVSCKASDYIFTKYDINWVRQAPGQGLEWMGWMS ANTGNTGYAQKFQGRVTMTRDTINTAYMELSSLTSGDTAVYFCARSSLFKTET APYYHFALDVWGQGTTVTVSSASTKGPSVFPLAPCSRSTSESTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLSSVVTVPSSSLGTKTYTCNVDH KPSNTKVDKRVESKYGPPEVQLVESGGVVQPGRLRLSCAASGFTFSSYGM HWVRQAPGKGLEWVAVISYDGSNKYYADSVKGRFTISRDNSKNTLYLQMNSLR AEDTAVYYCARALFTTIAMDYWGQGTLTVSS	SEQ ID NO: 285  Translation of SEQ ID NO: 284

**Table 4: Translations of all three VH genes and the common light chain gene**

Protein sequence	Description	SEQ ID NO
EVQLVESGGVVQPGRLRLSCAASGFTFSS YGMHWVRQAPGKGLEWVAVISYDGSNKYYA DSVKGRFTISRDNSKNTLYLQMNSLRAEDTAV YYCARALFTTIAMDYWGQGTLVT	Translation of MF1122	SEQ ID NO: 26
EVQLVESGGVVQPGRLRLSCAASGFTFSS YGMHWVRQAPGKGLEWVAVISYDGSNKYYA DSVKGRFTISRDNSKNTLYLQMNSLRAEDTAV YYCARALFTTIAMDYWGQGTLVT	Translation of MF1122	SEQ ID NO: 286
EVQLVESGGGLVQPGGSLRLSCAASGFTFSS YAMSWVRQAPGKGLEWVSAISGSGGSTYYA DSVKGRFTISRDNSKNTLYLQMNSLRAEDTAV YYCARADWWATFDYWGQGTLVT	Translation of MF1025	SEQ ID NO: 27
EVQLVESGGGLVQPGGSLRLSCAASGFTFSS	Translation of MF1025	SEQ ID

YAMSWVRQAPGKGLEWVSAISGSGGSTYYA DSVKGRFTISRDNSKNTLYLQMNSLRAEDTAV YYCARADWWATFDYWQGQTLTVSS		NO: 287
EVQLVETGAEVKKPGASVKVSCKASDYIFTKY DINWVRQAPGQGLEWMGWMMSANTGNTGYA QKFQGRVTMTRDTSINTAYMELSSLTSGDTAV YFCARSSLFKTETAPYYHFALDWGQGTTVT	Translation of MF1337	SEQ ID NO: 28
EVQLVETGAEVKKPGASVKVSCKASDYIFTKY DINWVRQAPGQGLEWMGWMMSANTGNTGYA QKFQGRVTMTRDTSINTAYMELSSLTSGDTAV YFCARSSLFKTETAPYYHFALDWGQGTTVT SS	Translation of MF1337	SEQ ID NO: 288
DIQMTQSPSSLSASVGDRVTITCRASQSISSYL NWYQQKPGKAPKLLIYAASSLQSGVPSRFSG SGSGTDFTLTSSLQPEDFATYYCQQSYSTPP TFGQGQTKVEIKRTVAAPSVFIFPPSDEQLKSGT ASVVCLLNNFYPREAKVQWKVDNALQSGNSQ ESVTEQDSKDSTYLSSTTLSKADYEKHKVY ACEVTHQGLSSPVTKSFNRGEC	Translation of common light chain (cLC)	SEQ ID NO: 29

The inserts and the vector were digested at 50°C for 2h using *Sfi* restriction enzyme, followed by 2h at 37°C with *Xho* I enzyme. The digested DNA was loaded onto a 0.8% agarose gel and run for 2 hours at 100 volts. The digested vector and inserts were subsequently isolated from the gel using Qiagen QIAquick Gel Extraction kit before overnight ligation at 16°C using T4 DNA ligase in 1/5 ratio (w/w vector/insert). 50 µL of DH5α-T1R competent *E. coli* were transformed in presence of 5µL of the ligation mix following a heat shock procedure of 30min on ice followed by 2 min at 42°C and 2 min on ice. Transformed bacteria were plated on LB agar supplemented with Ampicillin and incubated overnight at 37°C. Single colonies were picked and mixed with 100 µL of sterile deionised water and used for colony PCR using primers DO\_2130 and DO\_1056 to confirm the presence of the insert, followed by sequence PCR with the BigDye® Terminator v1.1 Cycle Sequencing Kit (Thermofisher) for clone confirmation using primer DO\_2130.

Single colonies of confirmed clones were used to inoculate 4mL of LB-Amp. The overnight cultures at 37°C were prepared in 24 wells format mini-prep using the QIAGEN Plasmid Mini Kit according to the manufacturer manual. After elution from the column, the purified DNA was precipitated by adding 0.7 volumes of room-temperature Isopropanol. The DNA pellet was washed with 1ml of 70% Ethanol and air dried in sterile conditions and resuspended in sterile Tris-EDTA buffer before storage at -20°C. The final constructs were dideoxy sequenced using primers DO\_1488, DO\_1056 and DO\_2130 for the insert as well as

primers DO\_0182 and DO\_0091 for the CH2/CH3 region using the BigDye® Terminator v1.1 Cycle Sequencing Kit.

All primer sequences are set out in Table 5.

Sequencing showed that all constructs had successfully been prepared.

5

Table 5: primer sequences

Primer	Sequence	Description	SEQ ID NO
DO_0091	CCTCATGCATCACGGAGCATG	CH3_rev	SEQ ID NO: 30
DO_0182	CAAAGGCCAAACTCTCCACTC	CH2 fwd	SEQ ID NO: 31
DO_1056	CGCTGTGCCCCAGAGGTGC	VH_rev	SEQ ID NO: 32
DO_1488	GTACCGGTGAATTGGCCGG	VH_fwd	SEQ ID NO: 33
DO_2130	GCGCCCTACTATCACTTCGCT CTGG	MF1337 CDR3 fwd	SEQ ID NO: 34

**Example 2: Transfection and IgG purification**

10 The expression vectors generated in Example 1 were combined with the vector MG1025C377 (Figure 4) expressing the second heavy chain of the base antibody portion of the multispecific antibodies, bearing the L351D-L368E mutations in the CH3 region (WO2013/157954 and WO2013/157953) and the Thyroglobulin Fab gene of antibody MF1025 (see Example 2 of WO2013/157953). Expression of the two heavy chains together with a  
15 common light chain leads to the production of the trispecific antibody as shown in Figure 2b.

FreeStyle 293-F cells (Thermofisher) were used for expression of the designed antibodies in a 24 wells plate format. Two days before transfection, FreeStyle 293-F cell stock was split in 293-F culture medium in a 1:1 ratio and incubated overnight at 37°C and 8%CO<sub>2</sub> at an orbital shaking speed of 155 rpm. Cells were diluted on the day before transfection to a  
20 density of 5 x 10<sup>5</sup> cells/mL 4ml of the suspension cells were seeded into a 24 deep wells plate, covered with a breathable seal and incubated overnight at 37°C and 8% CO<sub>2</sub> at an orbital shaking speed of 285 rpm. On transfection day, 4.8 ml 293-F culture medium were mixed with  
240 µg of polyethylenimine (PEI) linear (MW 25,000). For each IgG to be produced, 200 µL of the 293F culture medium-PEI mix was added to 8 µl of DNA (for IgG heterodimers 4 µl of DNA  
25 encoding each heavy chain) as detailed in Table 6. The mixture was incubated for 20 minutes at room temperature before gently adding to the cells. On the day after transfection Penicillin-Streptomycin (Pen Strep) diluted in 500 µL 293F medium was added to each well. The plates were incubated at 37°C and 8% CO<sub>2</sub> at an orbital shaking speed of 285 rpm until harvest seven  
30 days after transfection. Plates were centrifuged 5 min at 500g, supernatants containing IgGs were filtered using 10-12 µm melt blown polypropylene filter plates and stored at -20°C prior to purification.

Various control antibodies were also expressed, namely:

Bivalent anti-Tetanus Toxoid antibody using Fab MF1337 (using vector MG1337C057)

Bivalent anti-thyroglobulin antibody using Fab MF1025 (using vector MG1025C059)

Bivalent anti-Fibrinogen antibody using Fab MF1122v (using vector MG1122C057)

5 MG1337C057 indicates a construct expressing the VH region of MF1337 from vector MV1057. MG1025C059 indicates a construct expressing the VH region of MF1025 from vector MV1059. MG1122C057 indicates a construct expressing the VH region of MF1122 from vector MV1057. MV1057 (Figure 6) and MV1059 are vectors expressing monospecific-bivalent human IgG1 molecules. MV1057 and MV1059 are essentially the same vectors resulting in expression 10 of identical IgG1 molecules.

Bispecific anti-thyroglobulin x anti-Tetanus Toxoid antibody combining Fab MF1337 and Fab MF1025 (using MG1025C377 x MG1337C260)

Bbispecific anti-thyroglobulin x anti Fibrinogen antibody combining Fab MF1122 and Fab MF1025 (using MG1025C377 x MG1122C260)

15 MG1025C377 (Figure 4) expresses the heavy chain variable domain of antibody MF1025 in the context of a human IgG1 heavy chain containing the L351D, L368E (DE) mutations. MG1337C260 (Figure 7) expresses the heavy chain variable domain of antibody MF1337 in the context of a human IgG1 heavy chain containing the L351K, T366K (KK) mutations. MG1122C260 expresses the heavy chain variable domain of antibody MF1122 in 20 the context of a human IgG1 heavy chain containing the L351K, T366K (KK) mutations.

**Table 6: Transfection scheme for IgG production.**

construct 1	construct 2
MG1025C377	MF1337xIgG4 UHxMF1122
MG1025C377	MF1337xIgG1 UHxMF1122
MG1025C377	MF1337xIgG2A G4SSxMF1122
MG1025C377	MF1337xIgG2A MHxMF1122
MG1025C377	MF1337xIgG2B MHxMF1122
MG1025C377	MF1337xIgG3 UHxMF1122
MG1025C377	MF1337xIgG4 MHxMF1122
MG1025C377	MF1337xIgG2A ULxMF1122
MG1025C377	MF1337xIgG2B ULxMF1122
MG1025C377	MF1337xIgG4 ULxMF1122
MG1025C377	MF1337xIgG1 MHxMF1122
MG1025C377	MF1337xIgG1 G4SxMF1122
MG1025C377	MG1337C260
MG1025C377	MG1122C260
MG1337C057	-*

<b>MG1122C057</b>	-*
<b>MG1025C059</b>	-*
<b>MG1025C377</b>	MF1337xIgG2 G4SLxMF1122
<b>MG1025C377</b>	MF1337xIgG1 ULxMF1122
<b>MG1025C377</b>	MF1337xIgG2A HxMF1122
<b>MG1025C377</b>	MF1337xIgG2 BHxMF1122
<b>MG1025C377</b>	MF1337xIgG3 ULHxMF1122
<b>MG1025C377</b>	MF1337xIgG4 HxMF1122
<b>MG1025C377</b>	MF1337xIgG1HxMF1122
<b>MG1025C377</b>	MF1337xIgG2A RxMF1122
<b>MG1025C377</b>	MF1337xIgG2B RxMF1122
<b>MG1025C377</b>	MF1337xIgG4 RxMF1122
<b>MG1025C377</b>	MF1337xIgG1 RxMF1122
<b>MG1025C377</b>	MF1337xIgG3 RxMF1122
<b>*= bivalent controls</b>	

Following harvesting, antibodies were purified in 24 well format as follows: supernatants were mixed with 50  $\mu$ L 1M Trizma pH8 and 100  $\mu$ L of ProteinA Sepharose CL-4B beads (50% v/v, G.E Healthcare Life Sciences) and incubated at 25°C for 2h at 600 rpm orbital shaking.

5 Beads were vacuum filtered and washed 2 times with 3 mL PBS pH7.4. The elution of the antibodies was performed by adding 200  $\mu$ L of citrate buffer 0.1M, pH3 followed by neutralization with 300  $\mu$ L 1M Trizma pH8. Purified IgG fractions were immediately buffer exchanged to PBS pH7.4. IgG samples were transferred into a 30 kDa 96 well filter plate, polyethersulfone membrane and centrifuged at 1500g 4°C until a volume of 10  $\mu$ L was left per well. 200  $\mu$ L of PBS were added in each well, samples were mixed at 500 rpm for 3 min before IgGs are collected for storage at 4°C. IgG concentration was determined by Octet and ProteinA biosensors (Pall ForteBio). Human IgG was used as standard in seven 2 folds dilutions starting from 192  $\mu$ g/ml to 3  $\mu$ g/ml. Concentrations of IgG samples were determined in duplicate. Reduced and Non-Reduced SDS-PAGE were performed for all thirty produced IgGs thus including the controls. The results are presented in Figure 8. In NR conditions, the expected product sizes of ~200 kDa for the trispecific, multivalent antibodies and ~150 kDa for the control monoclonal IgGs and Biclonics® were observed. For R conditions, product sizes of ~25 kDa (LC), ~50 kDa (HC/1VH) and ~75kDa (HC/2VH) for the trivalent antibodies were observed. The control IgGs band sizes of ~25 kDa (LC) and ~50 kDa (HC) were as expected. The results also showed bands at ~150 kDa for the trispecific constructs. These are the homodimers resulting from association of DE heavy chains which may be the result of higher expression levels of the DE containing shorter heavy chain over the longer KK containing heavy chain.

**Example 3: Binding activity of the Fab domains in the VH1, VH2 and VH3 position measured in ELISA**

Binding activity of the three Fab domain in each construct was checked by ELISA using the tetanus toxoid, fibrinogen and thyroglobulin antigens and huEGFR-Fc antigen as a negative 5 control (see Table 7 for coating conditions, supplier and catalogue numbers).

Each multispecific IgG sample was first diluted to 10 µg/ml in PBS and analysed in titration on Fibrinogen, Tetanus Toxoid and Thyroglobulin; in four 5-fold dilutions, from 10 to 0.08 µg/ml. All 30 samples were analyzed on huEGFR-Fc at 10 µg/mL. Appropriate amounts of antigen in PBS were prepared. 50 µl of diluted antigen solution were added per ELISA plate well 10 and coated o/n at 4°C. Plates were washed twice with wash buffer (PBS/Tween). Wells were blocked for 1 hour at RT with 300 µl / well block buffer (PBS/2%BSA). During incubation appropriate IgG dilutions were made in block buffer. Plates were emptied by inverting above sink followed by slapping on tissue. 50 µl of diluted IgG samples and controls were added to the wells of the blocked plate, covered with seal and incubated 60 minutes at RT. Plates were 15 washed three times with wash buffer (PBS/0.05%Tween). Diluted detection antibody (mouse anti-Human IgG HRP-conjugated; Becton Dickinson, cat. no. 555788), 1/2000 in block buffer was added at 50 µl per well. The plate was covered with seal and incubated for 60 minutes at RT. The plate was washed three times with wash buffer. TMB substrate solution (BD, OptEIA™ cat. no. 51-2606KC) was made by mixing reagents A and B in 1:1 ratio and adding 50 µl per 20 well and developed for (maximal) 10 minutes. 50 µl of 1 M H<sub>2</sub>SO<sub>4</sub> was added to each well to stop the staining reaction.

The plates were read at A450 using a 'BioTek Elx808 ELISA plate reader. Binding curves were plotted using GraphPad Prism 7 and Area Under Curve (AUC) calculated for each antigen ELISA of each construct and listed in Table 8. A small variation in AUC is seen for 25 binding to Thyroglobulin (VH1, on the DE-arm, see Figure 1) and Tetanus Toxoid (VH3, on the tip of the KK arm, see Figure 1) of 12% and 8%. A larger variation is seen for the Fibrinogen-arm (this is VH2, Figure 1). This indicates that accessibility or affinity of the Fab domain in the VH2 position depends on the linker that connects the Fab in the VH2 position to the Fab in the VH3 position. All linkers provide VH2s that are functional.

30

**Table 7. List of antigens used for ELISA**

Antigen	Coating concentration (µg/ml)	Coating Buffer	Supplier	Catalogue number
Fibrinogen	10	PBS	Sigma	F4753
Tetanus toxoid	2	PBS	Statens institute	T162-2
Thyroglobulin	10	PBS	Sigma	T1126-500MG

huEGFR-Fc	2.5	PBS	R&D Systems	344-ER
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**Table 8: Area Under Curve values resulting from ELISA binding assays of the 24 trispecific, multivalent antibodies listed in Table 6. The antibodies were titrated in ELISA for binding to three different antigens. The resulting AUC values were sorted based on the binding activity towards Fibrinogen, resulting from the Fab in the VH2 position. This identified that a range of binding activities of the Fab in the VH2 position exists. The fifteen constructs with the highest activity for the Fab in the VH2 position were prioritized for further experimentation, the six constructs with the lowest activity are indicated in Italics. ND means Not Done**

10

Construct	Tetanus Toxoid, VH3	Fibrinogen, VH2	Thyroglobulin, VH1
IgG1 MH	2.50	1.68	2.29
IgG1 H	2.55	1.64	2.44
IgG1 R	2.46	1.63	2.44
IgG1 G4S	2.54	1.56	2.19
IgG1 UH	2.33	1.42	2.31
IgG3 R	2.48	1.34	2.48
IgG3 UH	2.45	1.30	2.40
IgG2A R	2.51	1.30	2.40
IgG2A MH	2.43	1.25	2.31
IgG3 ULH	2.42	1.23	2.38
IgG2B R	2.51	1.21	2.42
IgG4 MH	2.46	1.18	2.35
IgG4 UL	2.50	1.17	2.25
IgG2A H	2.39	1.16	2.37
IgG2B H	2.42	1.16	2.36
IgG2A G4SS	2.34	1.12	2.32
IgG2B MH	2.48	1.11	2.25
IgG4 UH	2.31	1.08	2.33
IgG2A UL	2.36	1.07	2.27
IgG2B UL	2.46	1.03	2.36
IgG2 G4SL	2.29	1.03	2.31
IgG1 UL	ND	ND	ND
IgG4 H	ND	ND	ND
IgG4 R	ND	ND	ND

**Example 4: Stability of binding activity**

Stability of binding activity of the three Fab domains in the trivalent antibody constructs was analyzed following four accelerated stress conditions. The samples were diluted to 10

5 µg/ml in PBS and incubated for 1 month at 4°C. The samples were diluted to 10 µg/ml in D10F medium and incubated for 7 days at 40°C. The samples were also diluted to 10 µg/ml in D10F medium and incubated for 2 days at 50°C. The samples were also diluted in PBS and subjected to five freeze-thaw cycles (5XFT).

Following these accelerated stress conditions binding activity towards the antigens 10 recognized by the three Fab domains was analyzed in ELISA as described before. Area Under Curve were calculated and tabulated.

15 Stress applied at 40°C only significantly affected binding of the Fab in the VH2 position, binding to Fibrinogen to different degrees in the different constructs tested. Stress at 4°C, at 50°C and 5xFT affected binding of all three Fab domains to different degrees in the different constructs tested.

20 The binding activities were ranked for each antigen and stress condition and the 16 most optimal constructs under each stress condition identified for each Fab position. The number of times that a construct was amongst the 16 most optimal constructs was added up and used to rank all constructs based on conservation of binding activity of the three Fab under accelerated stress conditions.

The results are set out in Figure 9 indicate that there is a range of stabilities of the 25 different constructs under accelerated stress conditions. Stability of binding activity of the three Fab domains in the 21 produced trivalent antibody constructs was analyzed following four accelerated stress conditions. ELISA data (AUC) are tabulated. The binding activities were ranked and the 16 most optimal constructs under each stress condition identified for each Fab position. The number of times that a construct was amongst the 16 most optimal constructs was added up and used to rank all constructs based on conservation of binding activity of the three Fab under accelerated stress conditions.

All antibodies are stable and some are more stable than others.

30

**Example 5: Large scale transfection and IgG purification**

Eighteen constructs were selected for a large scale production for further analysis as follows: IgG1 MH, IgG1 H, IgG1 R, IgG1 G4S, IgG1 UH, IgG3 R, IgG3 UH, \*\*IgG2A R, IgG2A

35 MH, IgG3 ULH, IgG2B R, IgG4 MH, IgG4 UL, IgG2A H, IgG2B H, \*IgG1 UL, \*IgG4 H, \*IgG4 R. As a control, the following productions were included: Bispecific anti-thyroglobulin x anti-Tetanus Toxoid (using MG1025C377 x MG1337C260 described before in Example 2) and bispecific anti-thyroglobulin x anti-Fibrinogen (using MG1025C377 x MG1122C260 described before).

DNA of these constructs was prepared as described before. Multispecific IgGs were transfected as described before by co-transfection of the constructs listed in Table 6. Those constructs selected were produced at larger scale. Two days before transfection, FreeStyle HEK293-F cell stock was split in 293-F culture medium in a 1:1 ratio in 100ml final volume per 5 500 ml culture flask and incubated at 37°C and 8% CO<sub>2</sub>, at an orbital shaking speed of 155 rpm. One day before transfection cells were counted and a cell suspension with a density of 5.0x10<sup>5</sup> cells/ml prepared by diluting the cells with 293-F culture medium. Cells were then seeded at 100 ml cell suspension per T500 flask and incubated at 37°C and 8% CO<sub>2</sub> at an orbital shaking speed of 155 rpm. Next day the cells were transfected. A mix of 293-F culture 10 medium, PEI and DNA was prepared by mixing 7.5 ml 293-F culture medium, 187.5 µl PEI stock at 1 µg/µl) and 150 µl DNA at 0.5 µg/µl. This was incubated for 20 minutes at RT and then added to the cells that were then incubated at 37°C and 8% CO<sub>2</sub> at an orbital shaking speed of 155 rpm for 7 days.

Supernatant containing antibody protein was centrifuged at 1000 g for 10 minutes to 15 remove the cells. The supernatant was filtered using a 0.45 µm filter. The IgG was purified from the supernatant using an ÄKTAexplorer 100 system (GE Health Care) and protein A affinity chromatography followed by desalting. A HiTrap MabSelect SuRe 5 ml column and HiTrap 5 ml Desalting column (GE Health Care) were used according to the manufacturer's instructions. IgG concentration was determined by OD280 absorption. This yielded 0.8-4.9 mg IgG in PBS for all 20 constructs. The generated proteins were analyzed on SDS-PAGE, reduced and non-reduced, as described above in Example 2. The data confirmed the data as found in Example 2 and are not provided here.

HP-SEC was performed to establish expression ratio's between the two heavy chains that make up base antibody portion of the the multispecific antibodies. Because of the size 25 difference of the two heavy chains in the trivalent constructs the halfbodies and homodimers can be identified and quantified in High Performance Size Exclusion Chromatography (HP-SEC). HP-SEC was performed using a Dionex HPLC system equipped with a TSK guard column SWXL (Tosoh Bioscience cat# 08543) and a TSK-gel column G3000SWXL (Tosoh Bioscience cat# 08541). For each analysis, 20 µg protein sample in PBS was injected to the 30 column, which was run using 200 mM Sodium Phosphate, 50 mM NaCl as running buffer at a flow speed of 1 mL/min at 4°C. The chromatograms were analyzed for retention times and relative peak areas based on the UV280 results using Chromeleon 6.80 software. The ratio between the amounts of trivalent IgG/DEDE homodimer were calculated and are presented in Table 9 below. The constructs that have a Trivalent/DEDE ratio above average are presented in 35 Italics.

**Table 9:** HP-SEC was performed to establish expression ratio's between the two heavy chains that make up the base antibody portion of the multispecific antibodies The ratio between the amounts of trivalent IgG/DEDE homodimer were calculated and are tabulated

Construct	Trivalent versus DEDE ratio
<i>IgG1 MH</i>	6.1
<i>IgG1 H</i>	5.7
<i>IgG1 UH</i>	5.6
<i>IgG2B H</i>	5.6
<i>IgG2A MH</i>	5.4
<i>IgG1 UL</i>	5.3
<i>IgG2A H</i>	5.3
<i>IgG1 G4S</i>	5.1
<i>IgG2B R</i>	4.4
<i>IgG1 R</i>	4.3
<i>IgG4 H</i>	3.1
<i>IgG4 UL</i>	3.1
<i>IgG4 MH</i>	2
<i>IgG4 R</i>	2
<i>IgG3 R</i>	1.7
<i>IgG3 ULH</i>	1.7
<i>IgG3 UH</i>	1.7
<i>IgG2A R</i>	NA

The stability of the large-scale purified IgGs (including the two controls) was assessed after different stress conditions as described before; after five freeze-thaw cycles in PBS, after 5 one week incubation at 40°C in D10F medium, after one week incubation at 50°C in D10F medium and after one week incubation at 50°C in PBS. Performance of the stressed samples was compared to performance of the same samples after a one week incubation at 4°C (control). To that purpose, purified IgGs from the 100 ml productions were diluted in PBS at 0.2 mg/mL and divided in two batches: one for stability tests in PBS at 0.2 mg/ml (4°C, 3xFT and 10 50°C), one diluted to 0.1mg/ml in D10F for stress tests at 4°C, 40°C and 50°C as described above. For sample IgG1 H, a concentration of 0.194mg/mL was used.

Following these accelerated stress conditions binding activity towards the antigens recognized by the three Fab fragments was analyzed in ELISA as described before. Area Under Curve were calculated and tabulated (see Figure 10). Percentage remaining binding 15 activity after stress compared to activity of the control sample stored at 4°C was calculated for each binding activity. The samples were ranked based on these percentages for each binding activity against each target at each stress condition. The percentages above average were indicated and the number of times each sample performed above average was added up and is presented in the last column of the Table in Figure 10. This shows that there is a range in the 20 stability of the constructs as measured by the binding activity of the three Fab arms after stress.

**Example 6: Stability analysis of 18 multivalent IgG constructs**

5 Stability analysis was performed on the 18 multivalent constructs identified in Figure 10 and as previously described in Example 5. In addition, four control antibodies were used having heavy chain binding domains comprising MF6744 (SEQ ID NO:91), MF1337 (SEQ ID NO: 28 and SEQ ID NO:288) previously described in WO 2018/056821 A1, which is incorporated by reference, and MF1122 (SEQ ID NO:26 and SEQ ID NO:286), coupled to a cLC namely:

10 monospecific anti-CD137 antibody using Fab MF6744/cLC;  
 monospecific anti-Fibrinogen antibody using Fab MF1122/cLC;  
 monospecific anti-Tetanus Toxoid antibody using Fab MF1337/cLC; and  
 bispecific anti-Fibrinogen x anti-Tetanus Toxoid antibody containing Fab MF1122/cLC and Fab MF1337/cLC as a DEKK bispecific control.

15

The list of samples tested for stability analysis is provided below in Table 10.

**Table 10. List of samples tested for stability analysis: 18 multivalent antibodies and 4 control antibodies**

IgGs	CH3	Fab(s)	Comments
Ctrl AA	WT	MF6744	Monospecific IgG
Ctrl BB	WT	MF1122	Monospecific IgG
Ctrl CC	WT	MF1337	Monospecific IgG
Ctrl BC	DE-KK	MF1122-MF1337	Bispecific – DEKK
Sample 1	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG1 H (SEQ ID NO: 19)
Sample 2	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG2A MH (SEQ ID NO: 4)
Sample 3	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG4 UL (SEQ ID NO: 10)
Sample 4	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG1 G4S (SEQ ID NO: 12)
Sample 5	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG1 MH (SEQ ID NO: 11)
Sample 6	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG1 R (SEQ ID NO: 23)
Sample 7	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG1 UH (SEQ ID NO: 2)

Sample 8	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG1 UL (SEQ ID NO: 14)
Sample 9	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG2A H (SEQ ID NO: 15)
Sample 10	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG2A R (SEQ ID NO: 20)
Sample 11	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG2B H (SEQ ID NO: 16)
Sample 12	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG2B R (SEQ ID NO: 21)
Sample 13	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG3 R (SEQ ID NO: 24)
Sample 14	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG3 UH (SEQ ID NO: 6)
Sample 15	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG3 ULH (SEQ ID NO: 17)
Sample 16	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG4 H (SEQ ID NO: 18)
Sample 17	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG4 MH (SEQ ID NO: 7)
Sample 18	DE-KK	MF6744xMF1337:MF1122	Trispecifics – IgG4 R (SEQ ID NO: 22)

Stability of the 18 multivalent constructs and 4 control antibodies was analyzed following 4 different conditions. Accordingly, the 22 samples (18 trivalent + 4 controls) were diluted to 0.2 mg/ml in PBS and subjected to:

5                   1 month at 4°C (T0) seen as reference;  
7 days at 50°C;  
5x freeze-thaw (FT) cycles at -80°C; or  
4 hours of shaking at 400 rpm at room temperature.

10 After each of these four conditions, stability was analyzed using 7 different methods,  
namely:

UV-Vis absorption spectroscopy: after the subtraction of background buffer absorption and light-scatter due to aggregates, absorbance is tested at 350nm to provide information on the aggregation state of the samples as explained in Eckhardt, 1994; Mulinacci, 2011b and Peters, 2013.

90° light-scattering spectroscopy: in solution, the scatter intensity of light may be affected by different factors, such as protein concentration, refractive index, particle size and shape, and the wavelength of the incident light. This method is used to study protein aggregation as reported in Cappelle, 2005; Demeule, 2007a and b; Mulinacci, 2011a and 2013; 5 Luca, 2010; Patois, 2011 and 2012; Peters 2013.

Tryptophan intrinsic fluorescence emission, expressed as % change compared to T0: changes to the hydrophobicity and rigidity of the environment can be measured through the fluorescence emission of tryptophan (Capelle, 2005; Demeule, 2007a and b and 2009; Mulinacci, 2011a and b; Luca, 2010; Patois, 2011; Peters, 2013).

10 1,8-ANS fluorescence emission, expressed as % change compared to T0: as an uncharged small hydrophobic fluorescent probe, 1-anilinonaphthalene-8-sulfonic acid (1,8-ANS) becomes fluorescent in water when bound to electrostatic pockets in proteins, protein aggregates, detergent micelles, leachables, membranes and cellular components and can therefore be used to study membrane surfaces and proteins (Demeule, 2009; Mulinacci, 2011a 15 and b; Luca 2010).

Nile Red fluorescence emission, expressed as % change compared to T0: as an uncharged small hydrophobic fluorescence probe, Nile Red is influenced by the polarity of the environment and can be used to analyze protein degradation, protein aggregation, lipid structures, protein unfolding (Sackett and Wolff, 1987).

20 Nile Red fluorescence microscopy, where the number of particles/ 1ul is measured: Nile Red is used to stain the samples in order to visualize protein aggregates for fluorescence microscopy using a Leica DMi8 microscope (Demeule, 2007a and b, and 2009; Mulinacci, 2011b and 2013; Patois 2011).

25 Dynamic light-scattering, expressed as % monomer (intensity calculation) change compared to T0: dynamic light-scattering is measured using a NanoFlex instrument. The laser passing through the optical fiber is scattered and reflected from particles towards the detector which measures the scattered light intensity in order to determine the size distribution profile of the particles.

30 The readout of these methods relates to the aggregation, fragmentation and unfolding of the proteins which is a measure of the stability of the proteins. Results showed that 4 hours of shaking at 400 rpm had the biggest impact on the stability of the antibodies. The control monospecific PG1337/MF1337 antibody was the most affected by the stress conditions. Therefore, as all 18 multivalent constructs and the bispecific control contained the PG1337/MF1337 Fab, all results were normalized with PG1337 set as the threshold.

35 The stability of the molecules was calculated by combining the scores of all the 7 methods during the 4 different conditions. The results can be seen in Figure 12 and reveal that the constructs have a range of stabilities, which for an array of trispecific molecules demonstrated superior stability to the control bispecific IgG.

### Example 7: Bioinformatic Linker Characterization

Eight linkers as set out in Table 11 below and depicted in Fig. 13 were further characterized. A flexibility prediction was obtained for each of these sequences using the Karplus and Schulz flexibility Prediction method (which computes the average of the flexibility index for each amino acid in the sequence). The flexibility index is derived from the average properties of each amino acid in protein structures as described in Karplus PA, Schulz GE. Prediction of Chain Flexibility in Proteins - A tool for the Selection of Peptide Antigens. Naturwissenschaften 1985; 72:212-3; <http://tools.immuneepitope.org/bcell/>). In table 11 we designate a linker as rigid (R) if the KS score is 1.015 or less, partially flexible if the KS-score is from about 1.015 to 1.04. A flexible sequence, for the purposes of this invention, is a sequence having a Karplus and Schulz flexibility Prediction greater than 1.04..

**Table 11: Eight linkers and flexibility as determined according to Karplus and Schulz**

#	Linker	Sequence	Flexibility	SEQ ID NO
1	IgG1 G4S	EPKSCDGGGGSGGGGS F	F	12
2	IgG1 H	EPKSCDKTHTSPPSPAPELLGG	F	19
3	IgG1 MH	EPKSCDKTHTSPPSP	F	11
4	IgG1 UH	EPKSCDKTHT	Med	2
5	IgG2A H	ERKSSVESPPSPAPPVAG	F	15
6	IgG2A MH	ERKSSVESPPSP	R	4
7	IgG2B H	ERKCSVESPPSPAPPVAG	Med	16
8	IgG2B R	ERKCSVEEAAAKEAAAKAPPVAG	R	21

15 A second bioinformatic prediction of these linkers was obtained using Rosetta local structure prediction. Here Rosetta fragment picker was used to provide local structure predictions as described in Gront D, Kulp DW, Vernon RM, Strauss CEM, Baker D (2011) Generalized Fragment Picking in Rosetta: Design, Protocols and Applications. PLoS ONE 6(8): e23294. <https://doi.org/10.1371/journal.pone.0023294>. For this use of the prediction tool a minimum number of 40 residues is preferred, and accordingly glycine residues were introduced at the termini of the linkers to make the sequences each 40 residues long, with the linker sequence in the middle. These linkers were characterized for secondary structure, running the sequences through the Rosetta fragment pipeline which finds close local sequence matches in structures in the ProteinDataBank and using these close sequence-sequence matches to predict local structure. The centered fragment was then visualized for each of the 8 sequences above. Fig. 13.

20

25

The summary of the results is set out below in Table 12: F = flexible; M = medium; R = rigid; C = coil = flexible; H = helix = rigid; and E = strand = medium, which demonstrates general agreement with the Karplus and Schultz score based on predicted secondary structure.

Table 12: Summary of flexibility results.

#	Linker	SEQ ID NO	Karplus Schultz	Fragment Prediction
1	IgG1 G4S	12	F	CE
2	IgG1 H	19	F	CH
3	IgG1 MH	11	F	C
4	IgG1 UH	2	M	H
5	IgG2A H	15	F	C
6	IgG2A MH	4	R	H
7	IgG2B H	16	M	CE
8	IgG2B R	21	R	H

## 5 Example 8: Generation of Anti-CD3, PD-L1, and EGFR binding domains

Mice used for immunization.

For generation of human antibodies binding to CD3, EGFR and PD-L1, mice transgenic for the human common light chain and for a human heavy chain (HC) minilocus (comprising a 10 selection of human V gene segments, all human Ds and all human Js) (see W02009/157771 incorporated herein by reference) can be immunized with either DNA encoding the proteins or recombinant DNA, seen below. These mice are referred to as 'MeMo®' mice. For specific heavy chain variable regions, or trivalent multimers having the sequences disclosed herein, they can be produced by any means known to persons of ordinary skill in the art.

### 15 Protein immunizations

'MeMo®' mice were immunized by subcutaneous injections with recombinant protein and Gerbu adjuvant MM (Gerbu Biotechnik c#3001). Recombinant huPDL1-His (SinoBiological; cat.no. 10084-H08H) proteins were used for immunizations. Mice were immunized with 40 µg recombinant protein in PBS mixed with 40 µl of adjuvant in a total volume of 100 µl.

20 Subsequently mice were boosted on day 14 and 28 with 20 µg of recombinant protein in PBS together with 20 µl of adjuvant in a total volume of 50 µl. Mouse serum was collected at day 35 to determine serum titers. Mice with low serum titers received additional cycles of booster immunizations and serum analyses. Each cycle consisted of two weekly immunizations using 20 µg of recombinant protein in 50 µl PBS followed one week later by serum collection for titer 25 analysis. Mice showing high serum titers against the human and macaque target received a final boost immunization consisting of daily injections with 20 µg of recombinant protein in 50 µl PBS on three consecutive days. One day after the final injection mouse lymphoid tissue was collected.

### 28 DNA immunizations

30 MeMo®' mice were immunized by DNA tattooing using a micropigmentation device. DNA tattoo immunizations were performed with 20 µg plasmid DNA encoding the target antigen.. Mice were immunized with DNA encoding the human target PD-L1. For PD-L1

immunizations, Treg cells were depleted four days prior to the start of immunization by injection of mice with 0.5 mg anti-CD25 antibody PC61.5 to break tolerance. Mice were immunized at day 0, 3, 6, 14, 17, 28 and 31. Mouse serum was collected at day 35 to determine serum titers. Mice with low serum reactivity against the human and/or macaque target received additional 5 cycles of booster immunizations with human, rat or macaque DNA antigen and serum analyses. Each cycle consisted of two weekly DNA immunizations followed one week later by serum collection for titer analysis. Mice showing serum reactivity against cells expressing the human and macaque target received a final boost immunization followed after 3 days by collection of lymphoid tissue.

10 Recovery of lymphoid tissue

Spleen and draining lymph nodes were removed from all mice that were successfully immunized. Single cell suspensions were generated from both spleen and inguinal lymph nodes and subsequently these tissues were lysed in Trizol LS reagent and stored at -80°C until use.

15 Generation of 'immune' phage antibody repertoires by RT-PCR cloning of VH genes From successfully immunized mice, the inguinal lymph nodes were used for the construction of 'immune' phage antibody repertoires. To this end, RNA was extracted from Trizol LS lysed lymphoid tissue and 1µg of total RNA was used in a RT reaction using an IgG-CH1 specific primer. The resulting cDNA was then used to amplify the polyclonal pool of VH-encoding cDNA using in-house developed VH-specific primers essentially as described in Marks et al. (J Mol 20 Biol. 1991 Dec 5;222(3):581-97). The resulting PCR product was then cloned in a phagemid vector for the display of Fab fragments on phage, as described in de Haard et al. (J Biol Chem. 1999 Jun 25;274(26):18218-30) with the exception that the light chain was the same for every antibody and was encoded by the vector. After ligation, the phagemids were used to transform E.coli TG1 bacteria and transformed bacteria were plated onto LB-agar105 plates containing 25 ampicillin and glucose. All phage libraries contained  $>10^6$  transformants and had an insert frequency of  $> 80\%$ . Bacteria were harvested after overnight growth and used to prepare phage according to established protocols (de Haard et al., J Biol Chem. 1999 Jun 25;274(26):18218-30).

30 Targeting antibodies

EGFR- and PD-L1 cLC antibodies were obtained using previously described methods from phage antibody repertoires generated from successfully target-immunised MeMo® mice. Further, methods to generate antibody variable domain VH chains for the EGFR antibodies, including synthetic human anti-EGFR antibodies, have also been described in pending applications that are incorporated herein by reference: WO 2015/130173 A1; WO 2015/130172

35 A1.

Immunization of Memo® mice with CDR3

For generation of human antibodies binding to CD3, mice transgenic for the human common light chain and for a human heavy chain (HC) minilocus (comprising a selection of human V gene segments, all human Ds and all human Js) (see WO2009/157771 incorporated

herein by reference) were immunized with TCR/CD3 containing lipoparticles (Intergal Molecular). These mice are referred to as 'MeMo®' mice. For specific heavy chain variable regions, or trivalent multimers having the sequences disclosed herein, they can be produced by any means known to persons of ordinary skill in the art.

5

MeMo® mice were immunized with Hek293T-derived human 5D5M TCR/CD3 containing lipoparticles, followed by human T-cells for the generation of an anti-TCR/CDR3 immune response and anti-TCR/CD3 antibody panel generation.

10

Lipoparticles concentrate conformationally intact membrane proteins directly from the cell surface, permitting these complex proteins to be manipulated as soluble, high-concentration proteins for antibody immunization and screening

15

The lipoparticles used in the present study for immunisation contain the 5D5M TCR $\alpha\beta$  combination. The amino acid sequences (SEQ ID NO: 289 and SEQ ID: 290)

Hek293T-derived TCR/CD3 lipoparticles of the 5D5M TCR $\alpha\beta$  combination were synthesized, cloned and used to generate lipoparticles containing this TCR/CD3 combination by transient transfection into HEK293T cells (Intergal Molecular).

20

5D5M TCR $\alpha$  (SEQ ID NO: 289)

MWGVFLYVSMKMGTTGQNIQPTEMTATEGAIVQINCTYQTSGFNGLFWYQQHA  
GEAPTFLSYNVLGEEKGRFSSFLSRSKGYSYLLKELQMKDSASYLCAVMDSNYQLIWGAG  
TKLIIPDIQNPDPAVYQLRDSKSSDKSVCLTFDSQTNVSQSKDSDVYITDKTVLDMRSMDF  
KSNSAVAWSNKSDFACANAFNNSIIPEDTFFPSPESSCDVKLVEKSFETDTNLNFQNLSVIGFRI  
25 LLLKVAGFNLLMTLRLWSS

5D5M TCR $\beta$  (SEQ ID NO: 290)

MRIRLLCCVAFSLLWAGPVIAGITQAPTSQILAAGRRTLRCTQDMRHNAMEWYRQDL  
GLGLRLIHYSNTAGTTGKGEVPDGYSVRANTDDFPLTLASAVPSQTSVYFCASSEAGGNTE  
30 LFFGEGSRLTVLEDLNKVFPPPEVAVFEPSEAEISHTQKATLVCLATGFFPDHVELWWVNGKEV  
HSGVSTDPPQPLKEQPALNDSRYCLSSRLRVSATFWQNPRNHFRCQVQFYGLSENDEWTQDR  
AKPVTQIVSAEAWGRADCGFTSVSYQQGVLSATILYEILLGKATLYAVLVSALVLMAMVKRKDF

35

MeMo® mice were used for immunizations using TCR/CD3 lipoparticles and primary human T cells

The immunization schedule contains points on day 35, 56, 77 and 98, where the antigen-specific Ig serum titer was determined by ELISA using QTG-derived 3SDX TCR/CD3 positive and -negative lipoparticles using anti mouse IgG detection and by ELISA using

CD3c5E-Fc fusion protein as a positive control. The reactivity was observed in sera drawn at day 35 will determine which mice developed a relevant anti-TCR/CD3 response.

For all immunized mice, lymphoid material for antibody discovery was collected and stored when:

5 Titors are 1/300 for human TCR/CD3 (in ELISA using lipoparticles), or:

Titors are <1/300 and >1/100 for human TCR/CD3 and did not increase during the last booster immunization.

Priming immunisation using lipoparticles

10 To prime the humoral immune response in the MeMo® mice for TCR/CD3, lipoparticles containing the human 5D5M TCR $\alpha\beta$  combination was used for immunization. Lipoparticles were used together with Gerbu adjuvant for the first and second injection.

Booster immunizations using polyclonal T-cells

15 Mice were immunised by sub-cutaneous injection of cell suspension. The first booster immunisations (day 28) comprised a mix of cells in PBS with adjuvant and all subsequent injections are only composed of cells in PBS. Mice that have developed at day 35 serum IgG titers of 1/300 against human TCR/CD3 (determined by ELISA using lipoparticles) received additional injections with cells on days 42, 43 and 44. Mice that failed to meet these criteria  
20 receive booster immunisations (day 42 and 49) with cells. All subsequent immunisations are given as sub-cutaneous injections of cells in PBS. After the final immunisation, mice are sacrificed, bled for serum and the spleen and left inguinal lymph nodes are collected.

Screening sera from immunised mice in ELISA

25 Interim serum IgG titers were screened by ELISA using TCR/CD3-containing lipoparticles and 'null' lipoparticles. Serum IgG titers were determined using anti-mouse IgG staining, as this staining was shown to be the most sensitive.

CD3 binding variable domains were made using the amino acid sequence of the heavy chain variable region of a CD3 MF of the CDR regions thereof as indicated in SEQ ID NO: 92-

30 154.

Re-cloning of VH-encoding cDNA's from the phagemid vector to IgG-expression vectors

The VH-encoding cDNA's of all target-specific clones were sequenced. A selection of unique clones based on sequence identity and cluster analysis was then re-cloned to different  
35 IgG expression vectors using Sfi1-BstEII or a Sfi1/Xhol digestion and ligation of the pool of digested cDNA's into the IgG expression plasmid was done according to standardised molecular biological techniques.

Purification of antibodies from culture supernatant

Medium containing antibodies is harvested and centrifuged to remove the cell debris. Subsequently Protein A Sepharose beads are added to the medium. Medium and Protein A Sepharose beads are incubated with the antibodies to allow binding.

5 After incubation the beads are isolated from the medium and washed, by a vacuum filter. The antibodies are eluted from the beads by incubation with elution buffer.

Optionally, the buffer of the purified IgG is exchanged/desalted.

#### Buffer exchange

10 In order to desalt the purified antibodies the antibody fraction is centrifuged using a filter plate or filter column. The plate or column is centrifuged to reduce the volume of the antibody fraction. Subsequently, PBS or the required buffer is added to the fraction to replace the buffer with a low salt buffer. Optionally this centrifugation step followed by adding buffer is repeated in order to further desalt the storage buffer of the antibodies.

15

#### **Example 9: Generation of trispecific antibodies with a tumor cell antigen on the short or long arm.**

Trispecific antibodies were generated by transient co-transfection of two plasmids encoding IgG with different VH domains, using CH3 engineering technology for efficient hetero-20 dimerisation and formation of trispecific antibodies. The common light chain is also co-transfected in the same cell, either on the same plasmid or on another plasmid. In our co-pending applications (e.g. WO2013/157954 and WO2013/157953; incorporated herein by reference) we have disclosed methods and means for producing multispecific antibodies from a single cell, whereby means are provided that favor the formation of multispecific antibodies over 25 the formation of monospecific antibodies. These methods can also be favorably employed in the present invention for the generation of multivalent multimers, including trispecific antibodies.

Specifically, preferred variations to predominantly produce trispecific full length IgG molecules are amino acid substitutions in reference to a human wild type sequence at positions 351 and 366, e.g. L351K and T366K (numbering according to EU numbering) in the first CH3 30 domain (the 'KK-variant' heavy chain) and amino acid substitutions at positions 351 and 368, e.g. L351D and L368E in the second CH3 domain (the 'DE-variant' heavy chain), or vice versa. It was previously demonstrated in our co-pending applications that the negatively charged DE-variant heavy chain and positively charged KK- variant heavy chain preferentially pair to form heterodimers (so-called 'DEKK' molecules). Homodimerization of DE-variant heavy chains (DE-35 DE homodimers) or KK-variant heavy chains (KK-KK homodimers) hardly occurs due to strong repulsion between the charged residues in the CH3-CH3 interface between identical heavy chains.

According to the present invention, the immune cell engaging binding domain or tumor antigen binding domain can be placed at any position on the multivalent molecule, including the

distal or interior position of the long arm or the short arm, and the heterodimerization technology can be utilized to favorably generate the trispecific molecule over the monospecific, bivalent homodimer, or quadrospecific homodimer.

First, it was demonstrated that a tumor cell antigen binding domain may be placed at  
5 either the distal or interior region of the long arm or the short arm.

For each of the trispecific and/or trivalent antibodies described herein, expression is  
accomplished through suspension growth adapted 293 cells were cultivated in T125 flasks on a  
shaker plateau until a density of  $3.0 \times 10^6$  cells/ml. Cells were seeded at a density of  $0.3-0.5 \times$   
10 $^6$  viable cells/ml in each well of a 24-deep well plate. The cells were transiently transfected  
with a mix of two plasmids encoding different antibodies, cloned into the proprietary vector  
system. Seven days after transfection, the cellular supernatant was harvested and filtered  
through a 0.22  $\mu$ M filter. The sterile supernatant was stored at 4°C until purification of the  
trispecific antibodies.

For an example of a trispecific molecule having an immune engaging binding domain at  
15 the interior position of the long arm and a tumor cell antigen at the short arm, DNA encoding the  
VH gene for the CD3 binding domain (MF8078), a linker of the invention and a tetanus toxoid  
(TT) binding domain (MF1337) are cloned into a vector encoding the positively charged CH3  
domain (KK), where DNA encoding the VH gene for the EGFR binding domain (MF8233) is  
cloned into a vector encoding the negatively charged CH3 domain (DE) encoding a trispecific  
20 molecule of EGFR x CD3:TT. The heavy chain variable regions for the three binding domains  
are set out in Table 13, with activity for these trispecific molecules described in Figs. 15a and  
16a. For these trispecific molecules, each heavy chain variable region pairs with a common light  
chain. SEQ ID NO: 29.

Table 13.

EGFR x CD3 : Mock					
DE arm	KK arm	Linker used	Linker SEQ ID	Purpose	
MF8233	MF8078	IgG1 G4S	12	MF1337	sample
MF8233	MF8078	IgG1 H	19	MF1337	sample
MF8233	MF8078	IgG2A MH	4	MF1337	Sample
MF8233	MF8078	IgG1 MH	11	MF1337	sample
MF8233	MF8078	IgG1 R	23	MF1337	sample
MF8233	MF8078	IgG1 UH	2	MF1337	sample
MF8233	MF8078	IgG1 UL	14	MF1337	sample
MF8233	MF8078	IgG2A H	15	MF1337	sample
MF8233	MF8078	IgG2A R	20	MF1337	sample
MF8233	MF8078	IgG2B H	16	MF1337	sample
MF8233	MF8078	IgG2B R	21	MF1337	sample
MF1025	MF8078	IgG1 H*	19	MF1337	Neg Contr
MF1025	MF8078	IgG2 AMH	4	MF1337	Neg Contr

25 For an example of a trispecific antibody having an immune engaging binding domain at  
the interior position and a tumor cell antigen at the distal position of the long arm, DNA encoding

the VH gene for the CD3 binding domain (MF8078), a linker of the invention and DNA encoding the VH gene for the EGFR binding domain (MF8233) are cloned into a vector encoding the positively charged CH3 domain (KK), where DNA encoding the VH gene for the Thyroglobulin binding domain ("Thyro") (MF1025) is cloned into a vector encoding the negatively charged CH3 domain (DE) encoding a trispecific molecule of Thyro x CD3:EGFR. Fig. 14b. The heavy chain variable regions for the three binding domains are set out in Table 14, with activity for these trispecific molecules described in Figs. 15b and 16b. For these trispecific molecules, each heavy chain variable region pairs with a common light chain. SEQ ID NO: 29.

10

Table 14.

Mock x CD3 : EGFR					
DE arm	KK arm	Linker used	Linker SEQ ID		Purpose
MF1025	MF8078	IgG1 G4S	12	MF8233	sample
MF1025	MF8078	IgG1 H	19	MF8233	sample
MF1025	MF8078	IgG2A MH	4	MF8233	sample
MF1025	MF8078	IgG1 MH	11	MF8233	sample
MF1025	MF8078	IgG1 R	23	MF8233	sample
MF1025	MF8078	IgG1 UH	2	MF8233	sample
MF1025	MF8078	IgG1 UL	14	MF8233	sample
MF1025	MF8078	IgG2A H	15	MF8233	sample
MF1025	MF8078	IgG2A R	20	MF8233	sample
MF1025	MF8078	IgG2B H	16	MF8233	sample
MF1025	MF8078	IgG2B R	21	MF1337	Comparator type A
MF1025	MF8078	IgG1 H	19	MF1337	Comparator type A
MF1025	MF1122	IgG2 AMH	4	MF1337	Neg contr
MF1025	MF8078	IgG1 H	19	MF1337	Neg contr
MF1025	MF1122	IgG2 AMH	4	MF1337	Neg contr

15 **Example 10: Effect of positioning the tumor cell antigen binding domain on short or long arm for T-cell engaging trivalent molecules.**

#### Cell lines

BxPC3 is a human pancreatic cancer cell line.

HCT-116 is a human colon carcinoma cell line.

20

The above series of trispecific IgGs were generated at 24 well production which incorporated eleven different linkers in trispecifics containing an anti-EGFR binding domain on the short arm (Fig. 14a and Table 13) and an anti-EGFR binding domain on the long arm (Fig.

14b and Table 14) and an array of control antibodies. These molecules have been assessed for their capacity to cause T-cell activation and in a cytotoxicity assay.

Using Ficoll and EasySep human T cell isolation kit according to standard techniques resting T cells were isolated from whole blood from healthy donors, checked for > 95% T cell 5 purity by anti-CD3 antibody using flow cytometric analysis and subsequently cryopreserved. For a cytotoxicity assay the cryopreserved T cells were thawed and used if their viability was > 90% upon thawing, determined by standard Trypan Blue staining. Cytotoxicity assay in short, thawed resting T cells and BxPC3 (Fig 15) or HCT116 (Fig 16 and 17) target cells were co-cultured in an E:T ratio of 5:1 for 48 hours. For the trivalent antibodies, a 6-step 3 fold dilution 10 series starting at a concentration of 4 ug/ml was used. EGFRxCD3 bispecific antibodies were used as a positive control; MockxMock:Mock, MockxCD3:Mock and EGFRxMock:Mock trivalent antibodies were used for specificity controls. T cell activation was quantified using flow cytometry; CD4 and CD8 T cells were gated based on CD4 and CD8 expression and subsequently analyzed for their activation status by measuring CD25 and CD69 expression on 15 T cells. Target cell lysis was determined by measuring the fraction of alive cells by measuring ATP levels assessed by CellTiterGlo (Promega). ATP levels, measured by luminescence on an Envision Microplate reader results in Relative light unit (RLU) values, which were analyzed using GraphPad Prism.

Target cell lysis for each sample was calculated as follows:

20 %Killing = (100- (RLU sample/RLU no IgG) x 100).

These data pertaining to T-cell activation (see Fig. 15) and cytotoxicity (Fig. 16) demonstrate that the trispecific antibodies are functional. As shown in Figures 15 and 16, MockxCD3:EGFR and EGFRxCD3:Mock trispecific molecules are demonstrated to be capable 25 of inducing EGFR target specific T cell activation and cytotoxicity. When the anti-EGFR Fab was positioned on the distal position in the long arm the MockxCD3:EGFR trispecific showed enhanced activity over both the bispecific EGFRxCD3 and the trispecific having the EGFR binding domain on the short arm and the CD3 and mock TT (MF1337) binding domain on the long arm (see Fig. 14a), both in terms of T cell activation (Fig.15) and cytotoxicity (Fig. 16).

Further, based on the activity of the trispecific antibodies having EGFR and CD3 on the 30 long arm, the linkers can be binned into those that correlate with relatively high cytokine production (IgG1 UH (SEQ ID NO: 2), IgG1 MH (SEQ ID NO: 11), IgG2A MH (SEQ ID NO: 4) and IgG1 G4S (SEQ ID NO: 12)) and those that correlate with relatively low cytokine production (IgG1 UL (SEQ ID NO: 14), IgG2A H (SEQ ID NO: 15), IgG2B R (SEQ ID NO: 21), IgG2A R (SEQ ID NO: 20), IgG1 H (SEQ ID NO: 19), IgG1R (SEQ ID NO: 23)) as show in Figs. 35 17b-d. The change in linker use and impact on cytotoxicity was less pronounced. Fig. 17a.

**Example 11: Generation of trispecific antibodies with an immune cell engaging binding domain on the short or long arm.**

According to the present invention, the immune cell engaging binding domain can be placed at any position on the multivalent molecule, including the distal or interior position of the long arm or the short arm, and the heterodimerization technology can be utilized to favorably generate the trivalent molecule. See Figs. 18 (CD3 binding domain on the short arm), Fig. 19  
5 (CD3 binding domain on the interior long arm) and Fig. 25 (CD3 binding domain on the distal long arm).

For example, an immune engaging domain is positioned at the short arm, where DNA encoding the VH gene for the CD3 binding domain (MF8078) is cloned into a vector encoding the positively charged CH3 domain (KK), where DNA encoding the VH gene for the EGFR  
10 binding domain (MF9988 (SEQ ID NO:218) or MF9891 (SEQ ID NO:191)), the linker IgG2A MH and the VH gene for the PD-L1 binding domain (MF5380 (SEQ ID NO:173) or MF5444 (SEQ ID NO:164)) is cloned into a vector encoding the negatively charged CH3 domain (DE). Fig. 18 (CD3xPD-L1:EGFR).

15 Table 15

CD3xPD-L1:EGFR		
KK arm	DE arm	
MF8078	MF5380	MF9988
MF8078	MF5380	MF9891
MF8078	MF5444	MF9988
MF8078	MF5444	MF9891

Alternatively, an immune engaging domain is positioned at the interior position of the long arm, where DNA encoding the VH gene for a CD3 binding domain (MF8078), the linker IgG2A MH (SEQ ID NO:4) and a VH gene for a PD-L1 binding domain (MF5444 (SEQ ID NO:164), MF5380 (SEQ ID NO:173), MF5377 (SEQ ID NO:155)) is cloned into a vector  
20 encoding the positively charged CH3 domain (KK), where DNA encoding the VH gene for a EGFR binding domain (MF9886 (SEQ ID NO:200), MF9988 (SEQ ID NO:218), MF9891 (SEQ ID NO:191) or MF9873 (SEQ ID NO:209)) is cloned into a vector encoding the negatively charged CH3 domain (DE). Fig. 19 (EGFRxCD3:PD-L1).

25

Table 16

EGFRxCD3:PD-L1		
DE arm	KK arm	
MF9988	MF8078	MF5444
MF9988	MF8078	MF5380

MF9886	MF8078	MF5380
MF9988	MF8078	MF5377
MF9886	MF8078	MF5377
MF9891	MF8078	MF5377
MF9873	MF8078	MF5377
MF9891	MF8078	MF5380
MF9873	MF8078	MF5380
MF9891	MF8078	MF5444
MF9873	MF8078	MF5444
MF1337	MF8078	MF5377
MF1337	MF8078	MF5380
MF1337	MF8078	MF5444
MF9886	MF8078	MF1337
MF9988	MF8078	MF1337
MF9891	MF8078	MF1337
MF9873	MF8078	MF1337
Control Antibodies		
MF1337	MF8078	-
MF8233	MF8078	-

Alternatively, an immune engaging domain is positioned at the distal position of the long arm, where DNA encoding the VH gene for a CD3 binding domain (MF8078 (SEQ ID NO:110), MF8508 (SEQ ID NO: 128) or MF8057 (SEQ ID NO: 92)) a linker IgG 1H (SEQ ID NO:19) and a VH gene for a Fibrinogen binding domain ("Fibri") (MF1025) is cloned into a vector encoding the positively charged CH3 domain (KK), where DNA encoding the VH gene for a EGFR binding domain (MF8233) is cloned into a vector encoding the negatively charged CH3 domain (DE).

5 Fig. 24 (EGFRxFibri:CD3).

Table 17

EGFRxFibri:CD3				
DE arm	KK arm			
		Linker used	Linker seq	
MF8233	MF1122	IgG1H	19	MF8078
MF8233	MF1122	IgG1H	19	MF8508
MF8233	MF1122	IgG1H	19	MF8057

For each of the trispecific molecules described above, and set out in Figures 18, 19 and 24, each heavy chain variable region pairs with a common light chain. SEQ ID NO: 29.

5 As described further below, each CD3 binding domain placement was demonstrated to be effective at generating T-cell cytotoxicity or activation against cells expressing one or more extracellularly exposed tumor cell antigens.

10 **Example 12: Effective dual tumor antigen binding and T cell engagement via CD3 for the trispecific format of EGFRxCD3:PD-L1.**

Cell lines:

MDA-MB-231 cells (ATCC® HTB-26) are breast cancer cells; derived from metastatic site.

15 Trispecific antibodies were produced according to the format at Figure 19 to analyze the capacity of such molecules to achieve simultaneous tumor antigen targeting and T-cell engagement via cytotoxicity. These antibodies were generated by techniques described above.

20 Four anti-EGFR Fabs (MF9886, MF9988, MF9891, MF9873) with a range of affinities from relatively low to high were used for the short arm and were combined with different anti-PD-L1 Fabs (MF5444, MF5380 and MF5377) that also contain a range of affinities from low to high for the distal long arm. The anti-CD3 Fab and linker were kept constant, using MF8078, and linker IgG2 AMH (SEQ ID NO: 4).

25 Ranking for EGFR and PD-L1 affinities was based on binding data from Table 18 and 19, respectively; the ranking was based on binding relative to a reference anti-EGFR and anti-PD-L1 antibody as described below.

Table 18 EGFR panel Monospecific, Bivalent EGFR antibodies having the heavy chains MF9886, MF9988, MF9891, MF9873.

MF	MDA-MB-231 cells (%)
9886	10.7
9988	21.0
9891	44.2
9873	82.5
8233	100

The affinity ranking of EGFR heavy chains is based on the relative ability of these monospecific, bivalent antibodies to bind EGFR expressing cells as set out above as compared to the positive control of a monospecific, bivalent antibody having heavy chain MF8233.

5

Table 19. PD-L1 panel Monospecific, Bivalent PD-L1 antibodies, having the heavy chains MF5444, MF5380 and MF5377

Anti-PD-L1 heavy chain	Binds to huPD-L1	Ratio EC50/EC50 RG7446
MF5377	Yes	3.09
MF5380	Yes	3.67
MF5444	Yes	5.46

10

The relative affinity ranking of the PD-L1 heavy chain arms is based on capacity to bind human PD-L1 in ELISA. To this end, ELISA plates were coated with huPD-L1-His (Sinobiological) in an 8-step, 3-fold titration dilution range, starting at 10 µg/ml. Subsequently, binding of each PD-L1 Fab was evaluated as a PD-L1xTT IgG 5 µg/ml. EC50s for binding were determined, and normalized to the binding EC50 as determined for anti-PD-L1 RG7446 MPDL3280A, see US 2010/0203056, present on each ELISA plate.

15

These trispecific EGFRxCD3:PD-L1 molecules were then tested for their capacity to induce cytotoxicity based on methods previously described herein against two cell lines (HCT116 and MDA-MB-231) having different antigen densities for the tumor cell antigens. The expression profiles of these cell lines were determined using FACS staining by use of control antibodies (cetuximab for EGFR, and MPDL3280A for PD-L1) were considered positive for expression of the antigen of interest when the mean fluorescence intensity (MFI) was 3x higher than the background signal. Triplicates were performed for PD-L1 and quadruplicates for EGFR, and results were reported as MFI as set out below.

Table 20

Cell line	EGFR (MFI)	PD-L1 (MFI)
HCT116	178,523	10,876
MDA-MB-231	276,915	74,581

This study examining cytotoxicity of the trispecific molecules against these cell lines show all three binding domains of the EGFRxCD3:PD-L1 trispecific molecules are capable of simultaneous binding the two tumor cell antigens and CD3, such that both tumor antigen binding domains of the trispecific molecule contribute to cytotoxicity upon T-cell engagement. Figs. 20  
5 and 21 (against MDA-MB-231 cells) and Fig. 22 (against HCT116 cells). These trispecific molecules generally showed enhanced functional activity over the bispecific EGFRxCD3xMock and the MockxCD3xPD-L1 controls, with the trispecific molecule 9873x8078: 5377 showing the largest percent lysis. Fig. 21. The trispecifics tested here are more potent against MDA-MB-231  
10 cells which have relatively high target antigen levels compared to HCT116 cells. Fig. 21 and Fig.  
22.

**Example 13: Effective dual tumor antigen binding and T cell engagement via CD3 for trispecific format of CD3xPD-L1:EGFR.**

Trispecific antibodies were produced according to the format at Figure 18 to further  
15 show simultaneous tumor antigen targeting where the immune engaging domain is present on the short arm. These antibodies were generated by techniques described above. For this format, dual antigen targeting-correlated binding was demonstrated, such that with increasing PD-L1 affinity there was a continued enhancement of target cell binding as measured by FACs on MDA-MB-231 cells as observed for CD3xPD-L1:EGFR molecules. Fig. 23a. For certain of  
20 these trispecific molecules having the format of CD3xPD-L1:EGFR, simultaneous dual antigen binding and immune cell engagement have been demonstrated to have an additive effect on cytotoxicity of BxPC3 cells over molecules binding a single antigen and CD3 (heavy chain sequences not shown) verifying the capability of these molecules to engage all three binding arms simultaneously. Fig. 23b. The protocol for the cytotoxicity assay for these data have been  
25 described above.

**Example 14: Effective T cell activation via CD3 at the distal long arm, for trispecific format of EGFRxFibrinogen:CD3.**

30 Jurkat-NFAT-RE-luc2 cells (Promega) are a genetically engineered Jurkat T cell line that expresses a luciferase reporter driven by an NFAT-response element (NFAT-RE). HT29 (ATCC HTB-38) is a human colon cancer cell.

A study was performed for placing the CD3 binding domain on the distal region of the  
35 long arm, with a trivalent molecule of EGFR (MF8233)xFibrinogen(MF1122):CD3(MF8078). Fig. 24. Performing a T-cell activation assay on Jurkat-NFAT-RE-luc2 cells against target cell HT29 to establish functional T-cell activation capacity for this format. The reporter assay in short: Jurkat effector T cells were co-incubated with target cells in the presence of a concentration range of trispecific antibodies and control antibodies. After 5 hours of incubation the Luciferase

activity of the reporter cells was measured as a read-out for T cell activation, using the Bio-Glo Luciferase Assay System (Promega). Luminescence activity was measured on an Envision Microplate reader resulting in Relative light unit (RLU) values, which were analyzed using GraphPad Prism.

5 As shown in Figure 25, activation of T-cells by the EGFRxFibri:CD3 trispecific molecules was demonstrated at levels equal to or greater than the positive control, which is a EGFRxCD3 antibody previously demonstrated to engender T-cell activation in Fig. 15a.

**Example 15: Effective Tumor Antigen Binding and T cell Engagement via CD3 for an**

**10 Array of CD3 Binding Domains and Linkers**

A panel of EGFRxCD3:EGFR bispecific, trivalent molecules were generated (Fig. 26) to demonstrate the efficacy of tumor targeting and T-cell engagement across a variety of different CD3, immune-cell engaging binding domains and eight different linkers. Each trivalent molecule contained two of the same anti-EGFR binding domains (MF9891) at the short arm and distal 15 long arm position, with the CD3 binding domain at the interior, long arm position. For this study, a reporter cell line of Jurkat-NFAT-RE-luc2 cells and target cells HCT116 (intermediate EGFR expression) and MDA-MB-231 were used to measure T-cell activation, by methods previously described. For a negative control, trivalent molecules using a CD3 binding domain from different superclusters were produced MockxCD3xMock or the bispecific (EGFR (MF8233) 20 xTT(MF1337)) (4,000 ng/ml). The read-out relied on reporter activation after 5 hours of incubation.

Each CD3 tested from varying superclusters demonstrated reporter activity using HCT116 cells, with a CD3 binding domain from supercluster 7 demonstrating the lowest relative reporter activity, while evidencing a spectrum of activity based on linker arms. In contrast, the 25 two CD3 binding domains from supercluster 1 (MF8058 and MF8078) and a CD3 binding domain from supercluster 4 (MF8508) demonstrated relatively consistent activity irrespective of linker arm. Finally, one CD3 binding domain of supercluster 1 (MF8057) evidenced relatively low reporter activity, which provided some differentiation associated with different linkers. See Fig. 27. A review of these data indicate that trivalents containing the IgG1 MH linker appear 30 consistently to be the most potent across superclusters.

Similarly, each CD3 binding domain tested from varying superclusters demonstrated reporter activity using MDA-MB-231 cells, with the CD3 binding domains from supercluster 7 demonstrating relatively low reporter activity, with a spectrum of activity based on the linker used. In contrast, the three CD3 binding domains from supercluster 1 (MF8057, MF8058 and 35 MF8078) and CD3 binding domain from supercluster 4, demonstrated relatively similar activity irrespective of linker used. See Fig. 28.

---

Table 21

IgG heavy chains for the generation of bispecific molecules. Table 21A: CH1 region. Table 21B: hinge region. Table 21C: CH2 region. Table 21D: CH2 containing L235G and G236R silencing substitutions. Table 21E: CH3 domain containing substitutions L351K and T366K (KK). Table 21F: CH3 domain containing substitutions L351D and L368E (DE).

Table 21A

CH1:

gctagcaccaagggcccacggcttccccctggcacccctccaaagagcacccctgg

10 A S T K G P S V F P L A P S S K S T S G

ggcacagcggccctggctggcgtcaaggactacttccccgaaccgggtacgggtgtcg

G T A A L G C L V K D Y F P E P V T V S

tggaaactcaggcgccctgaccagcggcgtgacaccccccgtacagtcctca

W N S G A L T S G V H T F P A V L Q S S

15 ggactctactccctcagcagcgtcgaccgtgcctccacgcagctggcaccaggacc

G L Y S L S S V V T V P S S S L G T Q T

Tcacatctgcaacgtgaatcacaagcccagcaacaccaagggtggacaagagagt (SEQ ID NO: 291)

Y I C N V N H K P S N T K V D K R V (SEQ ID NO: 292)

20 Table 21B

Hinge:

Gagccaaatcttgtgacaaaactcacacatgcccaccgtgccc (SEQ ID NO: 293)

E P K S C D K T H T C P P C P (SEQ ID NO: 294)

25

Table 21C

CH2:

gcacctgaactccgggggaccgtcagtcttccttcccccaaaacccaaggacacc

A P E L L G G P S V F L F P P K P K D T

30 ctcatgtctccggaccctgagggtcacatgcgtgggtggacgtgagccacgaagac

L M I S R T P E V T C V V V D V S H E D

cctgaggtaacttcaactggtaactggacggcgtggagggtgcataatgcaagacaag

P E V K F N W Y V D G V E V H N A K T K

ccgcgggaggaggcgtacaacacgacacgttccgtggcgtacgttccgttccgttccgt

35 P R E E Q Y N S T Y R V V S V L T V L H

caggactggctgaatggcaaggaggtaactggtaactggtaagggtctccaaacaagccctcccagcc

Q D W L N G K E Y K C K V S N K A L P A

Cccatcgagaaaaccatctccaaagccaaa (SEQ ID NO: 295)

P I E K T I S K A K (SEQ ID NO: 296)

40

Table 21D

CH2 containing L235G and G236R silencing substitutions:

gcacctgaactcggcaggggaccgtcagtcttccttcccccaaaacccaaggacacc

A P E L G R G P S V F L F P P K P K D T

45 ctcatgtctccggaccctgagggtcacatgcgtgggtggacgtgagccacgaagac

L M I S R T P E V T C V V V D V S H E D

cctgaggtaacttcaactggtaactggacggcgtggagggtgcataatgcaagacaag

P E V K F N W Y V D G V E V H N A K T K

ccgcgggaggaggcgtacaacacgacacgttccgtggcgtacgttccgttccgttccgt

P R E E Q Y N S T Y R V V S V L T V L H  
 caggactggctgaatggcaaggagttacaagtgcacaggctccaaacaaggcccccagcc  
 Q D W L N G K E Y K C K V S N K A L P A  
 Cccatcgagaaaaccatctccaaagccaaa  
 5 P I E K T I S K A K (SEQ ID NO: 297)  
 (SEQ ID NO: 298)

## Table 21E

CH3: KK of DEKK

10 gggcagccccgagaaccacagggttacaccaaggcccccattccggggaggagatgaccaag  
 G Q P R E P Q V Y T K P P S R E E M T K  
 aaccagggtcagcctgaagtgcctggtaaaggcttcatcccagcgacatgcctggag  
 N Q V S L K C L V K G F Y P S D I A V E  
 tgggagagcaatggcagccggagaacaactacaagaccacgcctccgtgctggactcc  
 15 W E S N G Q P E N N Y K T T P P V L D S  
 gacggctcccttccttatagcaagctaccgtggacaagagcagggtggcagcaggggg  
 D G S F F L Y S K L T V D K S R W Q Q G  
 aacgttctcatgtccgtatgcacggctctgcacaaccactacacgcagaagagc  
 N V F S C S V M H E A L H N H Y T Q K S  
 20 ctctccctgtctccgggttga (SEQ ID NO: 299)  
 L S L S P G - (SEQ ID NO: 300)

## Table 21F

CH3: DE of DEKK

25 gggcagccccgagaaccacagggttacaccaaggcccccattccggggaggagatgaccaag  
 G Q P R E P Q V Y T D P P S R E E M T K  
 aaccagggtcagcctgcgcaggtaaaggcttcatcccagcgacatgcctggag  
 N Q V S L T C E V K G F Y P S D I A V E  
 tgggagagcaatggcagccggagaacaactacaagaccacgcctccgtgctggactcc  
 30 W E S N G Q P E N N Y K T T P P V L D S  
 gacggctcccttccttatagcaagctaccgtggacaagagcagggtggcagcaggggg  
 D G S F F L Y S K L T V D K S R W Q Q G  
 aacgttctcatgtccgtatgcacggctctgcacaaccactacacgcagaagagc  
 N V F S C S V M H E A L H N H Y T Q K S  
 35 ctctccctgtctccgggttga (SEQ ID NO: 301)  
 L S L S P G - (SEQ ID NO: 302)

**CLAIMS**

1. A multivalent antibody which comprises:

5

- a base antibody portion which comprises two binding domains; and
- at least one additional binding domain,

10 wherein the base antibody portion is connected by a linker to the at least one additional binding domain,

wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region, and

15 wherein the linker comprises a hinge sequence or a sequence derived from a hinge sequence.

2. A multivalent antibody according to claim 1, wherein at least one additional binding domain comprises a Fv domain, a Fab domain or a modified Fab domain.

20

3. A multivalent antibody according to claim 1, wherein at least one additional binding domain comprises a CH1 region and is connected to the base antibody portion by said linker, linking a variable region of the base antibody portion and the CH1 region .

25 4. A multivalent antibody according to any one of the preceding claims, wherein the common variable region is in the form of a common light chain comprising VL-CL.

5. A multivalent antibody according to claim 4, wherein the common light chain comprises the sequence of SEQ ID NO: 29.

30

6. A multivalent antibody according to any one of claims 1 to 3, wherein the common variable region is in the form of a common heavy chain comprising VH-CH1.

35 7. A multivalent antibody according to any one of the preceding claims, wherein each binding domain of the base antibody portion binds a different epitope.

8. A multivalent antibody according to any one of the preceding claims which binds at least three different epitopes.

40 9. A multivalent antibody which comprises:

- a base antibody portion which comprises two binding domains; and
- at least one additional binding domain,

5 wherein at least one additional binding domain comprises a CH1 region and is connected to the base antibody portion by said linker, linking a variable region of the base antibody portion and the CH1 region, and

wherein the multivalent antibody binds to at least three different epitopes.

10

10. A multivalent antibody according to claim 9, wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region

15 11. A multivalent antibody according to claim 9 or 10, wherein at least one additional binding domain comprises a Fv domain, a Fab domain or a modified Fab domain.

12. A multivalent antibody according to claim 10 or 11, wherein the common variable region is in the form of a common light chain comprising VL-CL.

20

13. A multivalent antibody according to claim 12, wherein the common light chain comprises the sequence of SEQ ID NO: 29.

14. A multivalent antibody according to claim 10 or 11, wherein the common variable region 25 is in the form of a common heavy chain comprising VH-CH1.

15. A multivalent antibody according to any one of claims 10 to 14, wherein each binding domain of the base antibody portion binds a different epitope.

30 16. A multivalent antibody according to any one of the preceding claims, wherein the linker comprises a sequence as set out in any one of SEQ ID NOs:1 to 24 or a polypeptide having at least about 85% sequence identity to any one thereto.

17. A multivalent antibody according to any one of the preceding claims, wherein the base 35 antibody portion is a full length immunoglobulin.

18. A multivalent antibody according to any one of the preceding claims, wherein the binding domains of the base antibody portion are Fab domains.

19. A multivalent antibody according to any one of the preceding claims, wherein the base portion of the antibody comprises a first CH3 domains that dimerizes with a second CH3 domain, the first of which comprises an amino acid residue lysine at positions 351 and 366 or at positions corresponding thereto and the second of which comprises the amino acid residues of 5 aspartic acid at 351 and glutamic acid at 368 or at positions corresponding thereto.

20. A multivalent antibody according to any one of the preceding claims, wherein the common variable regions of the binding domains of said antibody are encoded by a nucleic acid that is obtained from, derived from or based on a nucleic acid encoded by a transgenic rodent 10 comprising a rearranged variable chain nucleic acid sequence in its germline.

21. A multivalent antibody according to any one of the preceding claims, wherein each Fab domain of the base antibody portion is connected via a linker to an additional binding domain.

15 22. A multivalent antibody according to any one of the preceding claims, wherein the N-terminal end of a binding domain of the base antibody portion is connected to the C-terminal end of one additional binding domain via a linker.

20 23. A multivalent antibody according to any one of the preceding claims, wherein an N-terminal end of the first binding domain of the base antibody portion is connected to a C-terminal end of a first additional binding domain via a linker and wherein an N-terminal end of the second binding domain of the base antibody portion is joined to a C-terminal end of a 25 second additional binding domain via a linker.

25 24. A multivalent antibody according to any one of claims 1 to 22, wherein an N-terminal end of the first binding domain of the base antibody portion is connected to a C-terminal end of a first additional binding domain via a linker and wherein a C-terminal end of a second additional binding domain is connected to an N-terminal end of the first additional binding domain via a 30 linker.

30 25. A multivalent antibody according to any one of the preceding claims which is capable of binding to at least four different epitopes.

35 26. A multivalent antibody according to any one of the preceding claims which is capable of binding to at least two different antigens.

27. A multivalent antibody according to any one of the preceding claims, wherein at least one additional binding domain is a Fab domain and said additional Fab domain comprises a CH1 is of a different subclass compared to the CH1 of the base antibody portion.

28. A multivalent antibody according to any one of the preceding claims, wherein the linker is short, long, charged, rigid or flexible.
- 5 29. A multivalent antibody according to any one of the preceding claims, wherein the amino acid sequence of the linker comprises a naturally-occurring sequence or comprises a sequence derived from a naturally-occurring sequence.
- 10 30. A multivalent antibody according to any one of the preceding claims, wherein the linker comprises a middle hinge region sequence.
31. A multivalent antibody according to any one of the preceding claims, wherein the linker comprises an upper and a lower hinge sequence.
- 15 32. A multivalent antibody according to any one of the preceding claims, wherein the linker comprises a helix-forming sequence.
33. A multivalent antibody according to any one of claims 1 to 27, wherein the linker comprises an amino acid sequence according to any one of SEQ ID NOS: 1 to 24.
- 20 34. A multivalent antibody according to any one of the preceding claims, wherein at least one additional binding domain comprises a CH1 domain.
35. A multivalent antibody according to any one of the preceding claims, wherein at least one of the binding domains specifically binds to an antigen on an immune effector cell.
- 25 36. A multivalent antibody according to claim 35, wherein the immune effector cell is a T cell.
- 30 37. A multivalent antibody according to claim 36, wherein the antigen on the said T cell is CD3.
38. A multivalent antibody according to any one of claims 35 to 37, wherein at least one of the binding domains specifically binds to an antigen on an aberrant cell.
- 35 39. A multivalent antibody according to claim 38, wherein at least two binding domains specifically bind to an antigen on an aberrant cell.
40. 40. A multivalent antibody according to claim 39, wherein the at least two binding domains specifically bind to at least two different antigens on an aberrant cell.

41. A multivalent antibody according to claim 40, wherein the at least two binding domains specifically binding to at least EGFR and PD-L1.

42. A multivalent antibody according to any one of claims 35 to 41, wherein a first binding domain specifically binds PD-L1; a second binding domain specifically binds CD3, and a third binding domain specifically binds to EGFR.

43. A multivalent antibody according to any one of claims 38 to 42, wherein the aberrant cell is a tumor cell.

10

44. A multivalent antibody according to any one of claims 35-43, wherein the base antibody comprises binding domains 1 and 2 and wherein the additional binding domain 3 is linked to binding domain 1 and wherein an optional additional binding domain 4 is linked to binding domain 2.

15

45. The multivalent antibody of claim 44, wherein binding domain 1 is a CD3 binding domain and binding domains 2 and 3 bind to different target cell antigens.

20

46. The multivalent antibody of claim 44, wherein binding domain 2 is a CD3 binding domain and binding domains 1 and 3 bind to different target cell antigens.

47. The multivalent antibody of claim 44, wherein binding domain 3 is a CD3 binding domain and binding domains 1 and 2 bind to different target cell antigens.

25

48. The multivalent antibody of any one of claims 44-47, comprising an binding domain 4 that binds yet a further different target cell antigen.

49. The multivalent antibody of any one of claims 44-48, wherein a first of said target cell antigen binding domain binds PD-L1, EGFR, CD137, CLEC12A, fibrinogen, or thyroglobulin.

30

50. The multivalent antibody of any one of claims 44-49, wherein said first and second target cell binding domain bind different antigens.

35

51. The multivalent antibody of claim 50, wherein a first and a second of said target cell antigen binding domains bind antigens selected from PD-L1, EGFR, CD137, CLEC12A fibrinogen, and thyroglobulin.

52. The multivalent antibody of any one of claims 44-51, wherein the CD3 binding domain comprises a heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino

acid sequence of the CDR1, CDR2 and CDR3 of MF8057, or of MF8058, or of MF8078, or of MF8397, or of MF8508, or of MF9249 or of MF9267.

53. The multivalent antibody of any one of claims 44-52, wherein the CD3 binding domain  
5 comprises a heavy chain variable region with the amino acid sequence of the VH of MF8057, of  
MF8058, of MF8078, of MF8397, of MF8508, of MF9249 or of MF9267 with 0-10, preferably 0-5  
amino acid insertions, deletions, substitutions, additions or a combination thereof at one or more  
positions other than the CDRs.

10 54. The multivalent antibody of any one of claims 44-53, wherein a target cell antigen  
binding domain is a PD-L1 binding domain.

15 55. The multivalent antibody of claim 54, wherein the PD-L1 binding domain comprises a  
heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid  
sequence of the CDR1, CDR2 and CDR3 of MF5377, or of MF5444, or of MF5380.

20 56. The multivalent antibody of claim 54 or claim 55, wherein the PD-L1 binding domain  
comprises a heavy chain variable region with the amino acid sequence of the VH of MF5377, of  
MF5444, or of MF5380 with 0-10, preferably 0-5 amino acid insertions, deletions, substitutions,  
additions or a combination thereof at one or more positions other than the CDRs.

57. The multivalent antibody of any one of claims 44-56, wherein a target cell antigen  
binding domain is a EGFR binding domain.

25 58. The multivalent antibody of claim 57, wherein the EGFR binding domain comprises a  
heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid  
sequence of the CDR1, CDR2 and CDR3 of MF8233, or of MF9891, or of MF9886, or of  
MF9873, or of MF9988.

30 59. The multivalent antibody of claim 57 or claim 58, wherein the EGFR binding domain  
comprises a heavy chain variable region with the amino acid sequence of the VH of MF8233, of  
MF9891, of MF9886, of MF9873, or of MF9988 with 0-10, preferably 0-5 amino acid insertions,  
deletions, substitutions, additions or a combination thereof at one or more positions other than  
the CDRs.

35 60. The multivalent antibody of any one of claims 44-59, wherein a target cell antigen  
binding domain is a CLEC12A binding domain.

61. The multivalent antibody of claim 60, wherein the CLEC12A binding domain comprises a heavy chain variable region comprising a CDR1, CDR2 and CDR3 with the amino acid sequence of the CDR1, CDR2 and CDR3 of MF4327.

5 62. The multivalent antibody of claim 60 or claim 61, wherein the CLEC12A binding domain comprises a heavy chain variable region with the amino acid sequence of the VH of MF4327 with 0-10, preferably 0-5 amino acid insertions, deletions, substitutions, additions or a combination thereof at one or more positions other than the CDRs.

10 63. The multivalent antibody of any one of claims 52-62, comprising a CD3 binding domain, an EGFR binding domain and a PD-L1 binding domain.

15 64. A method for the preparation of a multivalent antibody, which method comprises: providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into a multivalent antibody according to any one of claims 1 to 63; and cultivating said host cell under conditions to provide for expression of the polypeptides and for their assembly into a multivalent antibody.

20 65. A method for the preparation of a multivalent antibody, which method comprises: providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into: (i) a base antibody portion which comprises two binding domains; (ii) at least one additional binding domain; and (iii) at least one linker which comprises a hinge sequence or a sequence derived from a hinge sequence,

25 cultivating said host cell under conditions to provide for expression of the base antibody portion, the at least one additional binding domain and the at least one linker and for their assembly into a multivalent antibody,

30 wherein, upon assembly, the base antibody portion is connected by a linker to the at least one additional binding domain and wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region.

35 66. A method for the preparation of a multivalent antibody, which method comprises: providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into: (i) a base antibody portion which comprises two binding domains; (ii) at least one additional binding domain; and (iii) at least one linker,

cultivating said host cell under conditions to provide for expression of the base antibody portion, the at least one additional binding domain and the at least one linker and for their assembly into a multivalent antibody,

5        wherein, upon assembly, the base antibody portion is connected by a linker to the at least one additional binding domain, wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region and wherein the multivalent antibody binds to at least three different epitopes.

10      67.      A method for the preparation of a multivalent antibody, which method comprises:

providing a cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into: (i) a base antibody portion which comprises two binding domains; (ii) at least one additional binding domain; and (iii) at least one linker,

15      cultivating said host cell under conditions to provide for expression of the base antibody portion, the at least one additional binding domain and the at least one linker and for their assembly into a multivalent antibody,

20        wherein, upon assembly, the base antibody portion is connected by a linker to the at least one additional binding domain, wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region and wherein each binding domain includes a rearranged variable region which is obtained from, based on or derived from a nucleic acid that has undergone somatic recombination in response to antigenic stimulation to form a Fab domain with said common variable region.

25      68.      A method according to any one of claims 64 to 67, wherein at least one additional binding domain comprises a Fv domain, a Fab domain or a modified Fab domain.

69.      A method according to any one of claims 64 to 68, wherein at least one additional binding domain comprises a CH1 region.

30      70.      A method according to any one of claims 64 to 69, wherein the common variable region is in the form of a common light chain comprising VL-CL.

71.      A method according to claim 70, wherein the common light chain comprises the sequence of SEQ ID NO: 29.

35

72.      A method according to any one of claims 64 to 71, wherein the common variable region is in the form of a common heavy chain comprising VH-CH1.

73. A method according to any one of claims 64 to 72, wherein each binding domain of the base antibody portion binds a different epitope.

74. A cell which comprises one or more nucleic acid sequences encoding polypeptides 5 which are capable of assembly into a multivalent antibody according to any one of claims 1 to 63.

75. A cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into: (i) a base antibody portion which comprises two binding 10 domains; (ii) at least one additional binding domain; and (iii) at least one linker which comprises a hinge sequence or a sequence derived from a hinge sequence,

wherein, upon assembly, the base antibody portion is connected by a linker to the at 15 least one additional binding domain and wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region.

76. A cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into: (i) a base antibody portion which comprises two binding 20 domains; (ii) at least one additional binding domain; and (iii) at least one linker,

wherein, upon assembly, the base antibody portion is connected by a linker to the at least one additional binding domain, wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region and wherein the multivalent antibody binds to at least three different epitopes.

25

77. A cell which comprises one or more nucleic acid sequences encoding polypeptides which are capable of assembly into: (i) a base antibody portion which comprises two binding domains; (ii) at least one additional binding domain; and (iii) at least one linker,

wherein, upon assembly, the base antibody portion is connected by a linker to the at 30 least one additional binding domain, wherein each binding domain of the base antibody portion and each of the at least one additional binding domains all have a common variable region and wherein each common variable region has co-evolved with its cognate region.

78. A polypeptide comprising an amino acid sequence as set out in any one of SEQ ID 35 NOs: SEQ ID NOs: 3 to 5, 7 to 11 or 13 to 24 or a polypeptide having at least about 85% sequence identity to any one thereto.

79. A nucleic acid sequence encoding a polypeptide according to claim 78.

80. A vector comprising a nucleic acid sequence according to claim 79.
81. A pharmaceutical composition which comprises an antibody according to any one of claims 1 to 63 and a pharmaceutically acceptable carrier and/or diluent.
82. A multivalent antibody according to any one of claims 1 to 63 for use in the treatment of the human or animal body by therapy.
83. A method for the treatment of a human or animal suffering from a medical indication, which method comprises administering to the human or animal a therapeutically effective amount of an antibody according to any one of claims 1 to 63.

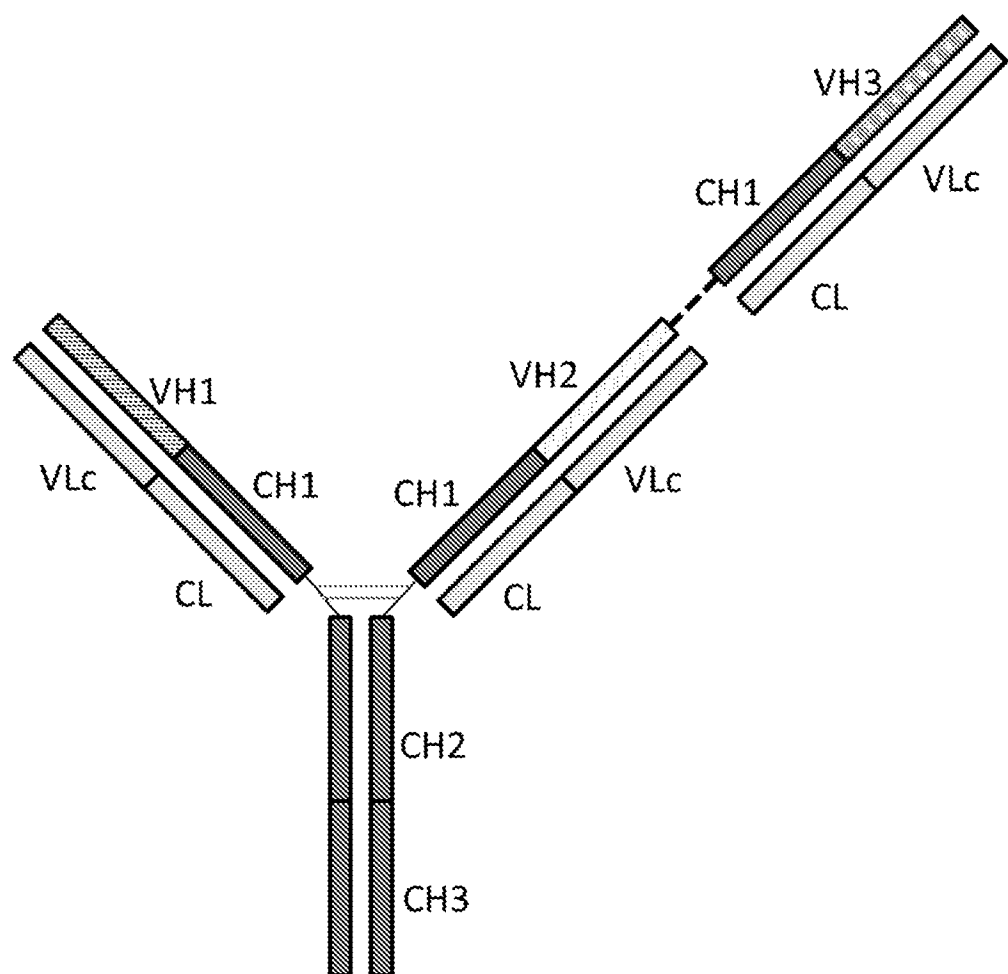


Fig. 1a

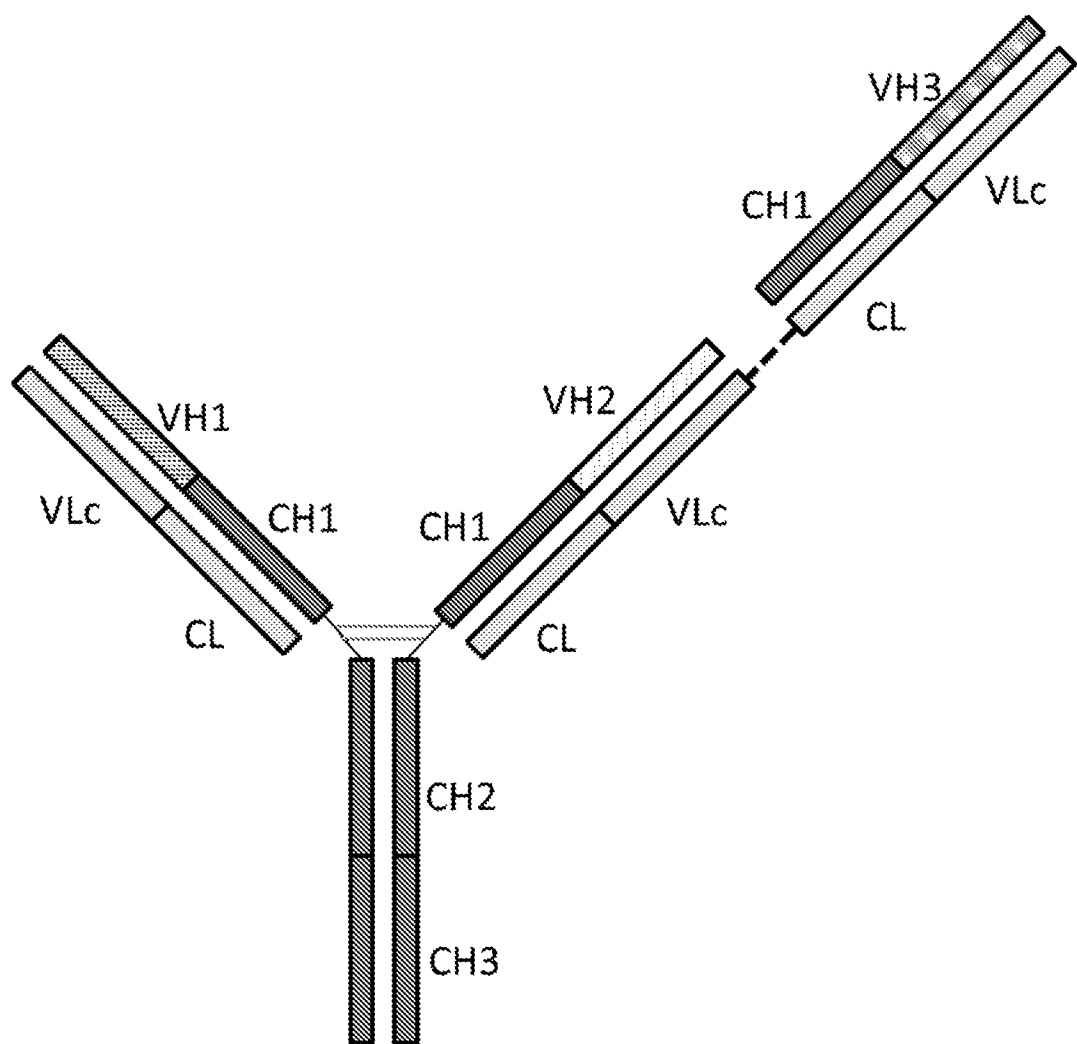


Fig. 1b

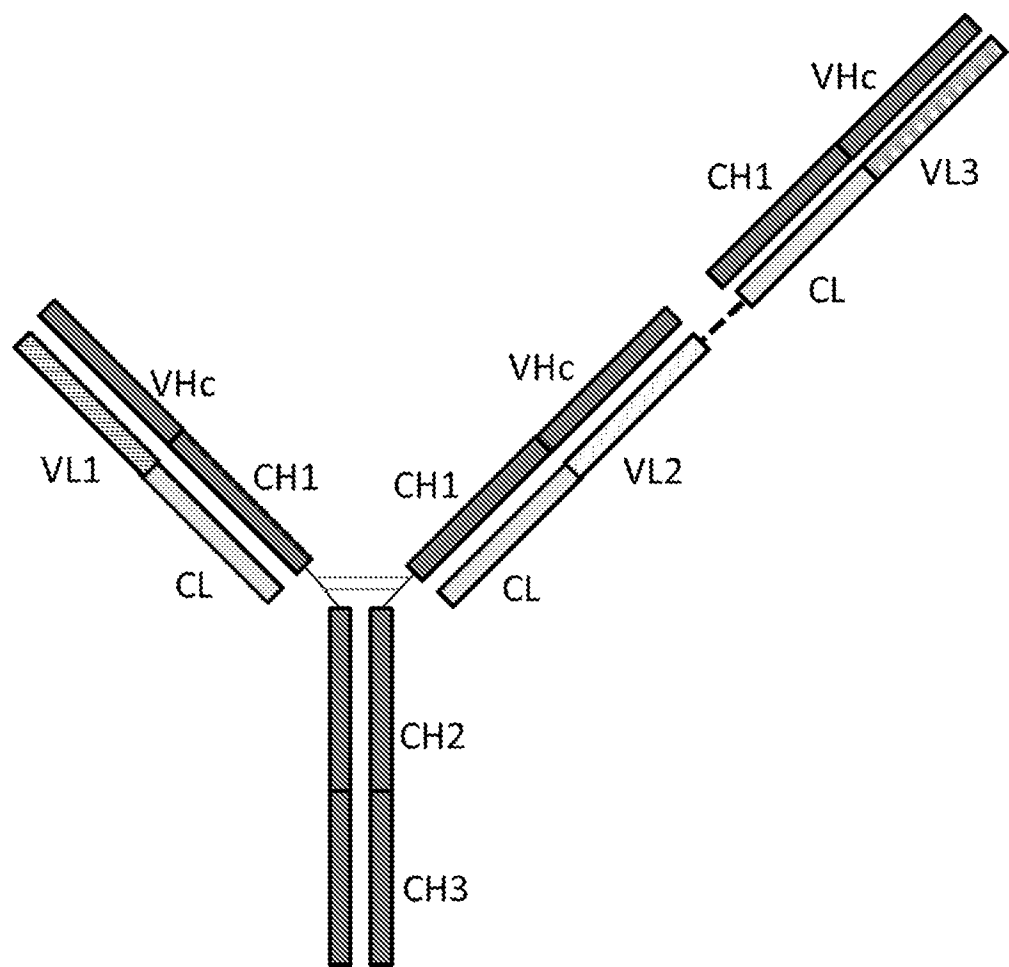


Fig. 1c

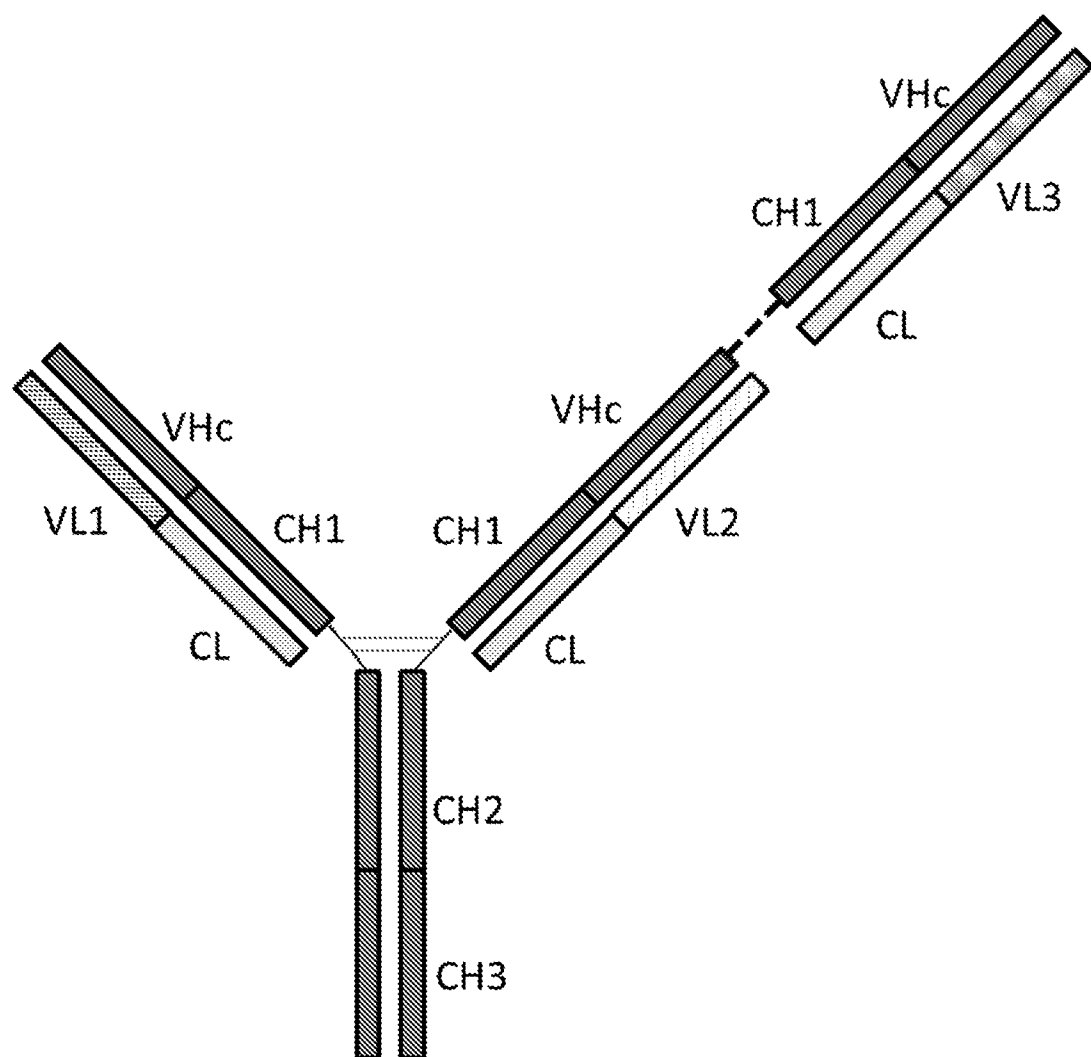


Fig. 1d

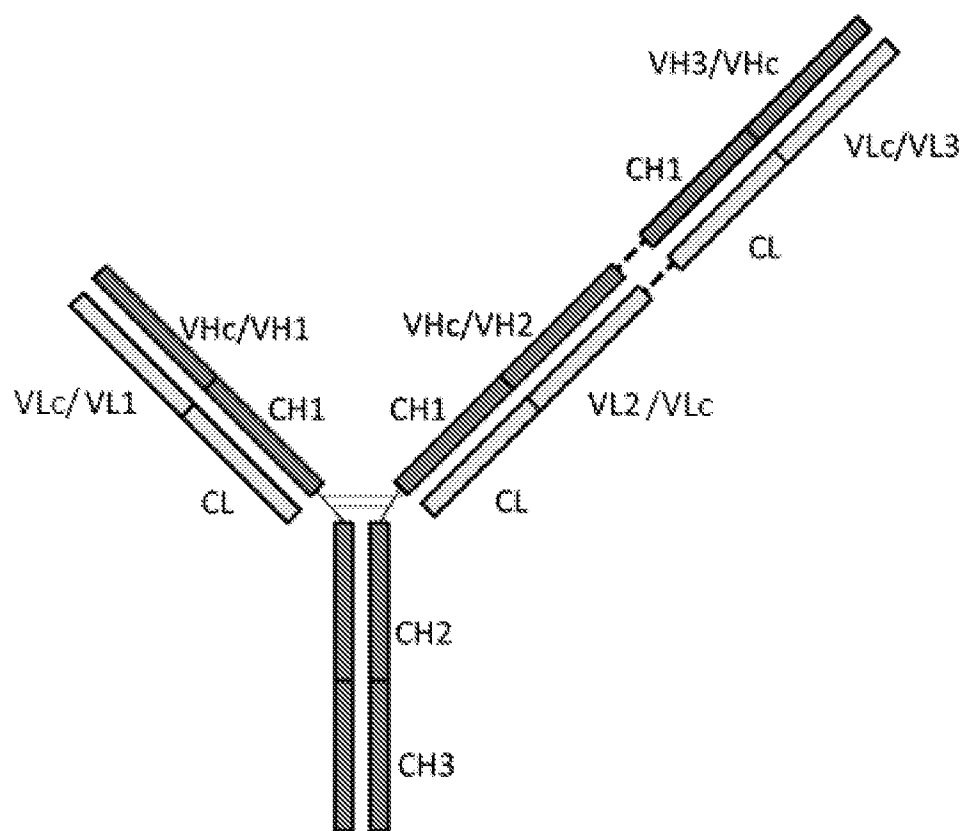


Fig. 1e

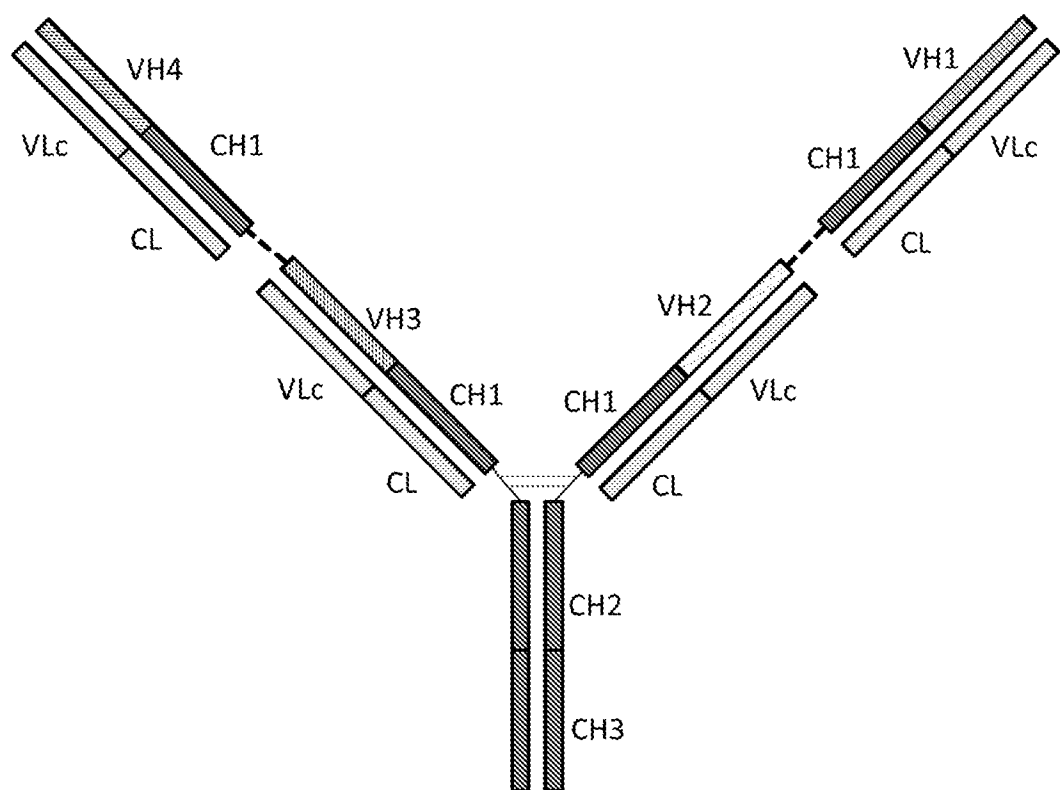


Fig. 1f

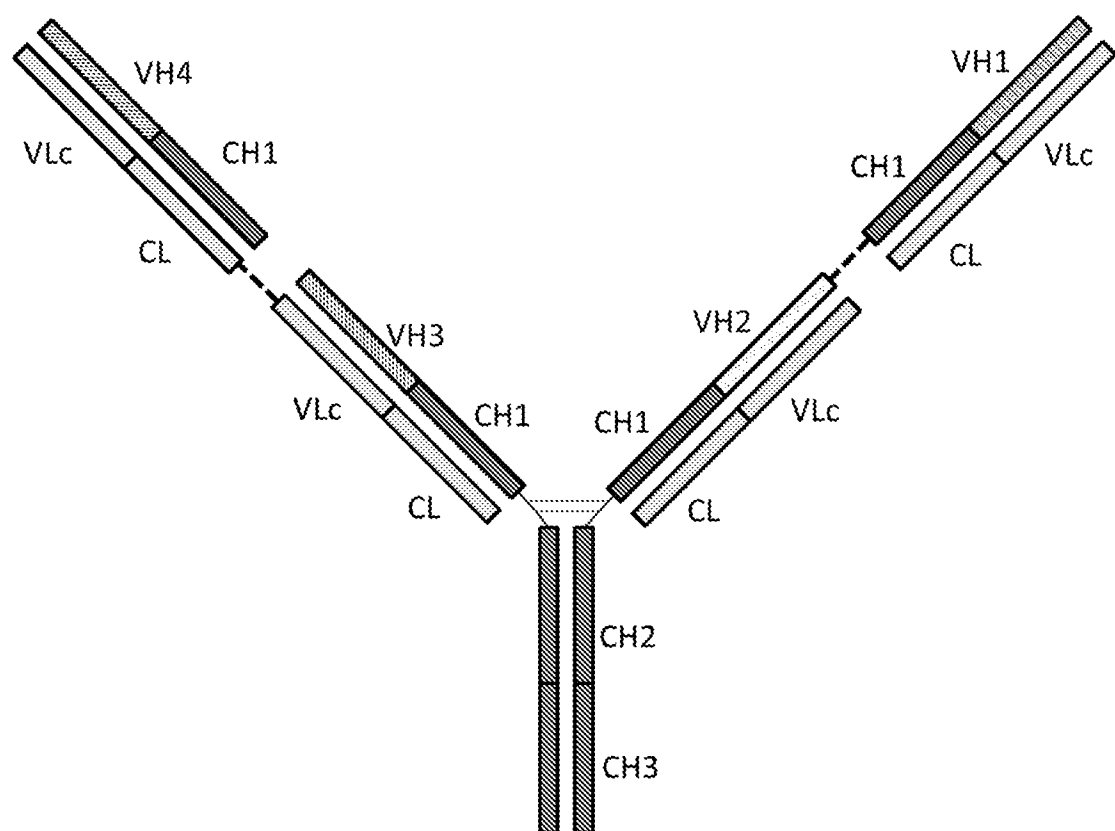


Fig. 1g

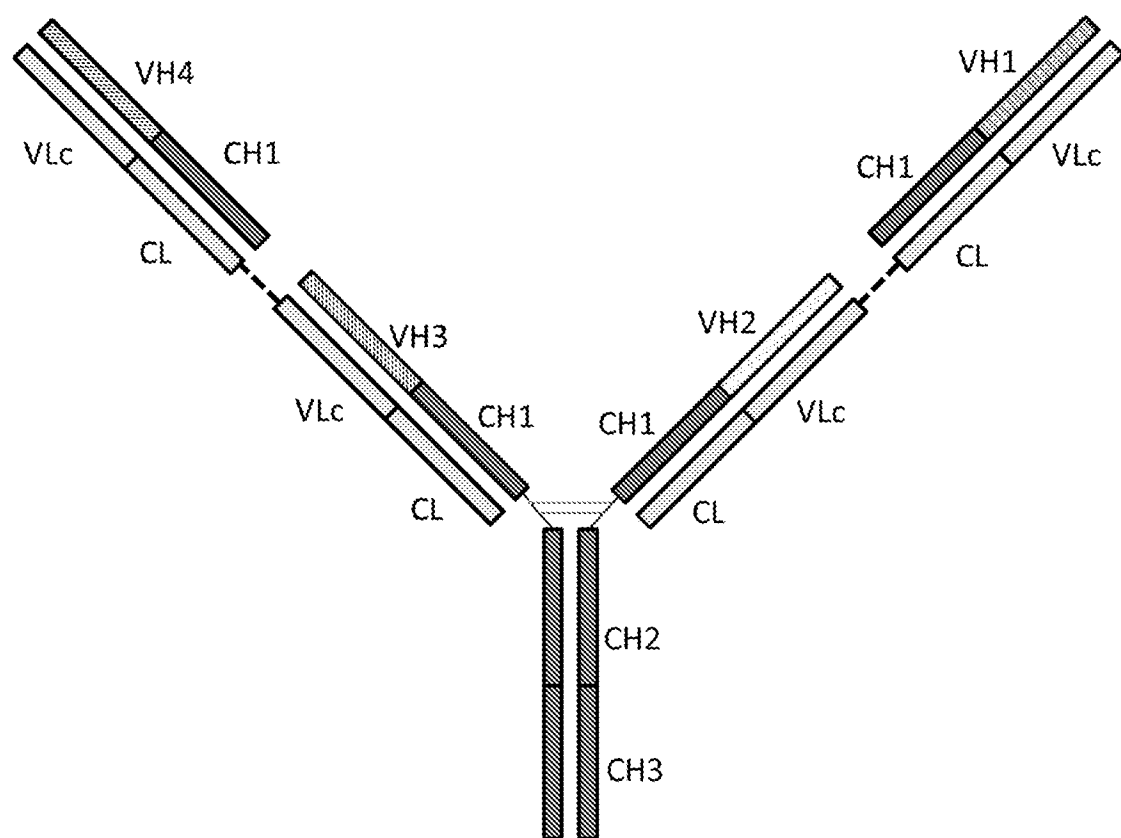


Fig. 1h

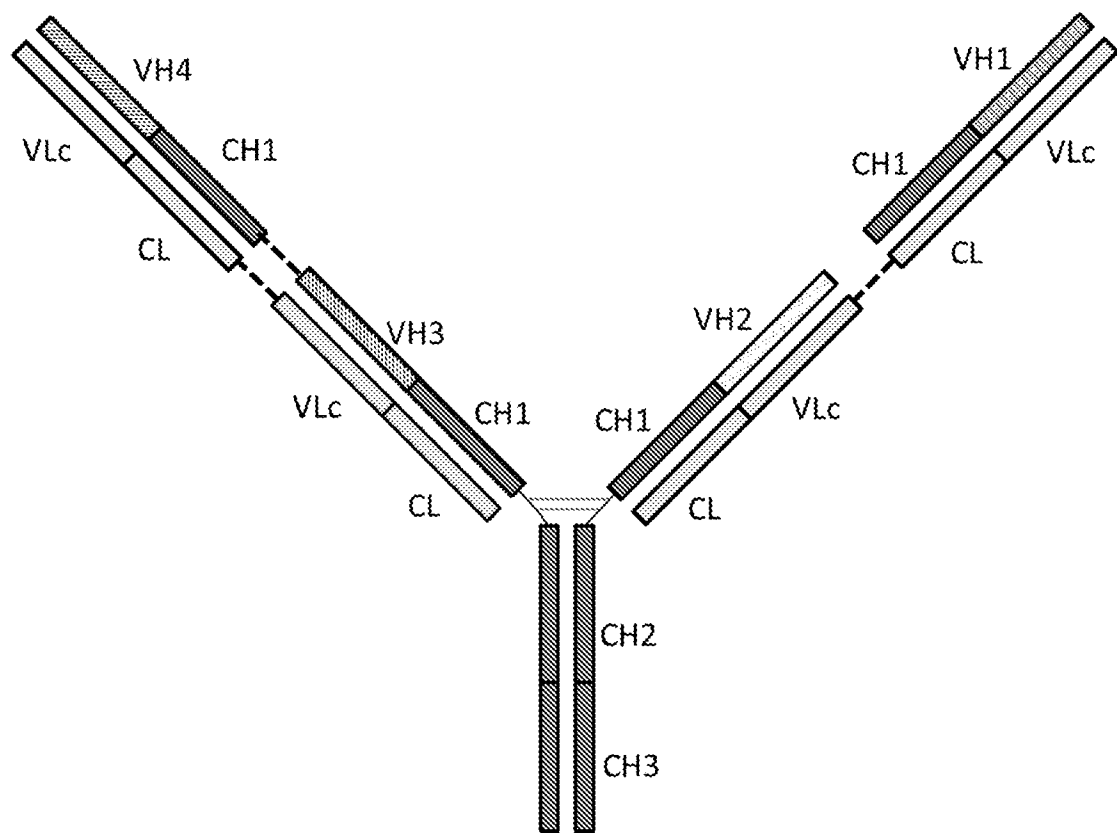


Fig. 1i

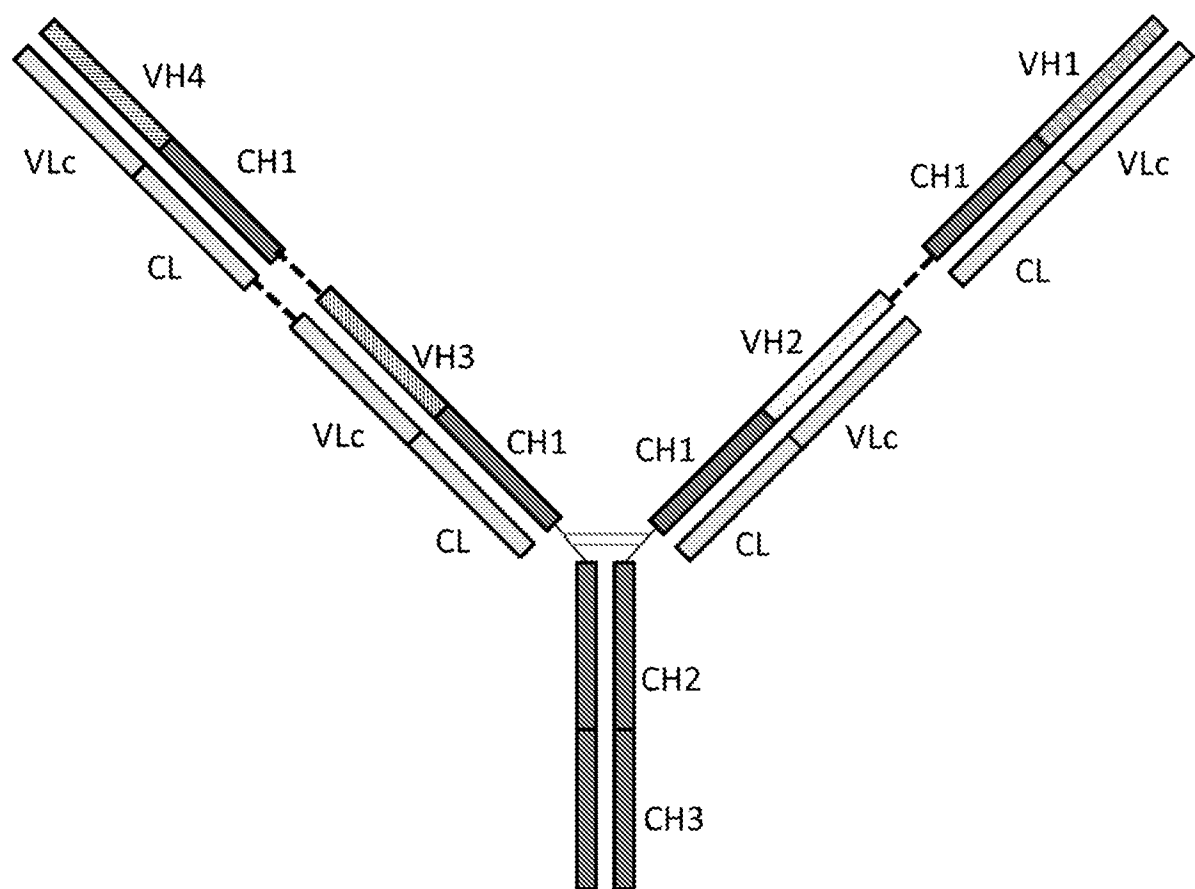


Fig. 1j

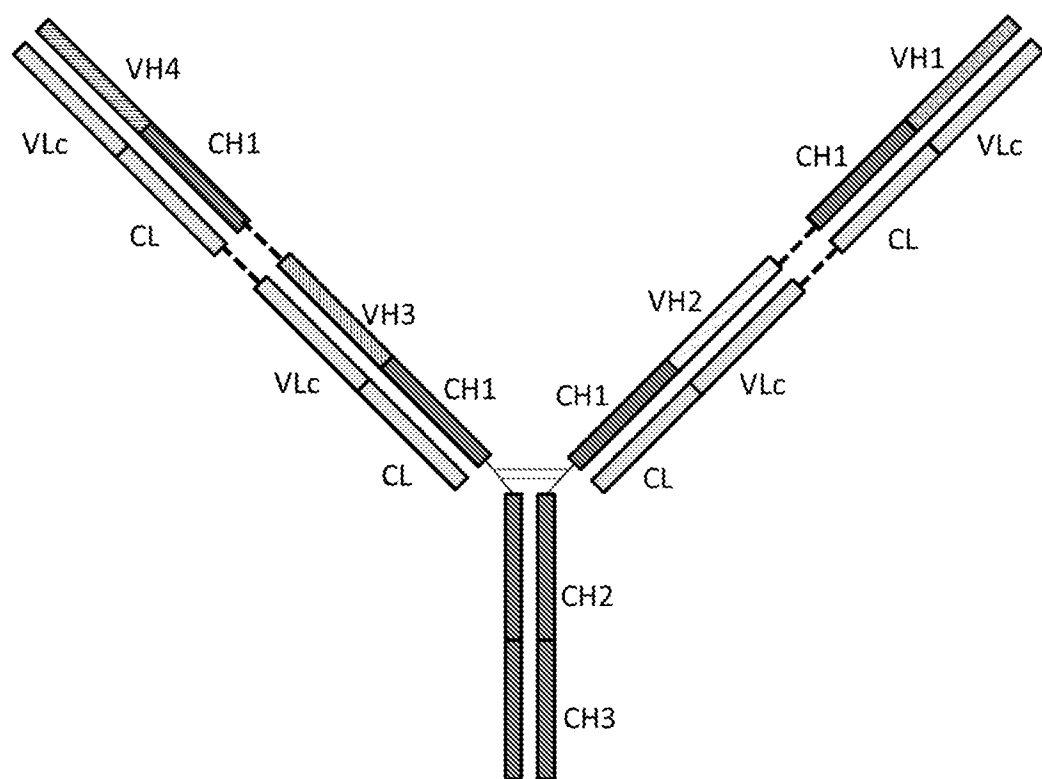


Fig. 1k

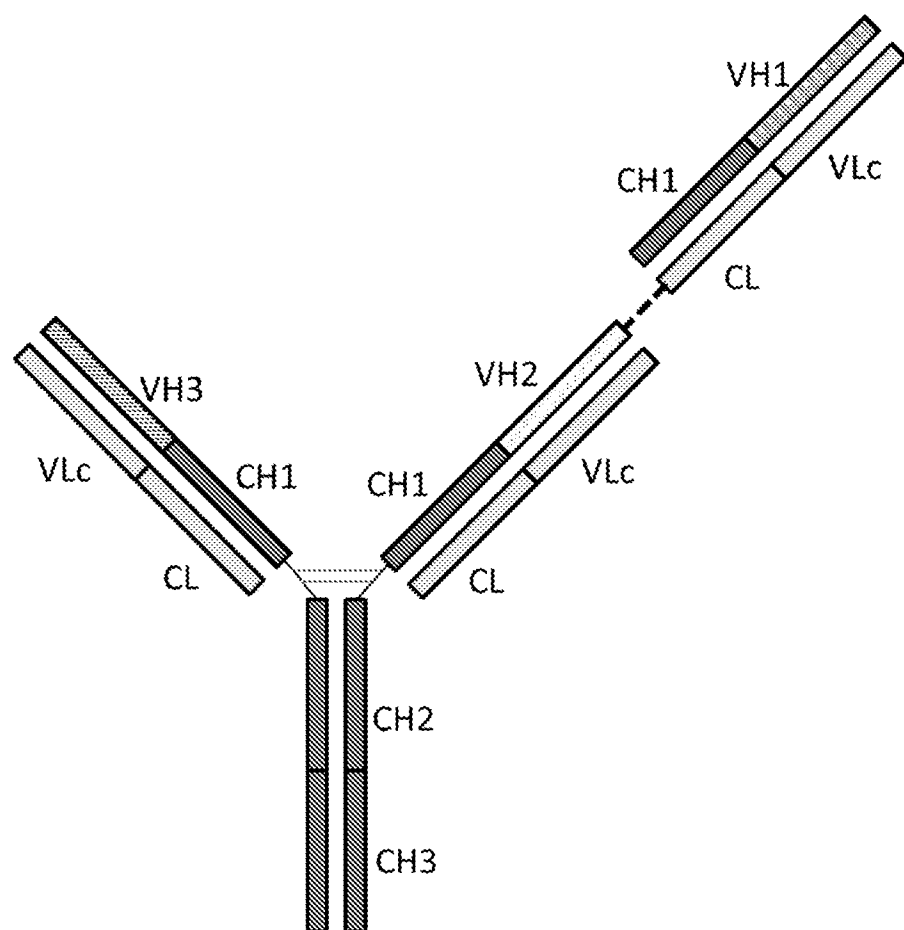


Fig. 11

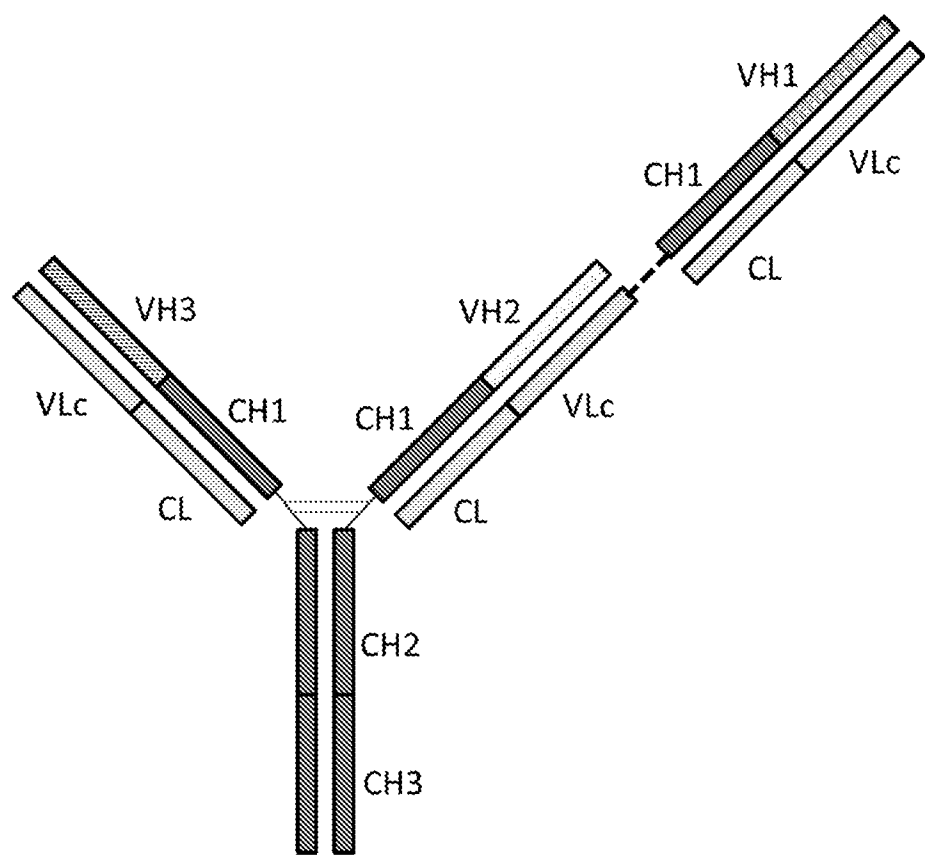


Fig. 1m

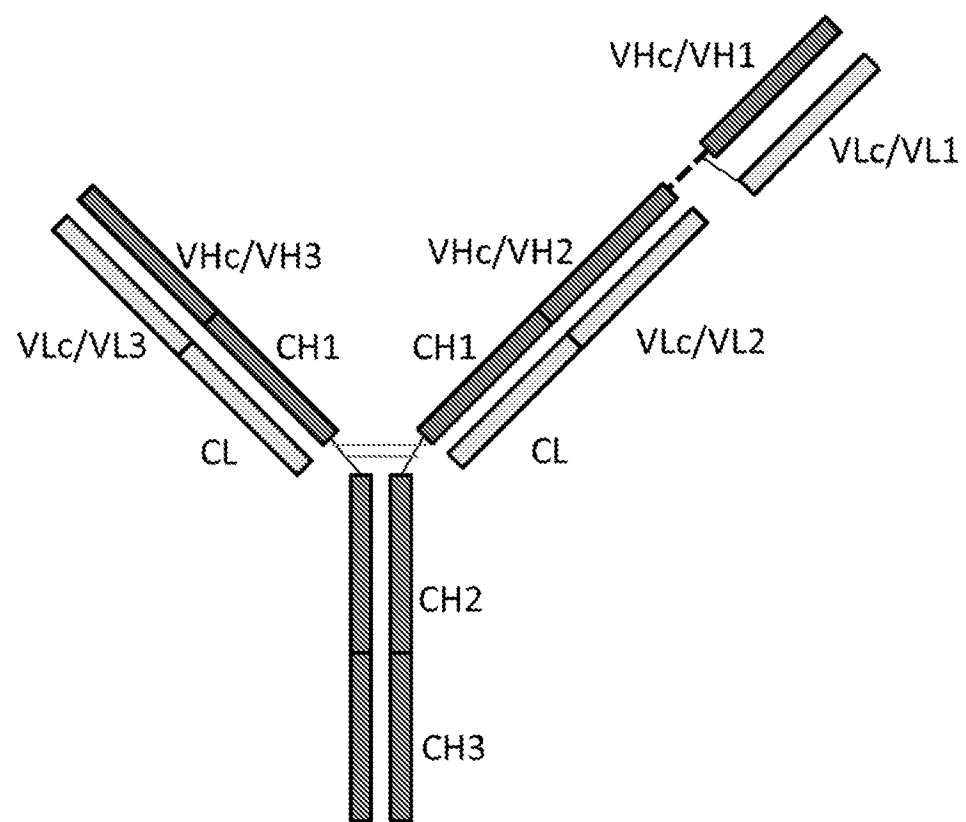


Fig. 1n

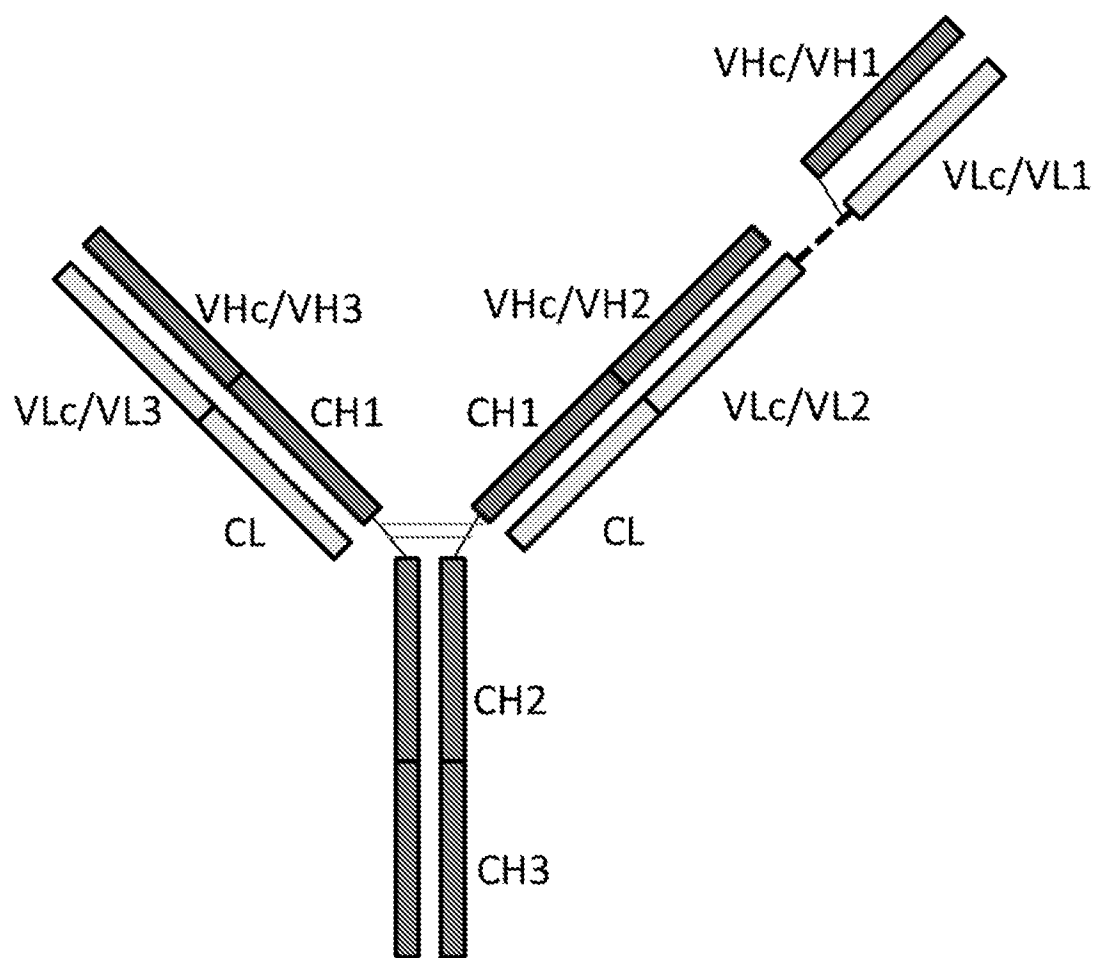


Fig. 1o

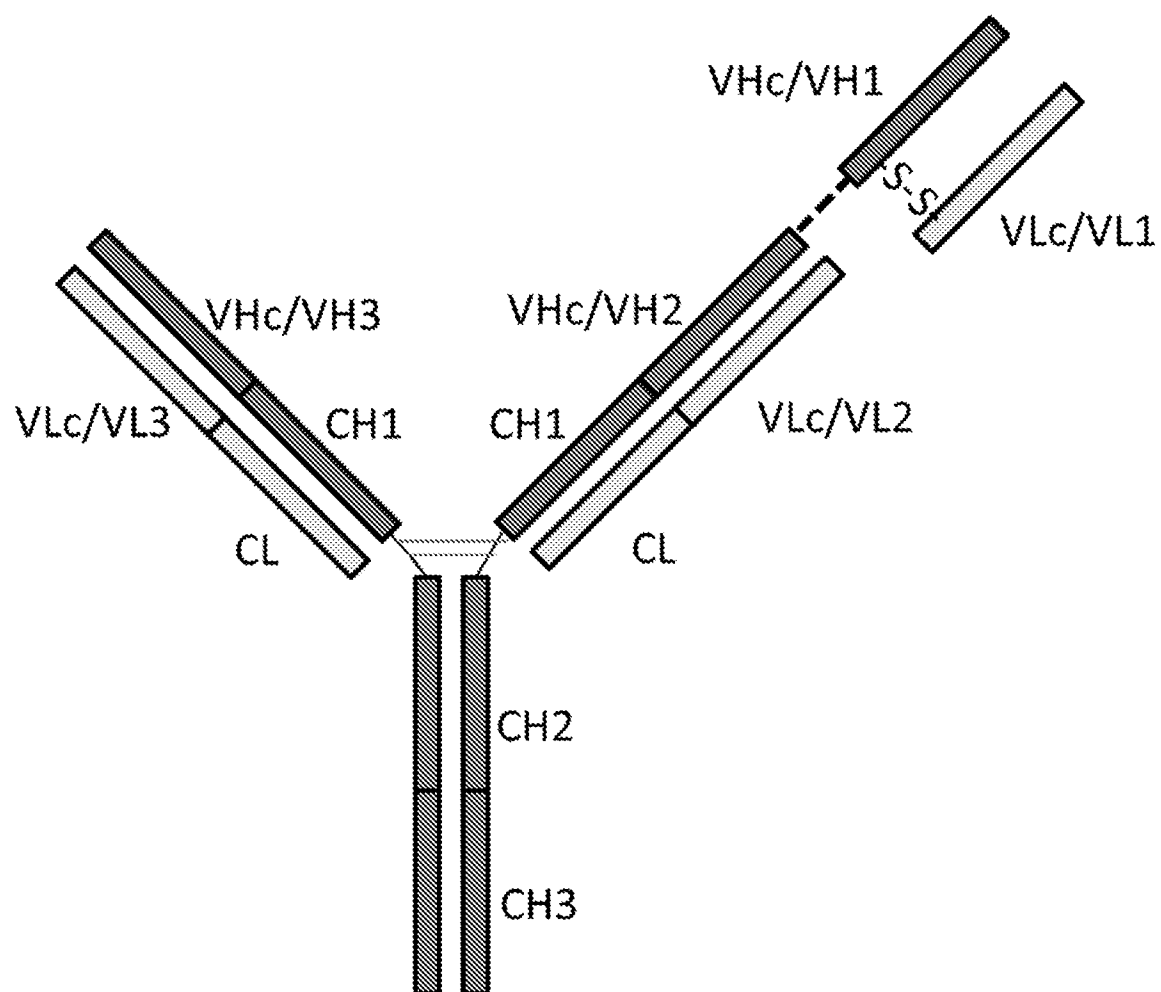


Fig. 1p

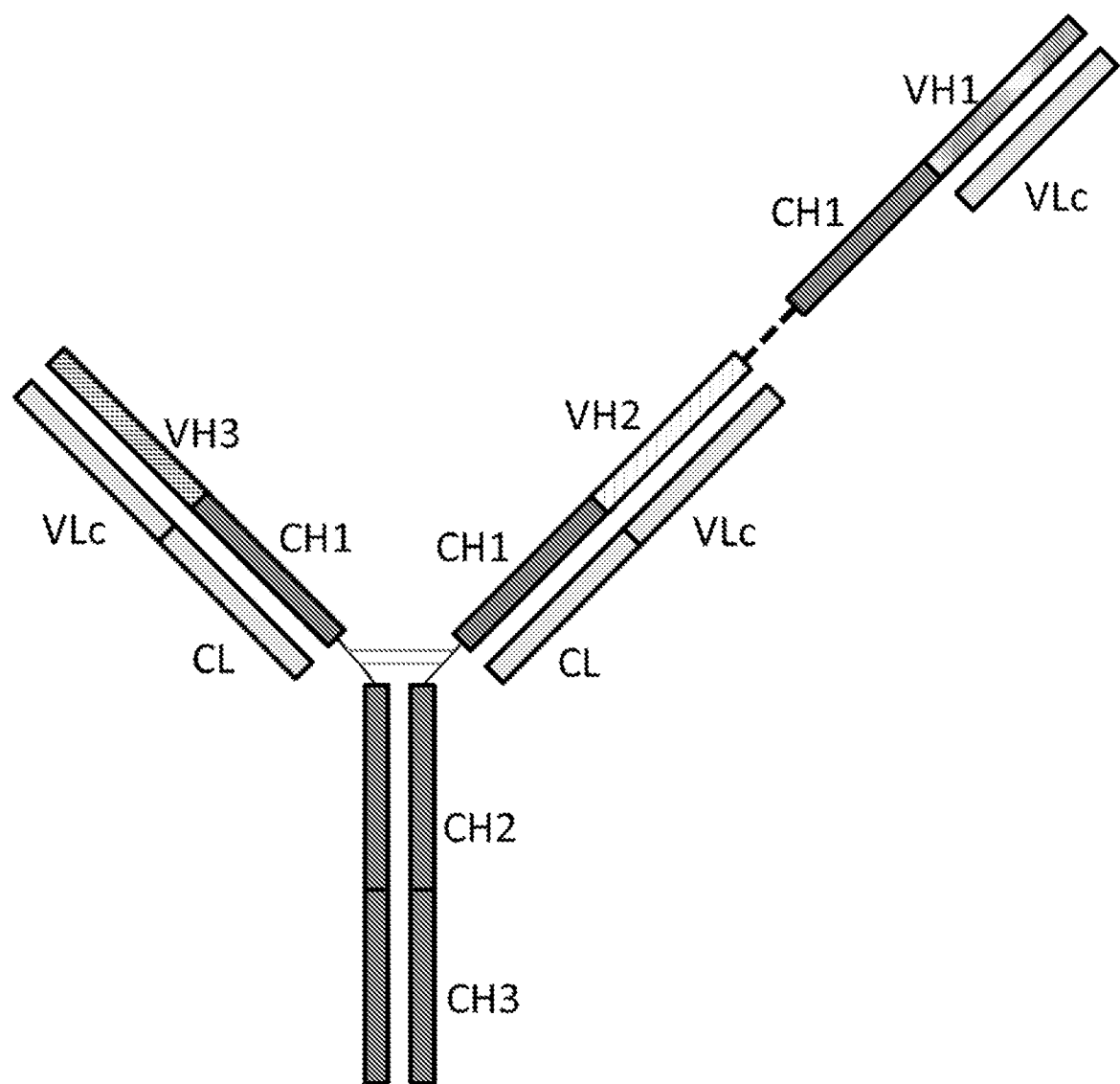


Fig. 1q

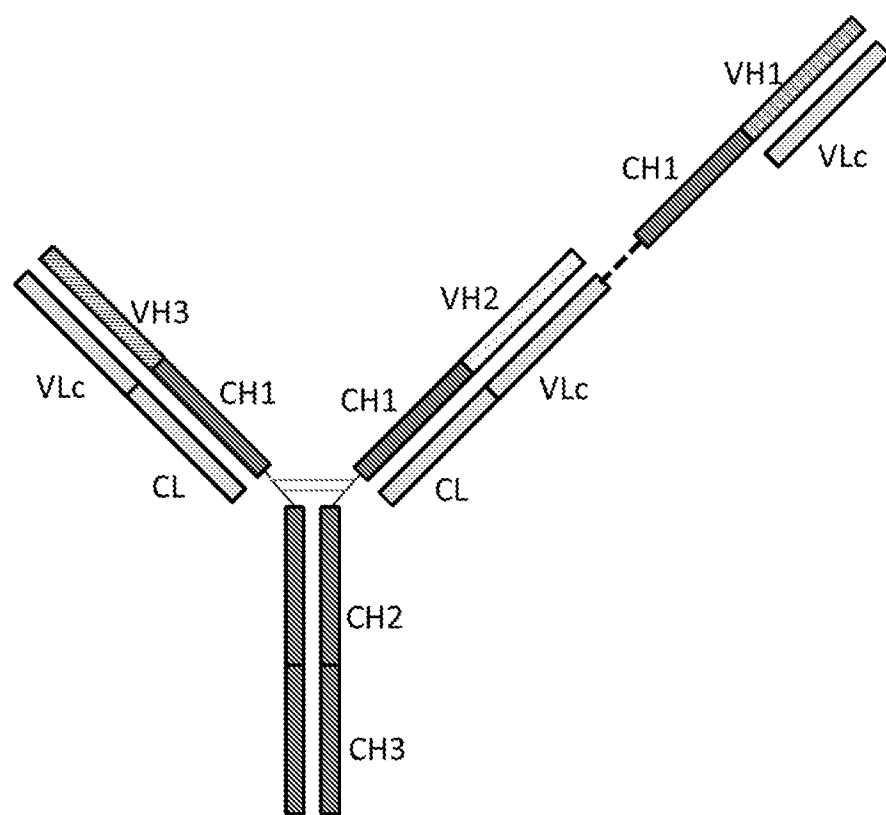


Fig. 1r

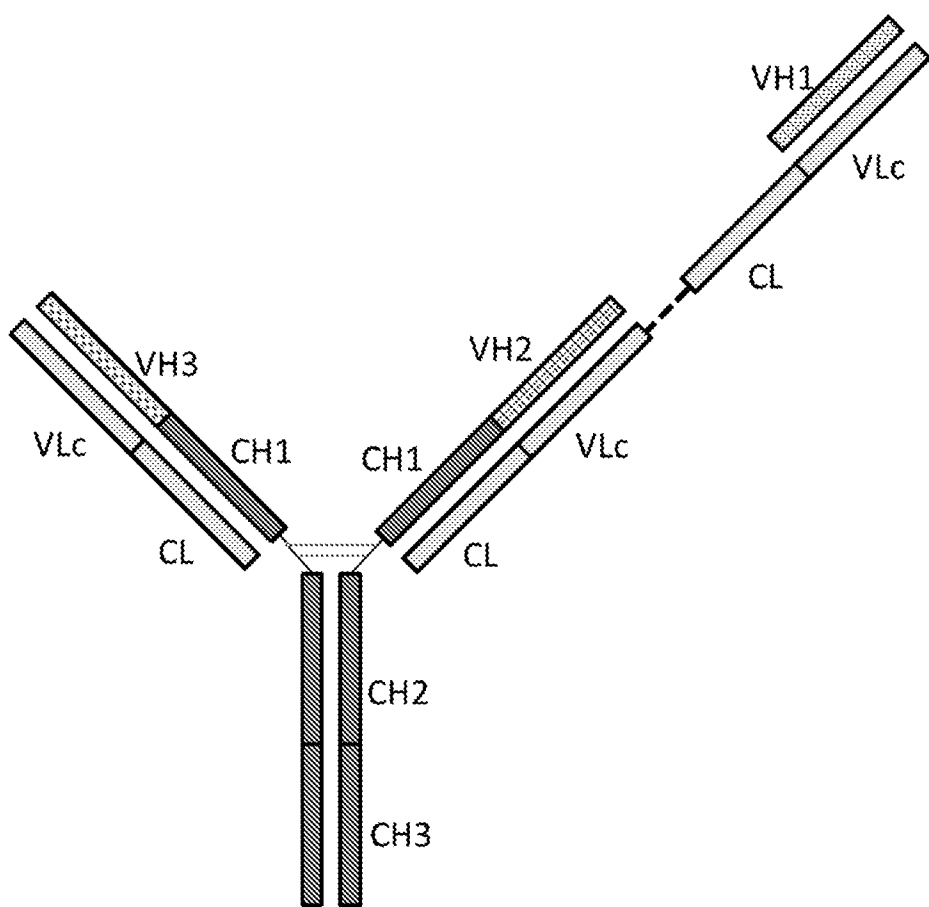


Fig. 1s

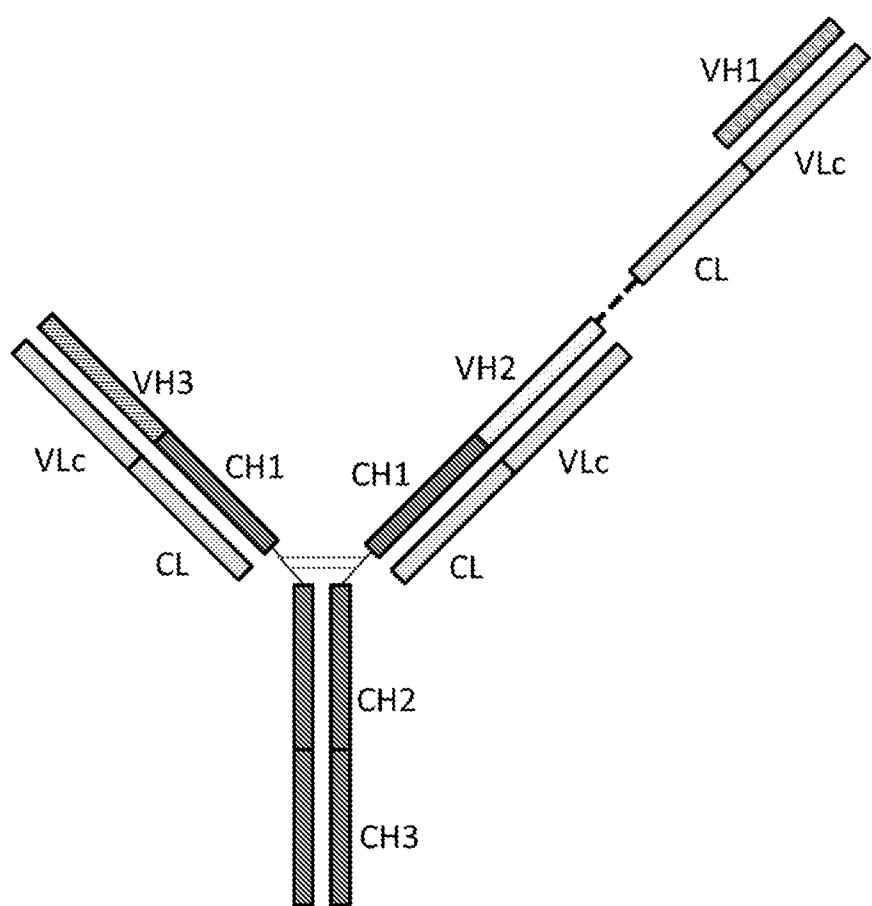


Fig. 1t

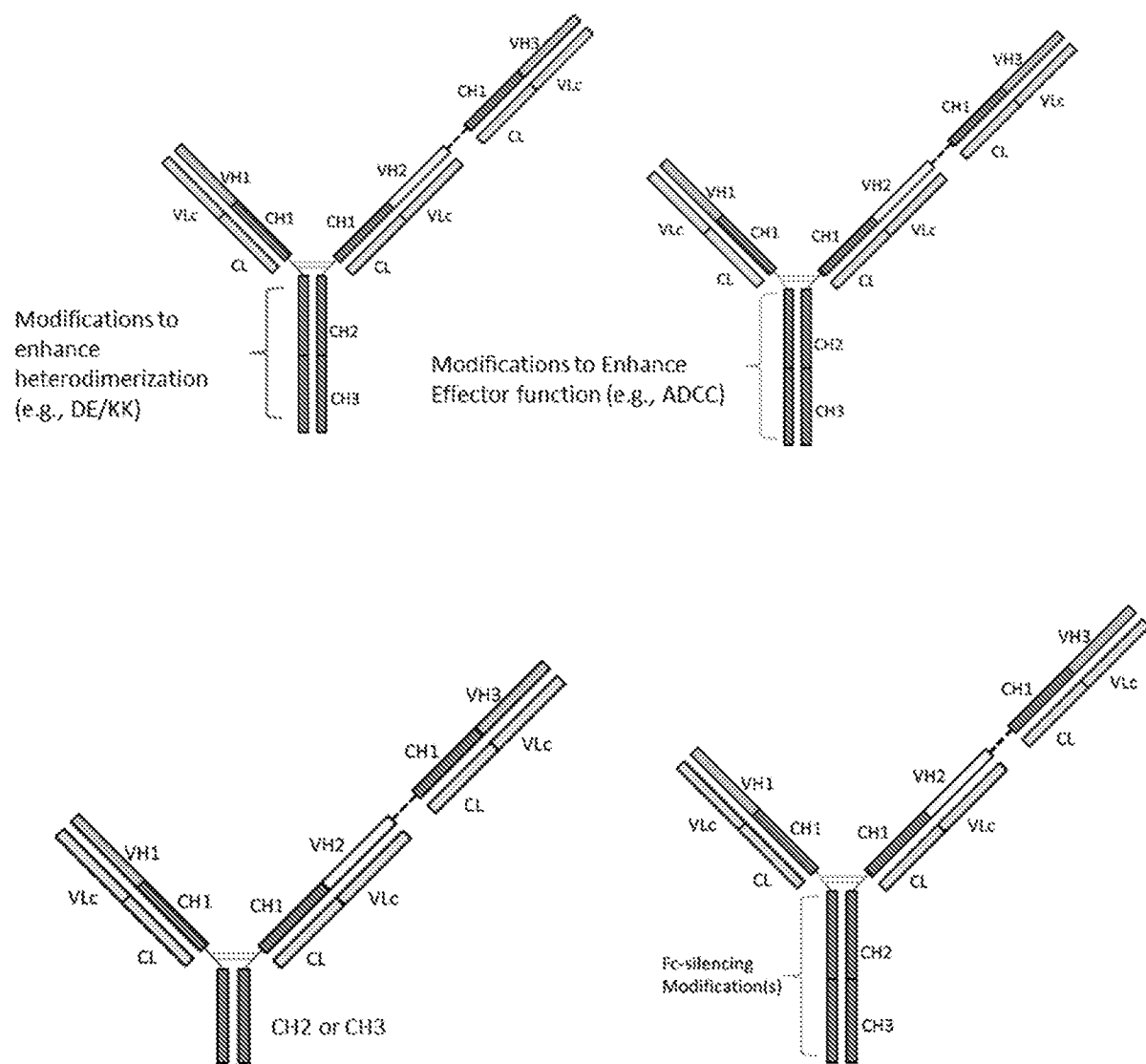


Fig. 1u

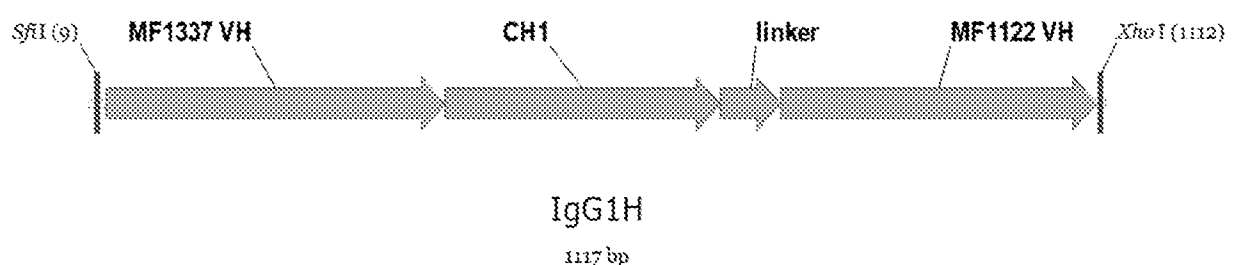


Fig. 2a

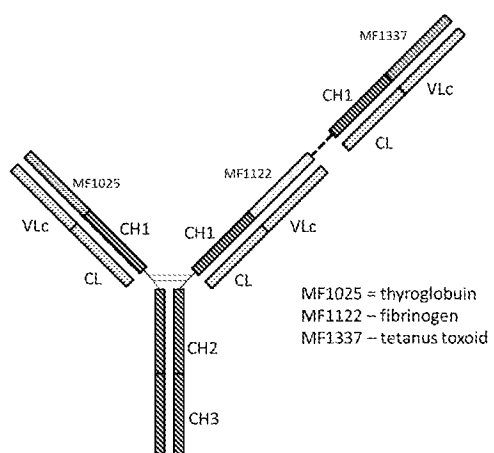


Fig. 2b

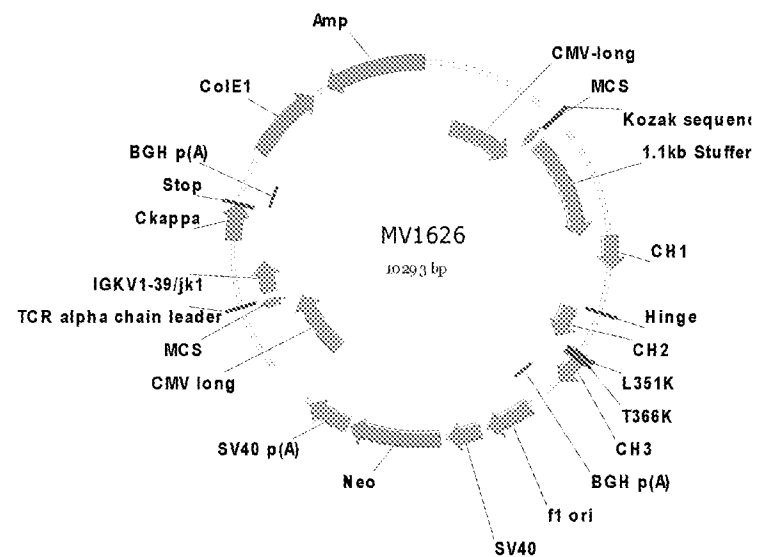


Fig. 3

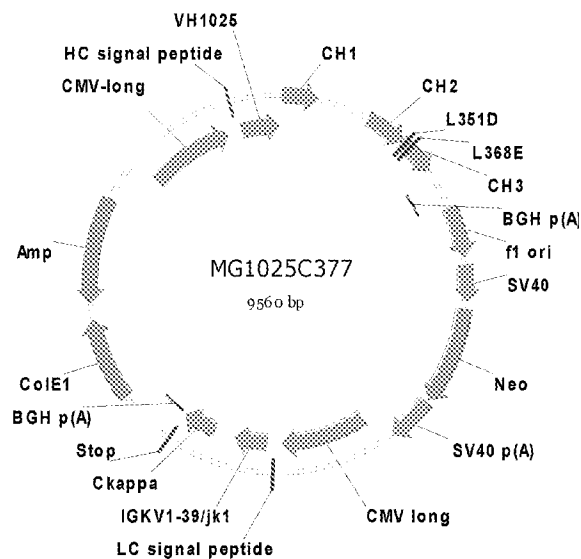


Fig. 4

SEQ ID NO.	Constituents
Translation of IgG1A	ASTKGPSVFLAPSSRKSTSGGTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPSPAPELUGG-----EVQL2.59
Translation of IgG1MH	ASTKGPSVFLAPSSRKSTGGTAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPSP-----EVQL2.60
Translation of IgG1UM	ASTKGPSVFLAPSSRKSTSGGTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPSP-----EVQL2.61
Translation of IgG1Q4S	ASTKGPSVFLAPSSRKSTSGGTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPSP-----EVQL2.63
Translation of IgG1EL	ASTKGPSVFLAPSSRKSTGGTAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPSP-----EVQL2.63
Translation of IgG1R	ASTKGPSVFLAPSSRKSTGGTAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPELLGG-----EVQL2.64
Translation of IgG2AH	ASTKGPSVFLAPSSRKSTGGTAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.65
Translation of IgG2AMH	ASTKGPSVFLAPSSRKSTGGTAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.66
Translation of IgG2AUL	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.67
Translation of IgG2MAGSS	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.68
Translation of IgG2A431	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.69
Translation of IgG2B8R	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.70
Translation of IgG2B4H	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.71
Translation of IgG2B3MH	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.72
Translation of IgG2B1UL	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.73
Translation of IgG3B8R	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.74
Translation of IgG3B1H	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.75
Translation of IgG3B4H	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.76
Translation of IgG3B8	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.77
Translation of IgG4B	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.78
Translation of IgG4B4H	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.79
Translation of IgG4B1H	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.80
Translation of IgG4R	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.81
Translation of IgG4UL	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.82
Constituents	ASTKGPSVFLAPCSRSTSESTAAALGCLVKDYFPEPYTVSNSGALTSGVHTFPALQSSGLYLSISVTVPESSSLGTTOTYICNVNHHKPSTKVDKRVEEKSCDKTHTSPPAPVAG-----EVQL2.83

Figure 5

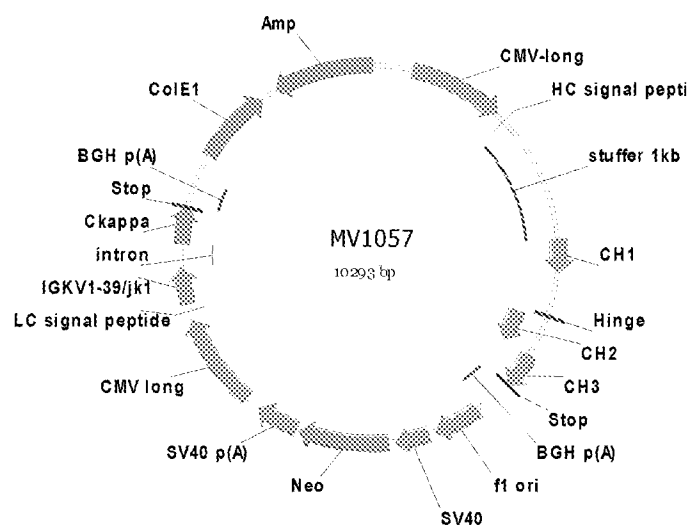


Figure 6

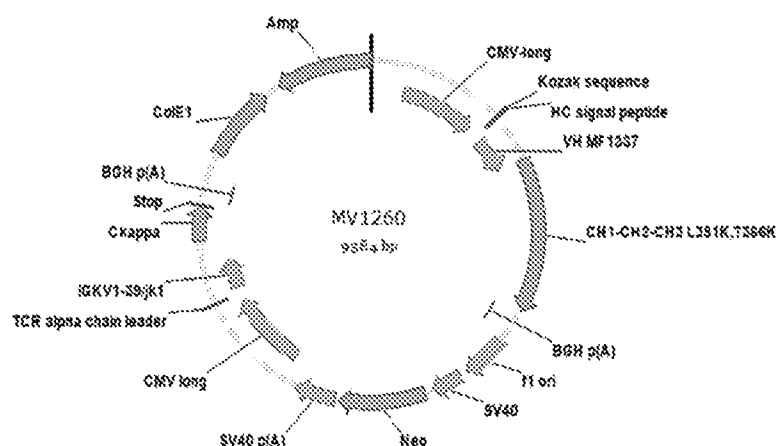


Figure 7

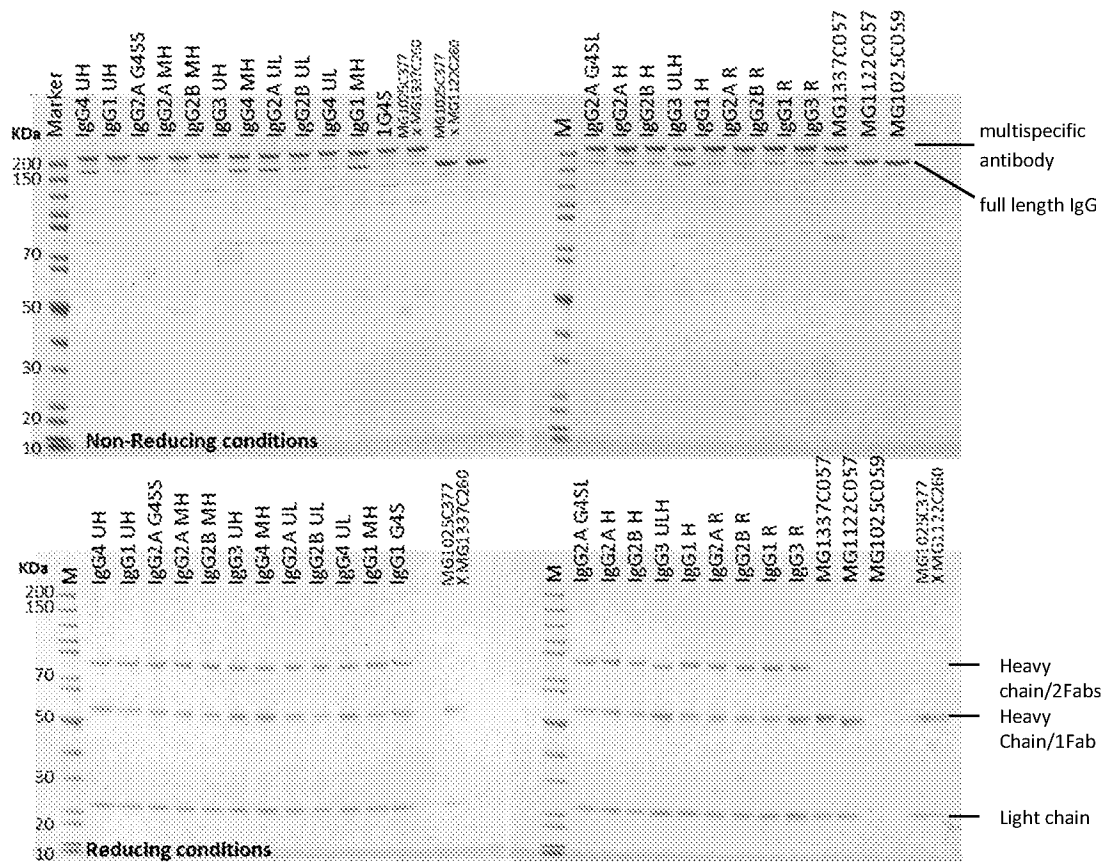


Fig. 8

Fig. 9

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Fig. 10

**Figure 11A (SEQ ID NO: 35)**

DIQMTQSPSSLSASVGDRVITCRASQSISSYLNWYQQKPGKAPKLLIYAASSLQSGVP  
SRFSGSGSGTDFTLTISLQPEDFATYYCQQSYSTPPTFGQGTKVEIK  
RTVAAPSVFIFPPSDEQLKSGTASVVCLNNFYPREAKVQWKVDNALQSGNSQESVTE  
QDSKDSTYSLSTTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC

**Figure 11B (SEQ ID NOs: 36 and 37)**

gacatccagatgacccagtctccatcctccctgtctgcatctgttaggagacagagtacc  
D I Q M T Q S P S S L S A S V G D R V T  
atcacttgccggcaagtcagagcattagcagctacttaattggtatcagcagaaacca  
I T C R A S Q S I S S Y L N W Y Q Q K P  
ggaaagccctaagctcctgatctatgctgcatccagttgcaaagtgggtccatca  
G K A P K L L I Y A A S S L Q S G V P S  
agttcagtggcagtggatctggacagattcacttcaccatcagcagtcgcaacct  
R F S G S G S G T D F T L T I S S L Q P  
gaagatttgcaacttactactgtcaacagagttacagtaccctccaacgttcggccaa  
E D F A T Y Y C Q Q S Y S T P P T F G Q  
ggaccaagggtggagatcaa  
G T K V E I K

**Figure 11C (SEQ ID NOs: 38 and 39)**

cgaactgtggctgcaccatctgtcttcatcttcccgccatctgtatggcagttgaaatct  
R T V A A P S V F I F P P S D E Q L K S  
ggaactgcctctgtgtgtgcctgctgaataactctatcccagagaggccaaagtacag  
G T A S V V C L L N N F Y P R E A K V Q  
tggaaagggtggataacgcgcctccaatcggttaactcccaggagagtgtcacagagcaggac  
W K V D N A L Q S G N S Q E S V T E Q D

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**Figure 11C (continued)**

agcaaggacagcacctacagcctcagcagcaccctgacgctgagcaaagcagactacgag  
S K D S T Y S L S S T L T L S K A D Y E  
aaacacaaaagtctacgcctgcgaagtccatcagggcctgagctcgccgtacaaaag  
K H K V Y A C E V T H Q G L S S P V T K  
agcttcaacaggggagagtgttag  
S F N R G E C -

**Figure 11D (SEQ ID NO: 40)**

DIQMTQSPSSLSASVGDRVITCRASQSISSYLNWYQQKPGKAPKLLIYAASSLQSGVP  
SRFSGSGSGTDFTLTISSLQPEDFATYYCQQSYSTPPITFGQQGTRLEIK

**Figure 11E (SEQ ID NO: 41)**

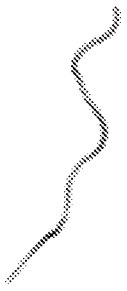
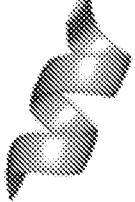
DIQMTQSPSSLSASVGDRVITCRASQSISSYLNWYQQKPGKAPKLLIYAASSLQSGVP  
SRFSGSGSGTDFTLTISSLQPEDFATYYCQQSYSTP

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Fig 12.

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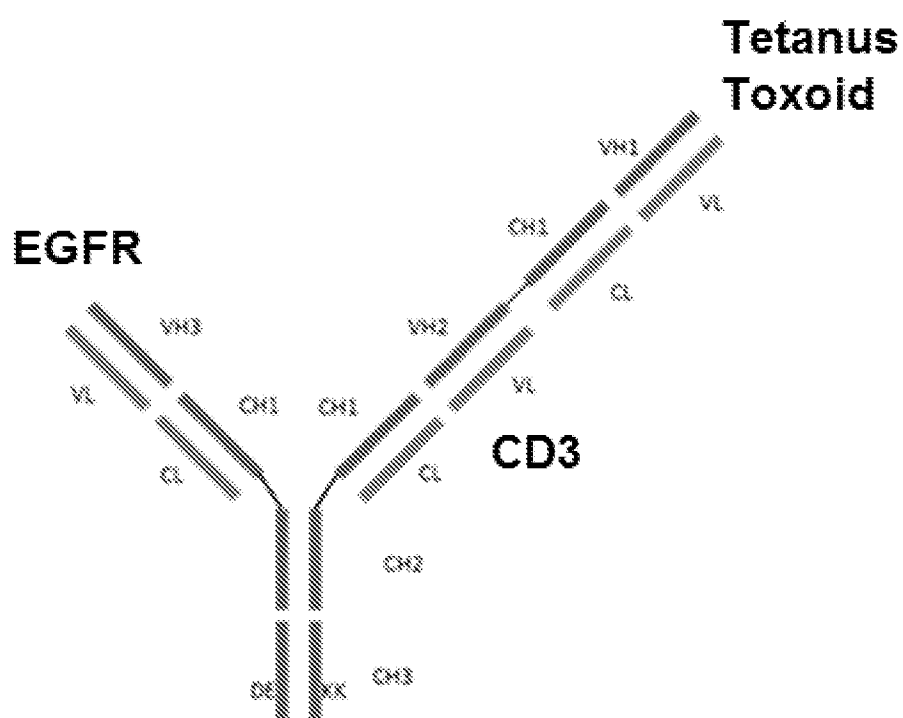
Fig 13.

IgG1 UH	SEQ ID NO: 2	IgG2A H	SEQ ID NO: 15
			
IgG2A MH	SEQ ID NO: 4	IgG2B H	SEQ ID NO: 16
			
IgG1 MH	SEQ ID NO: 11	IgG1 H	SEQ ID NO: 19
			
IgG1G4	SEQ ID NO: 12	IgG2B R	SEQ ID NO: 21
			

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Fig. 14

14a



14b

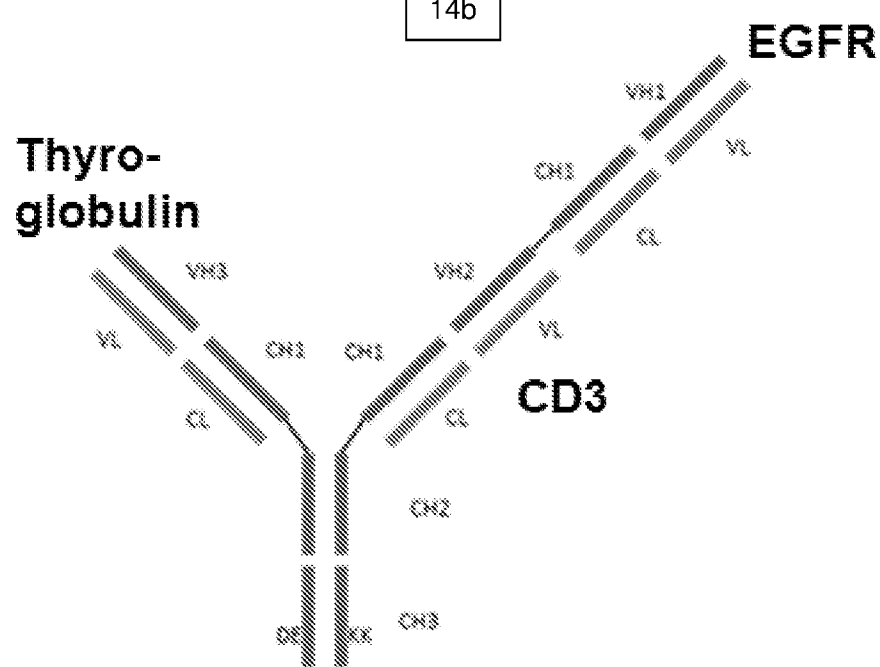


Fig. 15a

T cell Activation:

EGFRxCD3:TT

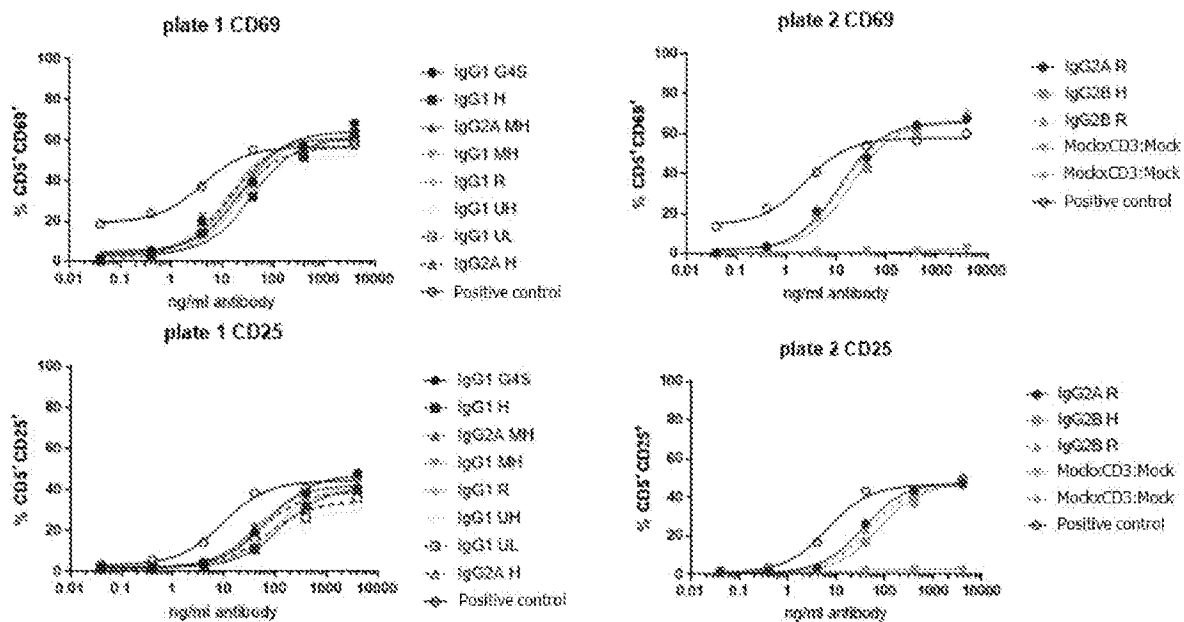


Fig. 15b

T cell Activation:

THYROxCD3:EGFR

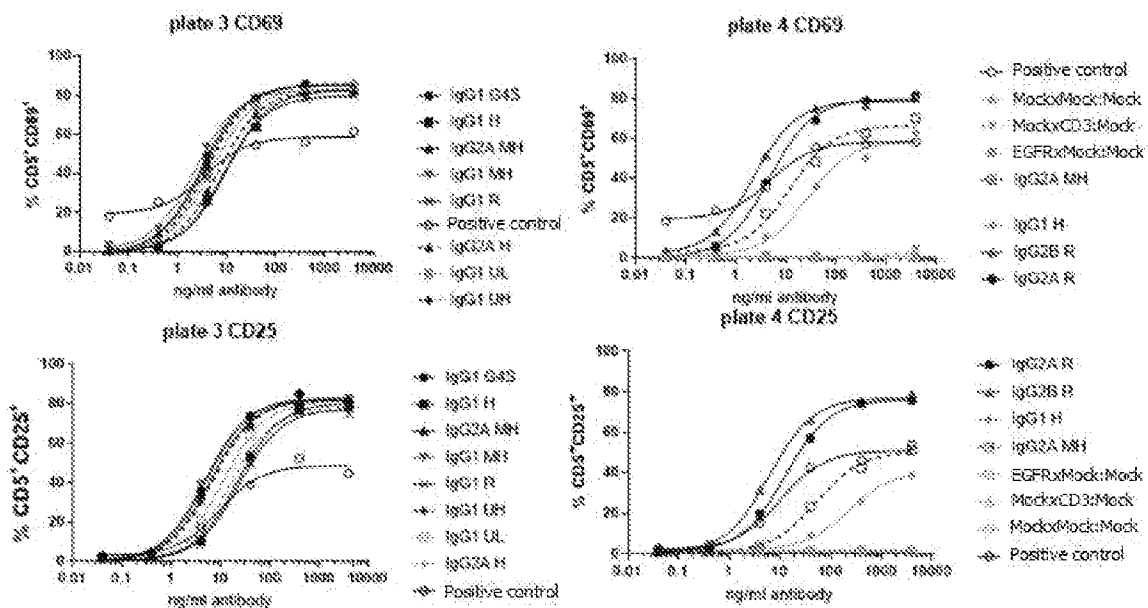


Fig. 16a

## T cell Cytotoxicity:

## EGFRxCD3:TT

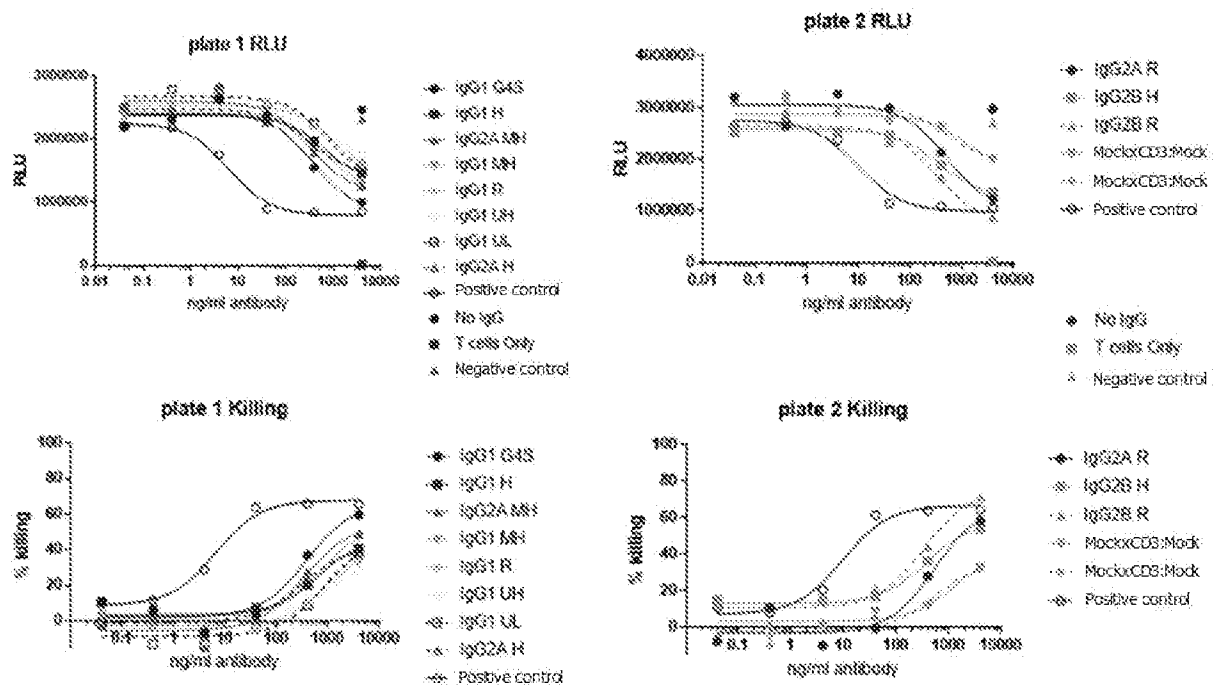
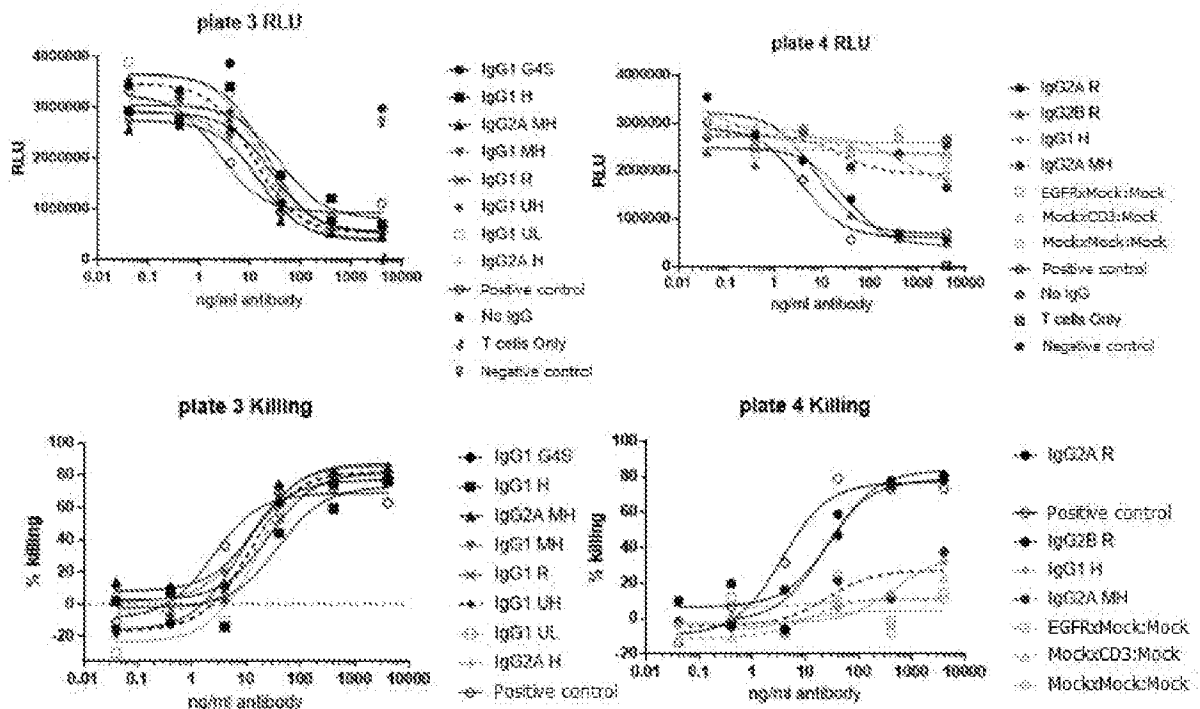


Fig. 16b

## T cell Cytotoxicity:

## THYROxCD3:EGFR



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Fig. 17a

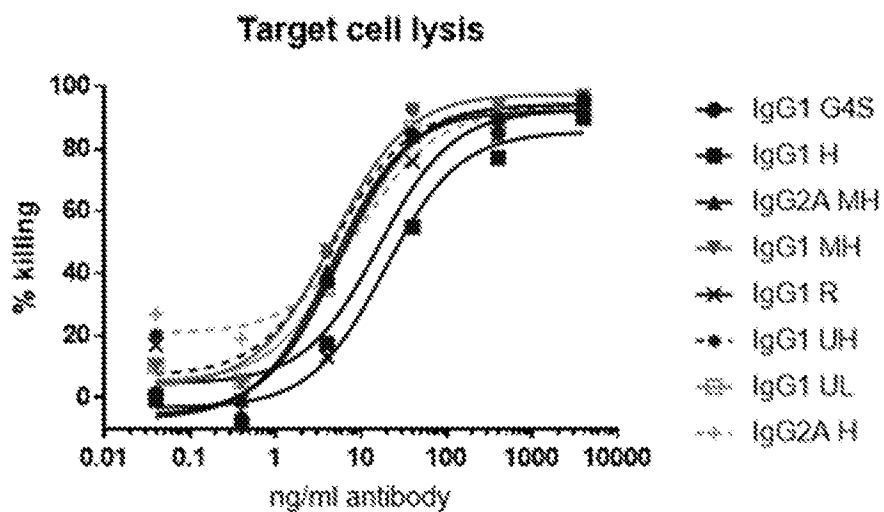


Fig. 17b

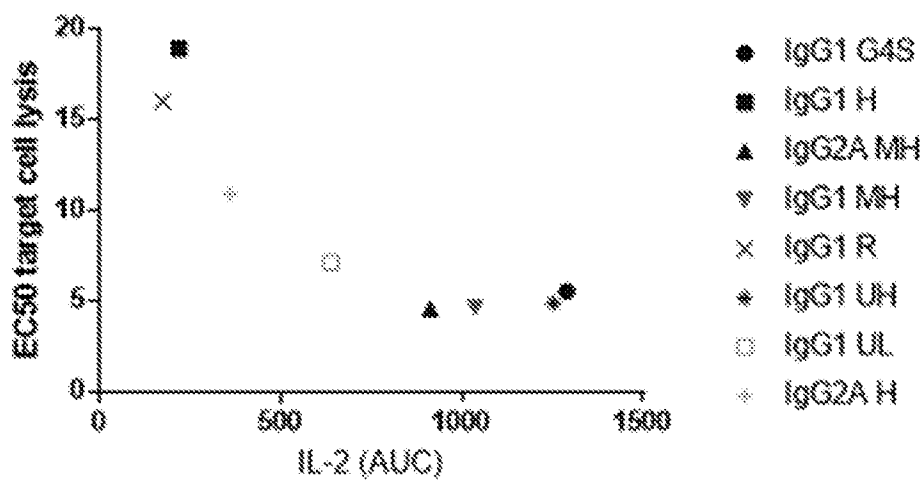


Fig. 17c

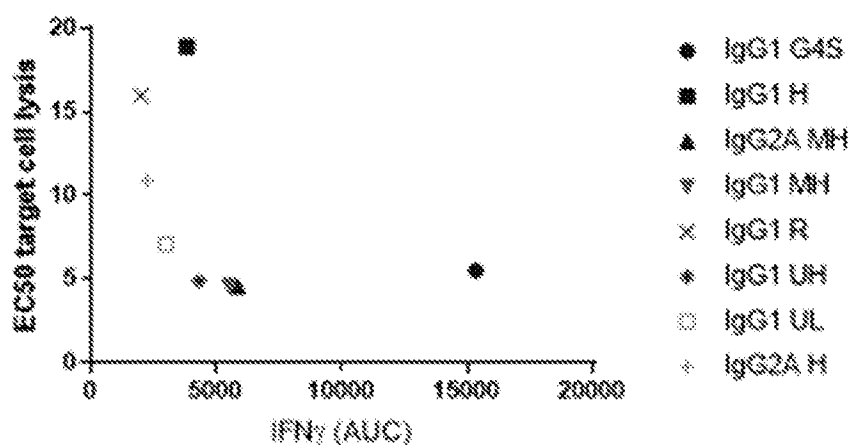
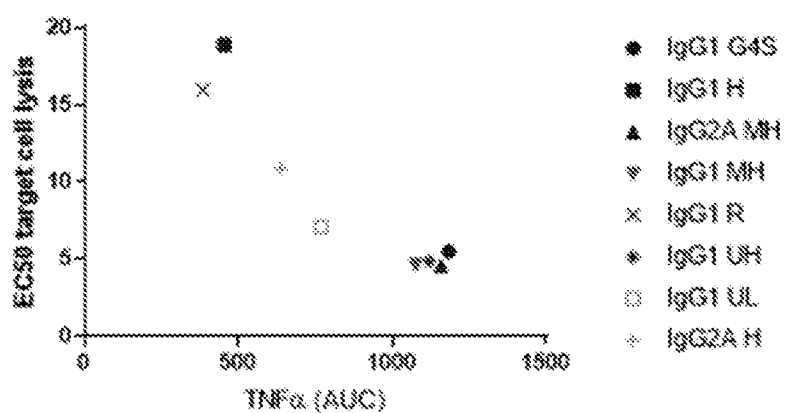
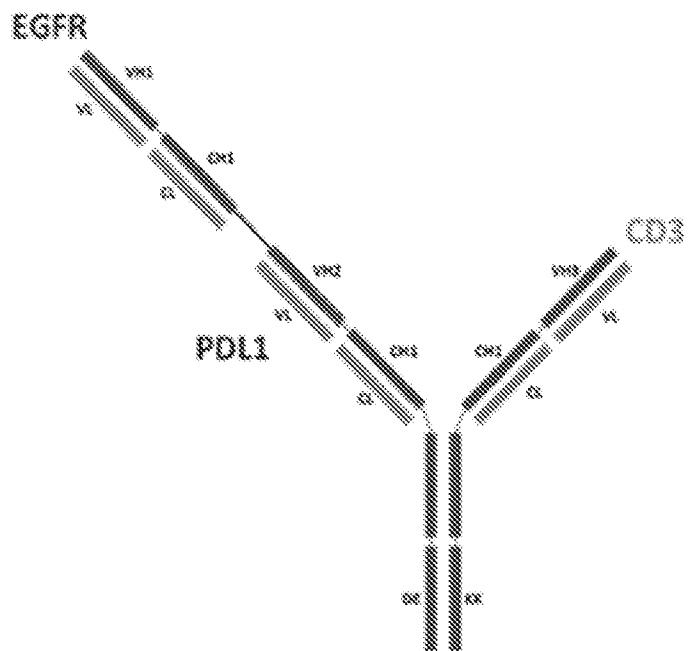


Fig. 17d



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Fig. 18



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Fig. 19

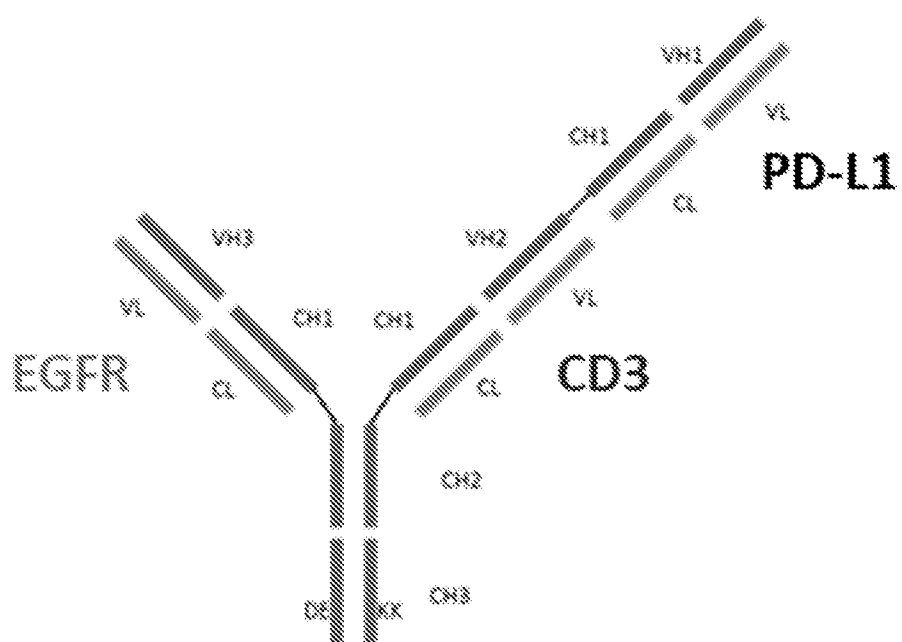


Fig. 20

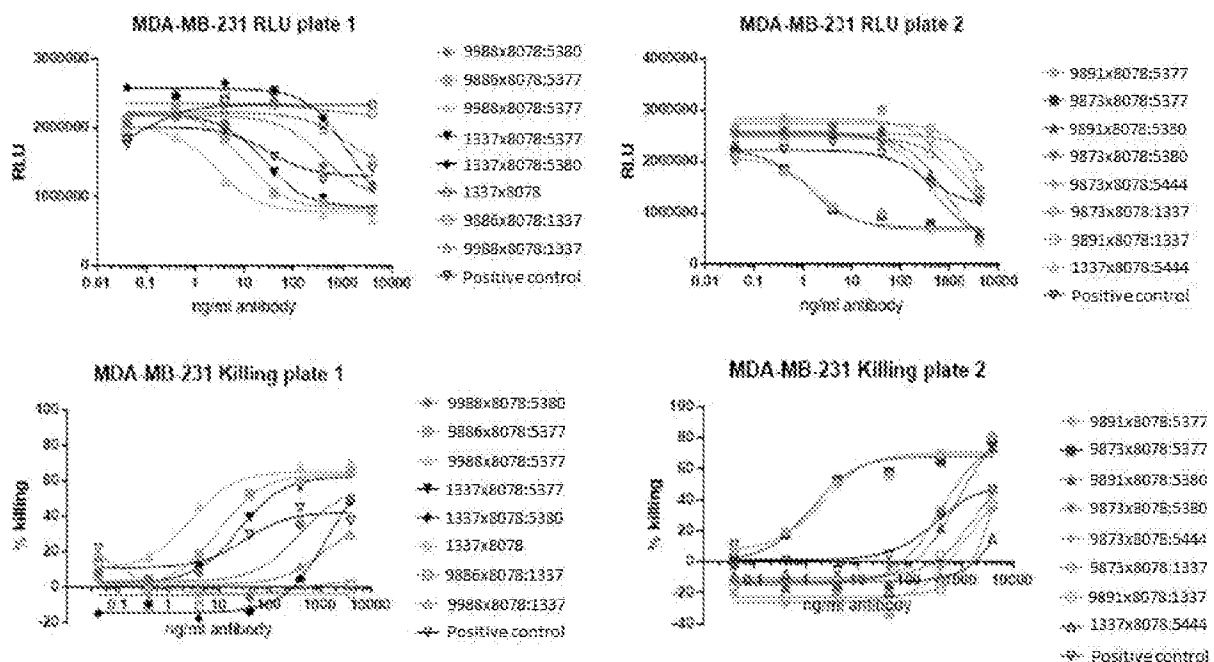


Fig. 21

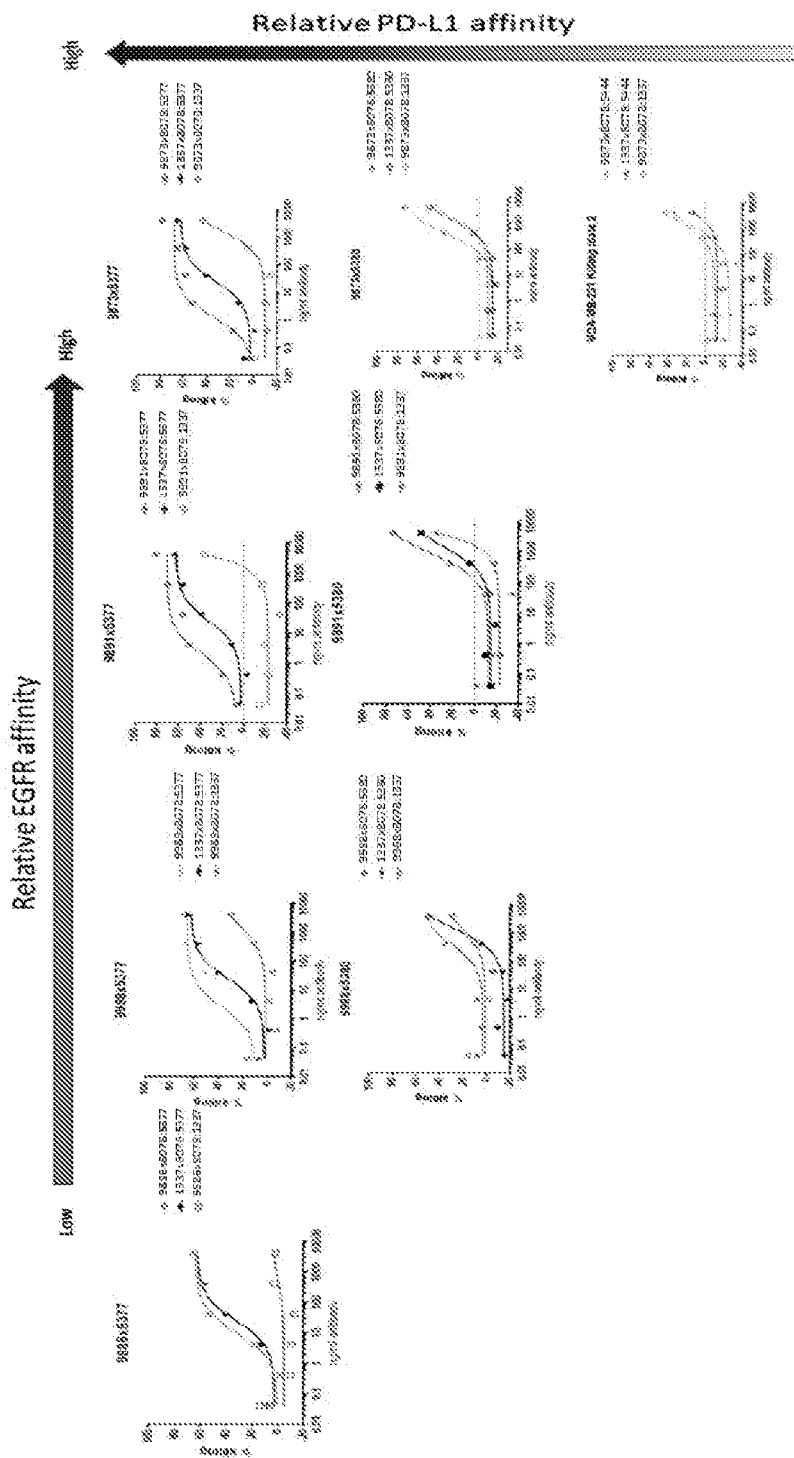


Fig. 22

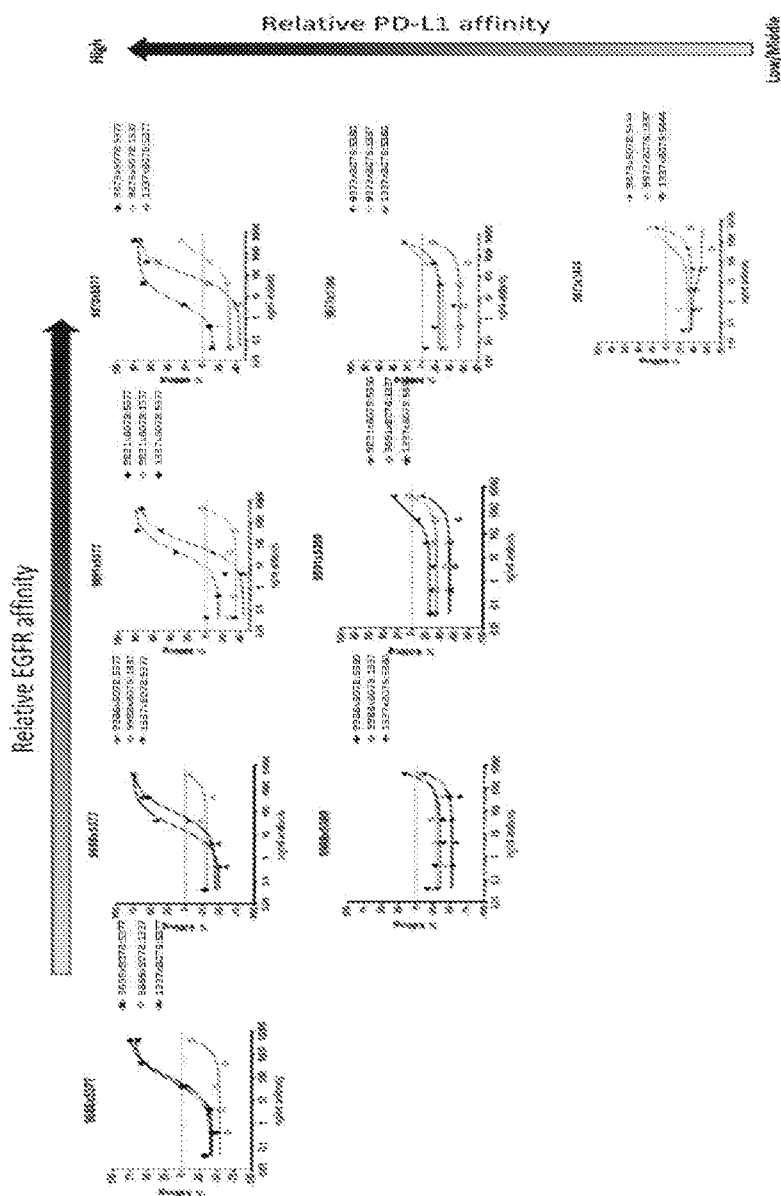
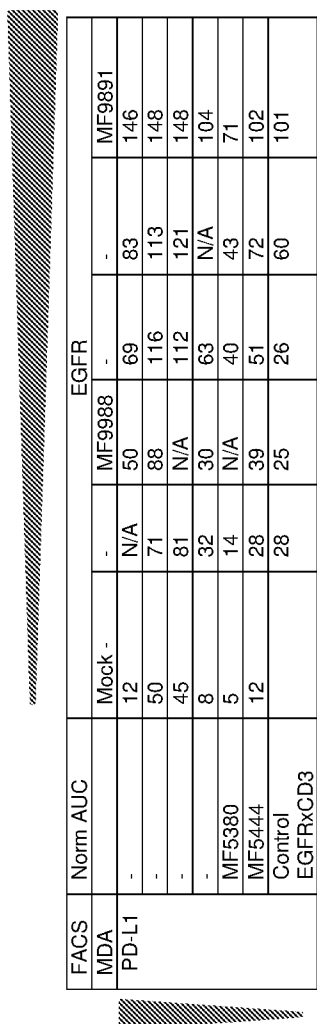


Fig. 23a



FACS	Norm AUC	MDA	Mock -	MF9891	EGFR	MF9891
PD-L1	-	12	N/A	50	69	-
	-	50	71	88	116	113
	-	45	81	N/A	112	121
	-	8	32	30	63	N/A
MF5380	5	14	N/A	40	43	104
MF5444	12	28	39	51	72	71
Control		28	25	26	60	102
EGFRxCD3						101

Fig. 23b

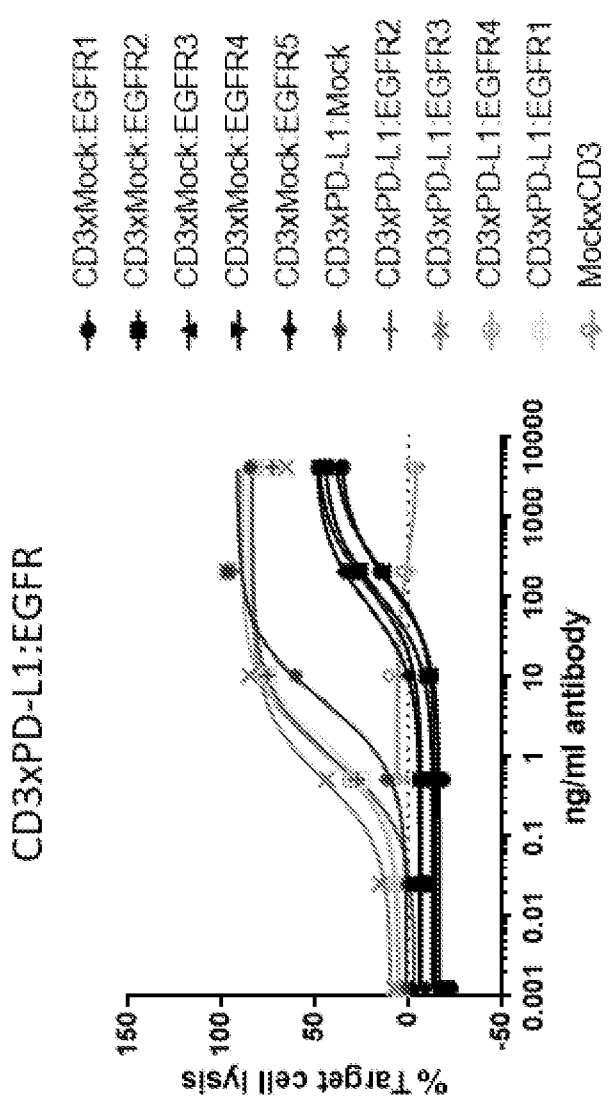


Fig. 24

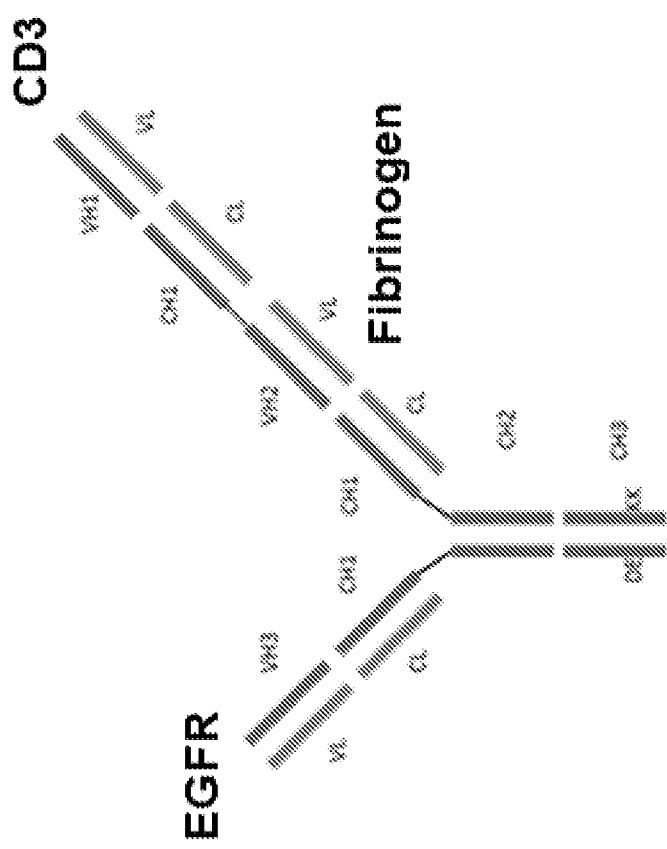


Fig. 25

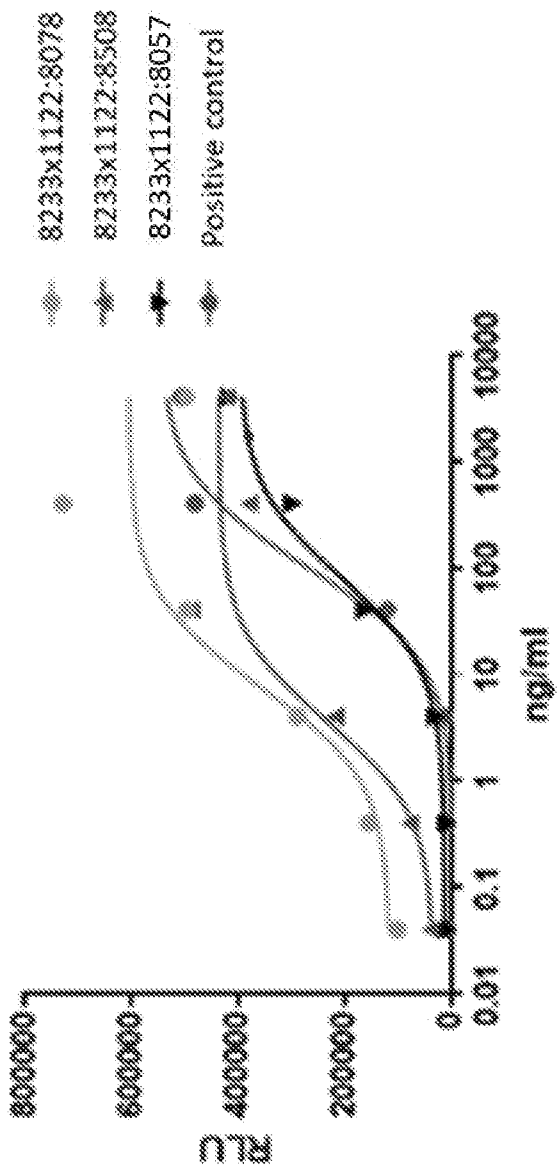
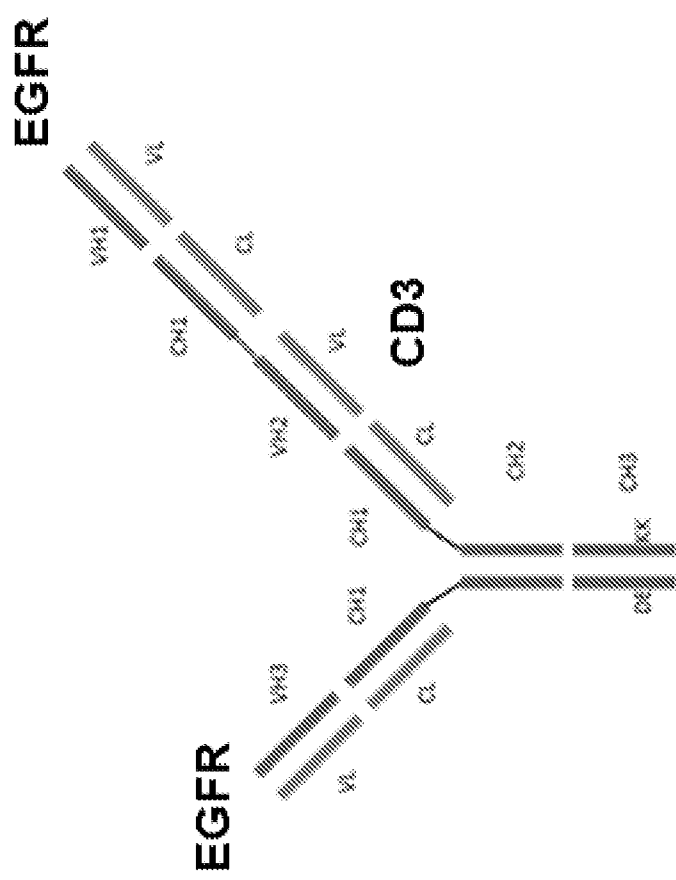


Fig. 26



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Fig. 27

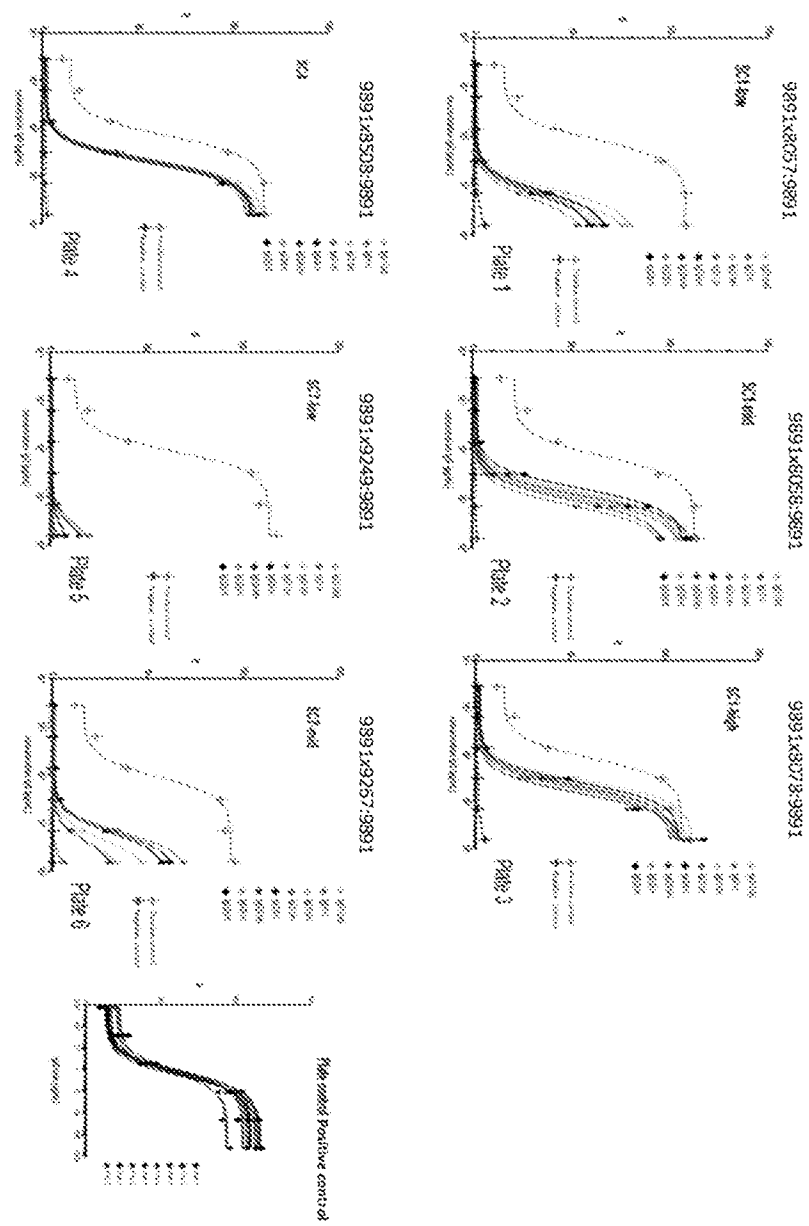


Fig. 28

