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(54) Title: MULTICHAIN MULTITARGETING BISPECIFIC ANTIGEN-BINDING MOLECULES OF INCREASED SELECTIVITY

(57) Abstract: The present invention provides multichain multitargeting bispecific antigen-binding molecules characterized by comprising a first and a second bispecific entity each comprising a domain binding to a target, a second domain binding to an extracellular epitope of the human and the Macaca CD3e chain, wherein both bispecific entities are linked to each other by a spacer which spaces apart the first and the second bispecific entity. Moreover, the invention provides a polynucleotide, encoding the multitargeting bispecific antigen-binding molecule, a vector comprising this polynucleotide, host cells, expressing the construct and a pharmaceutical composition comprising the same.



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MULTICHAIN MULTITARGETING BISPECIFIC ANTIGEN-BINDING MOLECULES OF INCREASED SELECTIVITY

TECHNICAL FIELD

- 5 [1] This invention relates to products and methods of biotechnology, in particular to multichain multitargeting antigen-binding molecules, their preparation and their use.

BACKGROUND

- [2] Redirecting T cell activity against tumor cells by means of bispecific molecules independent of T cell receptor specificity is a further evolving approach in immunooncology (Frankel SR, Baeuerle
10 PA. Targeting T cells to tumor cells using bispecific antibodies. *Curr Opin Chem Biol* 2013;17:385–92). Such new protein-based pharmaceuticals typically can simultaneously bind to two different types of antigen. They are known in several structural formats, and current applications have been explored for cancer immunotherapy and drug delivery (Fan, Gaowei; Wang, Zujian; Hao, Mingju; Li, Jinming (2015). "Bispecific antibodies and their applications". *Journal of Hematology & Oncology*. 8: 130).

- 15 [3] Bispecific molecules useful in immunooncology can be antigen-binding polypeptides such as antibodies, e.g. IgG-like, i.e. full-length bispecific antibodies, or non-IgG-like bispecific antibodies, which are not full-length antigen-binding molecules. Full length bispecific antibodies typically retain the traditional monoclonal antibody (mAb) structure of two Fab arms and one Fc region, except the two Fab sites bind different antigens. Non-full-length bispecific antibodies can lack an Fc region
20 entirely. These include chemically linked Fabs, consisting of only the Fab regions, and various types of bivalent and trivalent single-chain variable fragments (scFvs). There are also fusion proteins mimicking the variable domains of two antibodies. An example of such a format is the bi-specific T-cell engager (BiTE[®]) (Yang, Fa; Wen, Weihong; Qin, Weijun (2016). "Bispecific Antibodies as a Development Platform for New Concepts and Treatment Strategies". *International Journal of*
25 *Molecular Sciences*. 18 (1): 48).

- [4] Exemplary bispecific antibody-derived molecules such as BiTE[®] molecules are recombinant protein constructs made from two flexibly linked antibody derived binding domains. One binding domain of BiTE[®] antigen-binding molecules is specific for a selected tumor-associated surface antigen on target cells; the second binding domain is specific for CD3, a subunit of the T cell receptor
30 complex on T cells. By their particular design, BiTE[®] antigen-binding molecules are uniquely suited to transiently connect T cells with target cells and, at the same time, potentially activate the inherent cytolytic potential of T cells against target cells. An important further development of the first

generation of BiTE[®] antigen-binding molecules (see WO 99/54440 and WO 2005/040220) developed into the clinic as AMG 103 and AMG 110 was the provision of bispecific antigen-binding molecules binding to a context independent epitope at the N-terminus of the CD3 ϵ chain (WO 2008/119567). BiTE[®] antigen-binding molecules binding to this elected epitope do not only show cross-species specificity for the human and the Macaca, or *Callithrix jacchus*, *Saguinus oedipus* or *Saimiri sciureus* CD3 ϵ chain, but also, due to recognizing this specific epitope (instead of previously described epitopes of CD3 binders in bispecific T cell engaging molecules), do not demonstrate unspecific activation of T cells to the same degree as observed for the previous generation of T cell engaging antibodies. This reduction in T cell activation was connected with less or reduced T cell redistribution in patients, the latter being identified as a risk for side effects, e.g. in pasotuximab.

[5] Antibody-based molecules as described in WO 2008/119567 are characterized by rapid clearance from the body; thus, while they are able to reach most parts of the body rapidly, their in vivo applications may be limited by their brief persistence in vivo. On the other hand, their concentration in the body can be adapted and fine-tuned at short notice. Prolonged administration by continuous intravenous infusion is used to achieve therapeutic effects because of the short in vivo half-life of this small, single chain molecule. However, bispecific antigen-binding molecules are available which have more favorable pharmacokinetic properties, including a longer half-life as described in WO 2017/134140. An increased half-life is typically useful in *in vivo* applications of immunoglobulins, especially with respect to antibody fragments or constructs of small size, e.g. in the interest of patient compliance.

[6] One challenging ongoing problem in antibody-based immunooncology is tumor escape. Such tumor escape happens when the immune system -even if triggered or directed by some antibody-based immune-therapeutics- is not capable enough to eradicate tumors, which carry accumulated genetic and epigenetic alterations and use several mechanisms to be the victorious of the immunoediting process (Keshavarz-Fathi, Mahsa; Rezaei, Nima (2019) "Vaccines for Cancer Immunotherapy"). Generally, four mechanisms interfering with effective antitumor immune responses are known: (1) defective tumor antigen processing or presentation, (2) lack of activating mechanisms, (3) inhibitory mechanisms and immunosuppressive state, and (4) resistant tumor cells. Especially with respect to the first mechanism, tumor antigens might be present in a new form due to the genetic instability, mutation of the tumor and escape from immune system. Epitope-negative tumor cells remain hidden and consequently resistant to the immune rejection. They have been developed following the elimination of epitope-positive tumor cells, similar to Darwin's theory of natural selection. In consequence, antibody-based immune-therapy directed against an antigen on tumor cells is rendered ineffective when such tumor cells no longer express a respective antigen due to tumor escape. Said antigen loss is understood herein as driving force for tumor escape and thus, used interchangeably.

Accordingly, there is a need to provide improved antibody-based immunooncology which addresses the problem of antigen loss to effectively prevent tumor escape.

[7] A probably even more pressing challenge to the broad utilization of immunooncology with respect to T-cell engaging bispecific molecules is the availability of suitable targets (Bacac et al., Clin Cancer Res; 22(13) July 1, 2016). For example, solid tumor targets may be overexpressed on tumor cells but expressed at lower, yet significant levels on nonmalignant primary cells in critical tissues. In nature, according to Bacac et al, T cells can distinguish between high- and low-antigen expressing cells by means of relatively low-affinity T cell receptors (TCRs) that can still achieve high-avidity binding to target cells expressing sufficiently high levels of target antigen. T-cell engaging bispecific molecules that could facilitate the same, and thus maximize the window between killing of high- and low-target expressing cells, are thus highly desirable. One approach discussed in the art is the use of dual targeting of two antigens which may lead to improved target selectivity over normal tissues that express only one or low levels of both target antigens. This effect is thought to be dependent on the avidity component mediated by the concurrent binding of the bsAb to both antigens on the same cell. With respect to dual targeting as such, some multispecific monoclonal antibodies (mAb) or other immune constructs are known in the art. WO 2014/116846 teaches a multispecific binding protein comprising a first binding site that specifically binds to a target cell antigen, a second binding site that specifically binds to a cell surface receptor on an immune cell, and a third binding site that specifically binds to cell surface modulator on the immune cell. US 2017/0022274 discloses a trivalent T-cell redirecting complex comprising a bispecific antibody, wherein the bispecific antibody has two binding sites against a tumor-associated antigen (TAA) and one binding site against a T-cell.

[8] However, dual targeting alone as in molecules described above may not be sufficient for efficient target selectivity (Mazor et al, mAbs 7:3, 461-469; May/June 2015). Especially the configuration of the bsAb binding domains, namely monovalent vs. bivalent, is a critical factor. Even more, the provision of a bispecific molecule with several valences alone may not lead to clinically suitable therapeutic as also the potential risk profile in terms of significant immunological side effects such as cytokine release syndrome (CRS) has to be considered. Hence, despite the so-far achieved pre-clinical and clinical success of antibody-based immune-therapeutics, notable limitations remain including differential responses between individuals and cancer types. Not all patients will respond to therapy at available safe doses as dose-limiting toxicity can be a limiting factor for the efficacy of antibody-based immune-therapeutics. Hence, there is also a need to reduce dose-limiting toxicity in antibody-based immune-therapeutics to make such therapy available to more patients suffering from diverse proliferative diseases.

[9] While different multispecific antibodies or antibody fragments are known in the art, some of which address T-cells, no multitargeting bispecific molecules have been proposed before which both

addresses the need of overcoming dose-limiting toxicity in T cell redirecting immune-therapeutics by increasing the therapeutic window and are a stable and ready-to-use therapeutic system.

SUMMARY

[10] In view of the various unmet needs described above, it is an object of the present invention to provide a molecule which comprises at least two polypeptide chains, i.e. a multichain molecule, which molecule is preferably an antigen-binding molecule. The molecule of the present invention is further preferably bispecific, such as a T cell engaging molecule. Further, the molecule of the present invention is preferably multitargeting, e.g. it is typically capable to immune-specifically bind to at least two antigens on a target cell which are typically associated with one or more diseases. It is further preferred that a molecule of the present invention is typically capable to immuno-specifically bind to two antigens on an effector cell at the same time, preferably for use in the treatment of said one or more diseases. Accordingly, the present invention provides a preferably multitargeting bispecific antigen-binding molecule comprising at least one polypeptide, wherein the molecule is characterized by comprising at least five distinctive structural entities, i.e. (i.) a first domain binding to a target cell surface antigen (e.g. a first tumor associated antigen, TAA), (ii.) a second domain binding to an extracellular epitope of the human (and preferably non-human primate, e.g. *Macaca*) CD3 chain, wherein the first binding domain and the second binding domain together form a first bispecific entity, (iii.) a spacer which connects but also sufficiently spaces apart the first bispecific entity from a second bispecific entity comprising (iv.) a third domain binding to the same or preferably a different target cell surface antigen (e.g. a second TAA), and (v.) a fourth domain binding to an extracellular epitope of the human (and preferably non-human primate, e.g. *Macaca*) CD3 chain. Preferably, the domains are (i.) scFv domains comprised of VH and VL domains in amino to carboxyl orientation, respectively, wherein a flexible but short peptide linker links the VL of the first domain to the VH of the second domain and the VL of the third domain to the VH of the fourth domain, respectively, and/or (ii.) Fab domains comprising a first polypeptide monomer comprising a VL and CL domain, and a second polypeptide monomer comprising a VH and CH domain. Surprisingly, a multichain multitargeting bispecific antigen-binding molecule as described herein is typically capable to enable T-cells to distinguish between killing of cells expressing only one or both targets typically associated with a particular disease, respectively, thus opening a therapeutic window and reducing the risk for off-target toxicities and side effects. Moreover, the invention provides a polynucleotide encoding the multitargeting bispecific antigen-binding molecule, a vector comprising this polynucleotide, and host cells expressing the construct and a pharmaceutical composition comprising the same.

[11] In a first aspect, it is envisaged in the context of the present invention to provide a multichain multitargeting bispecific molecule comprising at least two polypeptide chains, wherein the molecule comprises

- (i.) a first binding domain which binds to a first target cell surface antigen (TAA1),

(ii.) a second binding domain which binds to an extracellular epitope of the human and/or the Macaca CD3 chain,

(iii.) a third binding domain which binds to a second target cell surface antigen (TAA2), and

5 (iv.) a fourth binding domain which binds to an extracellular epitope of the human and/or the Macaca CD3 chain,

wherein the first binding domain and the second binding domain form a first bispecific entity, and the third and the fourth binding domain form a second bispecific entity, and

wherein the molecule further comprises a spacer entity selected from

10 (1.) a dimerizing domain selected from

(a.) an Fc domain comprising a first and a second polypeptide monomer comprising a hinge, a CH2 domain and a CH3 domain, respectively, wherein the first and second polypeptide monomer form a heterodimer; wherein the heterodimer is formed by

15 - charged pair mutations selected from (i.) D399K, K409D, K392D, and E356K, (ii.) D399K, K409D, K392D, E357K, K370D, and E356K, (iii.) D399K, K409D, K392D, E356K and K439D, (iv.) D399K, K409D, and K392D, (v.) D399K, K409D, K392D, E357K and E370K, (vi.) D399K, K409D, K392D, E357K, K370E and K360E, (vii.) D399K, K409D, K392D, E357K, K370E, E356K, and K439E, and (viii.) D399K, K409D, K392D, E357K, K370E, K360E, E356K, and K439D,
20 preferably comprising a K392D, K409D and/or K439D mutation in the CH3 domain of the first polypeptide monomer and comprising a E356K and/or D399K mutation in the CH3 domain of the second polypeptide monomer, wherein the positions are according to EU numbering; or

25 - knobs-into-holes mutations comprising preferably a T366S, L368A and Y407V mutation in the first polypeptide monomer and a T366W mutation in the second monomer, wherein the positions are according to EU numbering;

(b.) a human serum albumin (HSA) domain comprising a first and a second polypeptide monomer, wherein the first and the second polypeptide monomer correspond to an HSA subdomain, respectively, wherein the first and second polypeptide monomer form a native HSA-like heterodimer; and
30

(c.) a Fab comprising a first and a second polypeptide monomer, wherein the first polypeptide monomer comprises a VL and CL domain and the second polypeptide monomer comprises a VH and CH1 domain, wherein the CL and CH1 domains are linked by a disulfide bridge;

5 wherein the dimerizing domain comprises two N-termini and two C-termini, respectively, whereof at least one N-terminus and one C-terminus, respectively, is linked to a bispecific entity, wherein any of the first, second, third and fourth domain can be selected from any form of binding domain, preferably selected from Fab and single chain domain, the single chain domain preferably selected from single chain Fv
10 (scFv) and scFab;

(2.) a single chain domain selected from ubiquitin , beta 2 microglobulin , , VH-only domain , PSI domain from Met-receptor, Fibronectin type III domain from tenascin , Granulocyte-macrophage colony-stimulating factor (GM-CSF) , interleukin-4 , CD137L Ectodomain , Interleukin-2 , PD-1 binding domain from human Programmed
15 cell death 1 ligand 1 (PDL1) , Tim-3 (AS 24-130), MiniSOG , a programmed cell death protein 1 (PD1) domain, human serum albumin (HSA), or a single chain Fc (scFc) domain comprising two polypeptide monomers comprising each a hinge, a CH2 and a CH3 domain a hinge and a further CH2 and a CH3 domain, wherein said two polypeptide monomers are fused to each other via a peptide linker,

20 wherein the single chain domain comprises one N-terminus and one C-terminus, which are, respectively, linked to a bispecific entity, wherein at least one of the first, second, third and fourth binding domain is a two-chained Fab, and any of the remaining up to at least three binding domains can be selected from any form of binding domain, is preferably selected from Fab and single chain domain, the single chain domain
25 preferably selected from scFv and, scFab;

wherein the distance between the C alpha atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus of the spacer entity are spaced apart by at least 30 Å, wherein the spacer entity spaces apart the first and the second bispecific entity by at least a distance of about 50 Å, wherein the indicated distance is preferably understood as the distance between centers of
30 mass (i.) of the first and the third binding domain or (ii.) the first and the second bispecific entity, and which spacer entity is positioned between the first and the second bispecific entity.

[12] Within said aspect, it is also envisaged in the context of the present invention to provide a multichain multitargeting bispecific antigen-binding molecule, wherein when the spacer is a single

chain domain, the arrangement of the binding domains in an amino to carboxyl order is selected from the group consisting of

- (i.) first and second domain, spacer, third and fourth domain
- (ii.) first and second domain, spacer, fourth and third domain
- 5 (iii.) second and the first domain, spacer, third and fourth domain, and
- (iv.) second and first domain, spacer, fourth and third domain.

[13] Within said aspect, it is also envisaged in the context of the present invention to provide a multichain multitargeting bispecific antigen-binding molecule, wherein when the spacer is a single chain domain, the arrangement of the binding domains in an amino to carboxyl order is selected from
10 the group consisting of

- (i.) first domain in the format of Fab, second domain in the format of scFv, spacer, third domain in the format of Fab and fourth domain in the format of scFv (e.g. Fig. 3B);
- (ii.) first domain in the format of Fab, second domain in the format of Fab, spacer, third domain in the format of Fab and fourth domain in the format of Fab (e.g. Fig. 3D);
- 15 (iii.) first domain in the format of scFv, second domain in the format of Fab, spacer, third in the format of scFv and fourth domain in the format of Fab (e.g. Fig. 3H);
- (iv.) first domain in the format of scFv, second domain in the format of scFv, spacer, third in the format of scFv and fourth domain in the format of Fab;
- (v.) first domain in the format of scFv, second domain in the format of scFv, spacer, third in the
20 format of Fab and fourth domain in the format of scFv;
- (vi.) first domain in the format of Fab, second domain in the format of scFv, spacer, third in the format of scFv and fourth domain in the format of scFv; and
- (vii.) first domain in the format of scFv, second domain in the format of Fab, spacer, third in the format of scFv and fourth domain in the format of scFv,
- 25 wherein each scFv comprises in an amino to carboxyl order VH, linker and VL or VL, linker and VH, preferably VH, linker and VL.

[14] Within said aspect, it is also envisaged in the context of the present invention to provide a multichain multitargeting bispecific antigen-binding molecule, wherein when the spacer is the spacer

is a dimerizing domain, the arrangement of the binding domains in an amino to carboxyl order is selected from the group consisting of

- 5 (i.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, second domain in the format of scFv, first polypeptide monomer of the spacer dimerizing domain, and a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain, and the fourth domain in the format of a scFv (e.g. Fig. 3A);
- 10 (ii.) first domain in the format of Fab, second domain in the format of Fab, spacer, third domain in the format of Fab and fourth domain in the format of Fab (e.g. Fig. 3C);
- (iii.) a first chain comprising the second domain in the format of scFv, the VH and CH1 of the first domain, forming a Fab together with the second chain, first polypeptide monomer of the spacer dimerizing domain, a second chain comprising the VL and CL of the first domain, a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain, and the fourth domain in the format of a scFv (e.g. Fig. 3E);
- 15 (iv.) a first chain comprising the second domain in the format of scFv, the VH and CH1 of the first domain, forming a Fab together with the second chain, first polypeptide monomer of the spacer dimerizing domain, a second chain comprising the VL and CL of the first domain, a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the fourth domain in the format of a scFv, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain (e.g. Fig. 3F);
- 20 (v.) a first chain comprising the second domain in the format of scFv, the VH and CH1 of the first domain, forming a Fab together with the second chain, first polypeptide monomer of the spacer dimerizing domain, a second chain comprising the VL and CL of the first domain, a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the fourth domain in the format of a scFv, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain (e.g. Fig. 3G);
- 25 30

wherein each scFv comprises in an amino to carboxyl order VH, linker and VL or VL, linker and VH, preferably VH, linker and VL.

[15] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein when the spacer is a dimerizing domain, the arrangement of the binding domains is in an amino to carboxyl order and is selected from the group consisting of

- 5 (i.) a first chain comprising the first domain in the format of scFv, first polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv and a second chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, fourth domain in the format of scFv (e.g. Fig 2A);
- 10 (ii.) a first chain comprising the first domain in the format of scFv, first polypeptide monomer of the spacer dimerizing domain, second domain in the format of scFv and a second chain comprising the third domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, fourth domain in the format of scFv (e.g. Fig. 2B);
- 15 (iii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv, and a third chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, fourth domain in the format of scFv (e.g. Fig. 2C);
- 20 (iv.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, second domain in the format of scFv, and a third chain comprising the fourth domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv (e.g. Fig 2D);
- 25 (v.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, and a third chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv, and fourth domain in the format of scFv (e.g. Fig. 2E);
- 30 (vi.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv, and fourth domain in the format of scFv, and a third chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain (e.g. Fig 2F);
- (vii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of

the spacer dimerizing domain, VH and CH1 of the third domain forming a Fab together with the third chain, a third chain comprising the VL and CL of the third domain, a fourth chain comprising the VH and CH1 of the second domain, forming a Fab together with the fifth chain, second polypeptide monomer of the spacer dimerizing domain, VH and CH1 of the third domain forming a Fab together with the sixth chain, a fifth chain comprising the VL and CL of the second domain, and a sixth chain comprising the VL and CL of the fourth domain (e.g. Fig. 2G);

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(viii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, second domain in the format of scFv, a third chain comprising the fourth domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, VH and CH1 of the third domain, forming a Fab together with the fourth chain, and a fourth chain comprising the VL and CL of the third domain (see e.g. Fig. 2H);

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(ix.) a first chain comprising the first domain in the format of a scFv, the first polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the third chain, the second polypeptide monomer of the spacer dimerizing domain, the VH and CH1 of the fourth domain, forming a Fab together with the fourth chain, a third chain comprising the VL and CL of the second domain, and a fourth chain comprising the VL and CL of the fourth domain (e.g. Fig. 2I);

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(x.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing domain, the fourth domain in the format of a scFv, and a fourth chain comprising the VL and CL of the second domain (e.g. Fig. 2J);

20

(xi.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, the second domain in the format of scFv, a third chain comprising the VH and CH1 of the fourth domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing domain, the third domain in the format of a scFv, and a fourth chain comprising the VL and CL of the fourth domain (e.g. Fig. 2K);

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(xii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing

domain, the third domain in the format of scFv, the fourth domain in the format of a scFv, and a fourth chain comprising the VL and CL of the second domain (e.g. Fig. 2L);

(xiii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, the fourth domain in the format of a scFv, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing domain, and a fourth chain comprising the VL and CL of the second domain (e.g. Fig. 2M);

wherein each scFv comprises in a N to C orientation VH, linker and VL or VL, linker and VH, preferably VH, linker and VL.

[16] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the spacer entity is a globular protein, wherein the distance between the C alpha atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus are spaced apart by at least 20 Å, preferably at least 30 Å, more preferably at least 50 Å, in order to effectively space apart the first and the second bispecific entity by preferably at least 50 Å.

[17] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein said spacer entity, when the spacer is single-chained, which effectively spaces apart the first and the second bispecific entity is selected from a group consisting of ubiquitin, beta 2 microglobulin, SAND domain, Green fluorescent protein (GFP), VHH antibody lama domain, PSI domain from Met-receptor, Fibronectin type III domain from tenascin, Granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-4, CD137L Ectodomain, Interleukin-2, PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1), Tim-3 (AS 24-130), MiniSOG, a programmed cell death protein 1 (PD1) domain, human serum albumin (HSA) or a derivate of any of the foregoing spacer entities, a multimer of a rigid linker, and a Fc domain or dimer or trimer thereof, each Fc domain comprising two polypeptide monomers comprising each a hinge, a CH2 and a CH3 domain a hinge and a further CH2 and a CH3 domain, wherein said two polypeptide monomers are fused to each other via a peptide linker or wherein the two polypeptide monomers are linked together by non-covalent CH3-CDH3 interactions and/or covalent disulfide bonds to form a heterodimer.

[18] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein said spacer entity when single chained is at least one Fc domain, preferably one domain or two or three covalently linked domains, which or each of which comprises in an amino to carboxyl order:

hinge-CH2-CH3-linker-hinge-CH2-CH3.

[19] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein each of said polypeptide monomers in the spacer entity has an amino acid sequence that is at least 90% identical to a sequence selected from the group consisting of:
5 SEQ ID NO: 17-24, wherein preferably each of said polypeptide monomers has an amino acid sequence selected from SEQ ID NO: 17-24.

[20] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the CH2 domain in the spacer comprises an intra domain cysteine disulfide bridge.

10 [21] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the single chain spacer entity comprises an amino acid sequence selected the group consisting of SEQ ID NO: 13 and 15 to 16 and 25 to 34, ubiquitin (SEQ ID NO: 1081), beta 2 microglobulin (SEQ ID NO: 1083), SAND domain (SEQ ID NO: 1084), Green fluorescent protein (GFP) (SEQ ID NO: 1085), VHH antibody lama domain (SEQ ID NO: 1086), PSI
15 domain from Met-receptor (SEQ ID NO: 1087), Fibronectin type III domain from tenascin (SEQ ID NO: 1088), Granulocyte-macrophage colony-stimulating factor (GM-CSF) (SEQ ID NO: 1089), interleukin-4 (SEQ ID NO: 1090), CD137L Ectodomain (SEQ ID NO: 1091), Interleukin-2 (SEQ ID NO: 1092), PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1) (SEQ ID NO: 1093), Tim-3 (AS 24-130) (SEQ ID NO: 1094), MiniSOG (SEQ ID NO: 1095), a programmed
20 cell death protein 1 (PD1) domain (SEQ ID NO: 16), human serum albumin (has, SEQ ID NO: 15) or an amino acid with at least 90%, preferably 95% or even 98% sequence identity thereof, preferably scFc (SEQ ID NO: 25).

[22] Within said aspect it is also envisaged that the first peptide monomer of the first peptide chain
25 is SEQ ID NO 35 and the second peptide monomer of the second peptide chain is SEQ ID NO 36, wherein the two peptide monomers preferably form a heterodimer.

[23] Within said aspect it is also envisaged that the antigen-binding molecule is characterized by

- (i) the first and third domain comprise two antibody-derived variable domains and the second and the fourth domain comprises two antibody-derived variable domains;
- 30 (ii) the first and third domain comprise one antibody-derived variable domain and the second and the fourth domain comprises two antibody-derived variable domains;

(iii) the first and third domain comprise two antibody-derived variable domains and the second and the fourth domain comprises one antibody-derived variable domain; or

(iv) the first domain comprises one antibody-derived variable domain and the third domain comprises one antibody-derived variable domain.

5 [24] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule comprising two polypeptide chains, wherein

the first polypeptide chain comprises a VH of the first domain, a VH second domain, the first polypeptide monomer comprising preferably a hinge, a CH2 and a CH3 domain, a VH of the third domain, and a VH of the fourth domain; and

10 the second polypeptide chain comprises a VL of the first domain, a VL second domain, the first polypeptide monomer comprising preferably a hinge, a CH2 and a CH3 domain, a VL of the third domain, and a VL of the fourth domain,

wherein preferably the first and second polypeptide monomer form a heterodimer, thereby connecting the first and the second polypeptide chain.

15 [25] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first, second, third and fourth binding domain each comprise in an amino to carboxyl order a VH domain and a VL domain, wherein the VH and VL within each domain is connected by a peptide linker, preferably a flexible linker which comprises serine, glutamine and/or glycine as amino acid building blocks, preferably only serine (Ser, S) or glutamine
20 (Gln, Q) and glycine (Gly, G), more preferably (G4S)_n or (G4Q)_n, even more preferably SEQ ID NO: 1 or 3.

[26] Within said aspect, it is also envisaged in the context of the present invention to provide peptide linker, wherein the peptide linker comprises or consists of S(G4X)_n and (G4X)_n, wherein X is selected from the group consisting of Q, T, N, C, G, A, V, I, L, and M, and wherein n is an integer
25 selected from integers 1 to 20, preferably wherein n is 1, 2, 3, 4, 5 or 6, preferably wherein X is Q, wherein preferably the peptide linker is (G4X)_n, n is 3, and X is Q.

[27] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the peptide linker between the first binding domain and the second binding domain and the third binding domain and the fourth binding domain is preferably a flexible
30 linker which comprises serine, glutamine and/or glycine or glutamic acid, alanine and lysine as amino acid building blocks, preferably selected from the group consisting of SEQ ID NO: 1 to 4, 6 to 12 and 1125.

[28] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the peptide linker between the first binding domain or the second binding domain and the spacer, and/or the third binding domain and the fourth binding domain and the spacer, respectively, is preferably a short linker rich in small and/or hydrophilic amino acids, preferably glycine and preferably SEQ ID NO: 5.

[29] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein any of the first target cell surface antigen and the second target cell surface antigen is selected from the group consisting of CS1, BCMA, CDH3, FLT3, CD123, CD20, CD22, EpCAM, MSLN and CLL1.

[30] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first target cell surface antigen and the second target cell surface antigen are not identical.

[31] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first target cell surface antigen and the second target cell surface antigen are identical.

[32] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first binding domains is capable of binding to the first target cell surface antigen and the third binding domain is capable of binding to the second target cell surface antigen simultaneously, preferably wherein the first target cell surface antigen and the second target cell surface antigen are on the same target cell.

[33] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule of claim 1, wherein the first target cell surface antigen and the second target cell surface antigen, respectively, are selected from the group consisting of CS1 and BCMA, BCMA and CS1, FLT3 and CD123, CD123 and FLT3, CD20 and CD22, CD22 and CD20, EpCAM and MSLN, MSLN and EpCAM, MSLN and CDH3, CDH3 and MSLN, FLT3 and CLL1, and CLL1 and FLT3.

[34] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule of claim 1, wherein the first target cell surface antigen and/or the second target cell surface antigen is human MSLN (selected from SEQ ID NOs: 1181, 1182 and 1183), and wherein the first and/or third binding domain of the antigen-binding molecule of the invention binds to human MSLN epitope cluster E1 (SEQ ID NO: 1175, aa 296-346 position according to Kabat) as determined by murine chimere sequence analysis as described herein, but preferably not to human MSLN epitope cluster E2 (SEQ ID NO: 1176, aa 247-384 position according to Kabat), E3 (SEQ ID

NO: 1177, aa 385-453 position according to Kabat), E4 (SEQ ID NO: 1178, aa 454-501 position according to Kabat) and/or E5 (SEQ ID NO: 1179 aa 502-545 position according to Kabat).

5 [35] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule of claim 1, wherein the first target cell surface antigen and/or the second target cell surface antigen is human CDH3 (SEQ ID NOs: 1170), and wherein the first and/or third binding domain of the antigen-binding molecule of claim 1 binds to human CDH3 epitope cluster D2B (SEQ ID NO: 1171, aa 253-290 position according to Kabat), D2C (SEQ ID NO: 1172 aa 291-327 position according to Kabat), D3A (SEQ ID NO: 1173 aa 328-363 position according to Kabat) and D4B (SEQ ID NO: 1174, aa 476-511 position according to Kabat), preferably D4B (SEQ ID NO: 10 1174, aa 476-511 position according to Kabat), as determined by murine chimere sequence analysis as described herein.

[36] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and the fourth binding domain (CD3 binding domains) both have (i.) an affinity lower than characterized by a KD value of about 1.2×10^{-8} M measured by surface plasmon resonance (SPR), or (ii.) an affinity characterized by a KD value of about 1.2×10^{-8} M measured by SPR. 15

[37] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and the fourth binding domain (CD3 binding domains) have an affinity characterized by a KD value of about 1.0×10^{-7} to 5.0×10^{-6} M measured by SPR, preferably about 1.0 to 3.0×10^{-6} M, more preferably about 2.5×10^{-6} M measured by SPR. 20

[38] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and the fourth binding domain (CD3 binding domains) have an affinity characterized by a KD value of about 1.0×10^{-7} to 5.0×10^{-6} M measured by SPR, preferably about 1.0 to 3.0×10^{-6} M, more preferably about 2.5×10^{-6} M measured by SPR.

25 [39] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein each of the second and the fourth binding domain (CD3 binding domains) individually has an at least about 10-fold, preferably at least about 50-fold or more preferably at least about 100-fold lower activity than one CD3 binding domain comprising a VH according to SEQ ID NO 43 and a VL according to SEQ ID NO 44 (i.e. in a mono targeting context in contrast to a dual targeting context). 30

[40] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and the fourth domain are effector binding domains binding to CD3 ϵ chain which comprise or consist of a VH region linked to a VL region, wherein

i) the VH region comprises:

a CDR-H1 sequence of X1YAX2N, where X1 is K, V, S, G, R, T, or I; and X2 is M or I;

a CDR-H2 sequence of RIRSKYNNYATYYADX1VKX2, where X1 is S or Q; and X2 is D, G, K, S, or E; and

5 a CDR-H3 sequence of HX1NFGNSYX2SX3X4AY, where X1 is G, R, or A; X2 is I, L, V, or T; X3 is Y, W or F; and X4 is W, F or Y; and

ii) wherein the VL region comprises:

a CDR-L1 sequence of X1SSTGAVTX2X3X4YX5N, where X1 is G, R, or A; X2 is S or T; X3 is G or S; X4 is N or Y; and X5 is P or A;

10 a CDR-L2 sequence of X1TX2X3X4X5X6; where X1 is G or A; X2 is K, D, or N; X3 is F, M or K; X4 is L or R; X5 is A, P, or V; and X6 is P or S; and

a CDR-L3 sequence of X1LWYSNX2WV, where X1 is V, A, or T; and X2 is R or L; and

iii) wherein one or more of CDR sequences of the VH region of i) and/or of the VL region of ii) comprise one amino acid substitution or a combination thereof selected from X24V and X24F in
15 CDR-H1;

D15, and X116A in CDR-H2;

H1, X12E, F4, and N6 in CDR-H3; and

X11L and W3 in CDR-L3.

20 **[41]** Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and the fourth binding domain comprise a VH region comprising CDR-H 1, CDR-H2 and CDR-H3 selected from SEQ ID NOs 37 to 39, 45 to 47, 53 to 55, 61 to 63, 69 to 71, 436 to 438, 1126 to 1128, 1136 to 1138, 1142 to 1144, 1148 to 1150, and 1217 to 1219 and a VL region comprising CDR-L1, CDR-L2 and CDR-L3 selected from SEQ ID NOs 40 to
25 42, 48 to 50, 56 to 58, 64 to 66, 72 to 74, 439 to 441, 1129 to 1131, 1139 to 1141, 1145 to 1147, 1151 to 1153, and 1220 to 1222, preferably 61 to 63 and 64 to 66 or 1217 to 1219 and 1220 to 1222.

[42] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and fourth binding domain comprise a VH region selected from SEQ ID NOs 43, 51, 59, 67, 75, 442, 1132 and 1223, preferably 67 or 1223.

30 **[43]** Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and fourth binding domain comprise a VL region selected from SEQ ID NOs 44, 52, 60, 68, 76, 443 1133 and 1224, preferably 68 or 1224.

[44] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the second and fourth binding domain comprising a VH region

selected from SEQ ID NOs 43, 51, 59, 67, 75, 442, 1132 and 1223, preferably 67, and a VL region selected from SEQ ID NOs 44, 52, 60, 68, 76, 443, 1133 and 1224, preferably 68, wherein when the VH region is 1132 and the VL region is 1133, the second and/or fourth binding domain as scFab domain additionally comprises a CH1 domain of SEQ ID NO: 1134 and a CLK domain of SEQ ID NO: 1135, and wherein the VH and VL region are linked to each other by a linker preferably selected from SEQ ID NO 1, 3 and 1125, or wherein the VH of the VH-CH1 of the second and forth domain is SEQ ID NO is SEQ ID NO 1223, and the CH1 of the VH-CH1 of the second and forth domain is SEQ ID NO is SEQ ID NO 1224, and the VL of the VL-CL of the second and forth domain is SEQ ID NO is SEQ ID NO 1225, and the CL of the VL-L of the second and forth domain is SEQ ID NO is SEQ ID NO 1226. Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or the third (target) binding domain bind to CDH3 and comprise a VH region comprising SEQ ID NO: 1154 as CDR-H 1 wherein X1 (the number behind the "X" indicates the numerical order of the "X" in respective amino acid sequence in N- to C-orientation in the sequence table) is S or N, X2 is Y or S, X3 is P or W, X4 is I or M and X5 is Y, N or H; SEQ ID NO: 1155 as CDR-H2 wherein X1 is K, V, N or R; X2 is A, D, R, Y, S, W or H; X3 is Y, S, P, G or T; X4 is S, G or K; X5 is A, V, D, K, G, or T; X6 is A, V, D, K, S, G or H; X7 is Y, G, or E; X8 is K, I, or N; X9 is A, S, or N; X10 is S, Q or G; X11 is S or K; X12 is F or V; and X13 is K or Q; and SEQ ID NO: 1156 as CDR-H3, wherein X1 is F or Q; X2 is R,K,S or W; X3 is G or D; X4 is Y, P or R; X5 is R, S, G, N or T; X6 is Y,A or H; X7 is F, L or M; X8 is A or V; and X9 is Y or V; and wherein the first and/or the third (target) binding domain bind to CDH3 and comprise a VL region comprising SEQ ID NO: 1158 as CDR-L 1 wherein X1 is K or R, X2 is A or S; X3 is Q,D,S,G or E; X4 is S,D or N; X5 is V,L or I; X6 is ,K,Y,S,or H; X7 is S or N; X8 is F,L or M; and X9 is A,N or H; SEQ ID NO: 1159 as CDR-L 2 wherein X1 is Y,G,W,N; X2 is T or A; X3 is S or K; X4 is T,N or R; X5 is L or R; X6 is E,A,V or H; and X7 is S or E; and SEQ ID NO: 1160 as CDR-L3 wherein X1 is Q or V; X2 is Q,N or H; X3 is F,L,Y,W,N, or H; X4 is A,D,Y,S or N; X5 is Q,R,S,G,W or M; X6 is T,Y or F; and X7 is F,Y or L.

[45] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or the third (target) binding domain bind to MSLN and comprise a VH region comprising SEQ ID NO: 1162 as CDR-H 1 wherein X1 (the number behind the "X" indicates the numerical order of the "X" in respective amino acid sequence in N- to C-orientation in the sequence table) is S,G or D; X2 is Y,A,G or F; X3 is I,W, or M; and X4 is V,S,G,T, or H; SEQ ID NO: 1163 as CDR-H 2 wherein X1 is A,S,N,W,Y,or V; X2 is Y,S or N; X3 is Y,G,P, or S; X4 is D,H,S, or N; X5 is G or S; X6 is E,G or S; X7 is G,S,N,F,T or Q; X8 is S,W,K,D,I or T; X9 is Y or N; X10 is A or N; X11 is A,P,N,D,E,I or Q; X12 is D,A,S or K; X13 is V,L, or F; X14 is K or Q; and X15 is G or S; and SEQ ID NO: 1164 as CDR-H 3 wherein X1 is D,E or V; X2 is R,G,or E; X3 is Y,A,or N; X4 is S,Y,V, or

H; X5 is A,P,F,Y, or H; X6 is R or S; X7 is E or G; X8 is Y or L; X9 is R,Y or L; X10 is Y or G; X11 is D or Y; X12 is R,Y, or F; X13 is M,S,F,D or Y; X14 is A,G,S,or T; X15 is L, M,or F; and X16 is Y,I or V; and wherein the first and/or the third (target) binding domain bind to MSLN and comprise a VL region comprising SEQ ID NO: 1166 as CDR-L 1 wherein X1
 5 is A or S; X2 is G or S; X3 is E or Q; X4 is G,S or K; X5 is I,L,V or F; X6 is R,G or S; X7 is D,S,N or T; X8 is A,S,K or T; X9 is Y or W; X10 is V or L; and X11 is Y or A; SEQ ID NO 1167 as CDR-L2 wherein X1 is A,G or Q; X2 is A or S; X3 is S or T; X4 is G,S,K,I or T; X5 is R or L; X6 is A,P or Q; and X7 is S or T; and SEQ ID NO 1168 as CDR-L 3 wherein X1 is A or Q; X2 is Y,S,A,or T; X3 is G,E,Y,H or Q; X4 is A or S; X5
 10 is S,T or F; X6 is -,P or T; X7 is R,A,L or F; and X8 is V or T.

[46] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or the third (target) binding domain bind to CDH3 and comprise a VH region of SEQ ID NO: 1157 wherein (the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence
 15 table) X1 is Q or E; X2 is V,L; X3 is Q,E ;X4 is A or G; X5 is G or E; X6 is V or L; X7 is K or V X8 isK or Q, X9 is A or G, X10 is V or L, X11 is K or R, X12 is V or L, X13 is A or K, X14 is Y or F, X15 is T or S, X16 is T or S, X17 is S or N, X18 is Y or S, X19 is P or W, X20 is I or M, X21 is Y,N or H, X22 is T or A, X23 is Q or K, X24 is V or M, X25 is S or G, X26 is K,V,N or R, X27 is A,D,R,Y,S,W or H, X28 is Y,S,P,Gr or T, X29 is
 20 S,K,or G, X30 is A,V,D,K, or ,T, X31 is A,-,D,K,S,G, or H, X32 is Y,G, or E, X33 is K,I, or N, X34 is A,S, or N, X35 is S,Q, or G, X36 is S or K, X37 is F or V, X38 is Q or K, X39 is F or V, X40 is I or M, X41 is T or S, X42 is V,I or R, X43 is T,K or N, X44 is T,A,S or K, X45 is S or N, X46 is A,V or L, X47 is L or M, X48 is Q or E, X49 is L or M, X50 is S or N, X51 is S or R, X52 is T or R, X53 is A or
 25 S, X54 is G,D,or E, X55 is T or S, X56 is T,K,or R, X57 is S,Q,W,or R, X58 is -,D,or G, X59 is Y,P,or R, X60 is F,S,G,N or T, X61 is Y,A,or H, X62 is A,-,or V, X63 is F or M, X64 is Y or V; X65 is T,L or M ; and a VL region of SEQ ID NO 1161 wherein X1 is D or E; X2 Q or V; X3 is L,M; X4 is A,S or D; X5 is F,S or T; X6 is A,S; X7is A,V; X8 is P,V,L; X9 is D,E; X10 is A,V; X11 is I,L; X12 is
 30 T,S,N; X13is K,R; X14 is A,S; X15 is Q,D,S,G or E; X16 is S,D,N; X17 is V,I or L; X18is -,K,Y,S or H; X19 is S,N; X20 is F,L,M; X21 is A,N,H; X22 is K,Q; X23 is A,P,V; X24 is K,R; X25 is I,V; X26 is Y,G,W,N; X27 is T,A; X28is S,K; X29 is T,N,R; X30 is L,R; X31is E,A,V,H; X32 is S,E; X33 is A,S,V,D; X34 is D,E; X35 is T,K; X36 is S,R; X37 is A,S,P; X38 is F,V; X39 is A,G; X40 is T,V; X41is Q,V;
 35 X42 is Q,N,H; X43 is F,L,Y,W,N,H; X44 is A,D,Y,S,N; X45 is Q,R,S,G,W,M; X46 is F,Y,T; X47 is F,Y,L; X48 is V,L; and X49 is D or E (wherein all aa per position are meant to be in the alternative “or” even if not explicitly stated).

[47] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or the third (target) binding domain bind to MSLN and comprise a VH region of SEQ ID NO: 1165 wherein (the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence table) X1 is E,Q; X2 is V,L,Q, X3 is E,Q; X4 is A,G,P; X5 is E,G; X6 is V,L; X7 is V,K; X8 is K,Q; X9 is G,S; X10 is E,A,G,R; X11 is S,T; X12 is V,L; X13 is R,S,K; X14 is V,L; X15 is S,T; X16 is A,K,T; X17 is A,V; X18 is Y,I,F; X19 is S,T; X20 is S,F; X21 is S,T; X22 is D,G,S; X23 is Y,G,A,F; X24 is I,W,M; X25 is G,S,V,T,H; X26 is I,V; X27 is A,P; X28 is M,K,Q; X29 is G,C; X30 is I,M,V,L; X31 is A,G,S; X32 is A,S,N,W,Y,V; X33 is Y,S,N; X34 is Y,G,P,S; X35 is D,H,S,N; X36 is G,S; X37 is E,G,S; X38 is G,S,N,F,T,Q; X39 is S,K,W,D,I,-,T; X40 is Y,N; X41 is A,N; X42 is A,P,N,E,D,I,Q; X43 is D,A,S,K; X44 is V,L,F; X45 is K,Q; X46 is G,S; X47 is V,F; X48 is I,M; X49 is S,T; X50 is R,V; X51 is N,T; X52 is A,S; X53 is I,K; X54 is S,N; X55 is S,T,Q; X56 is A,L,F; X57 is Y,S,F; X58 is L,M; X59 is E,K,Q; X60 is M,L; X61 is S,N; X62 is R,S; X63 is V,L; X64 is R,T; X65 is A,S; X66 is D,A,E; X67 is R,K; X68 is D,E,V,L; X69 is E,R,G,P; X70 is R,A,N,Y; X71 is G,S,Y,V,H; X72 is A,P,F,D,Y; X73 is R,G; X74 is M,R,S,D; X75 is E,G; X76 is Y,L; X77 is Y,F; X78 is Y,S,F; X79 is A,G,S,T,H; X80 is L,M,F; X81 is Y,I,V; and X82 is L,M,T ; and a VL region of SEQ ID NO 1169 (the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence table) X1 is E,S,D; X2 is Y,I,L; X3 is E,-,V,T; X4 is V,L,M; X5 is P,S; X6 is G,S; X7 is S,T; X8 is V,L; X9 is A,V,L; X10 is P,V; X11 is E,Q,D; X12 is R,T; X13 is A,V; X14 is S,T; X15 is I,L; X16 is S,T; X17 is A,S; X18 is G,S; X19 is E,Q; X20 is G,S,K; X21 is I,V,L,F; X22 is R,G,S; X23 is D,S,-; X24 is A,S,N,K,T; X25 is Y,W,M; X26 is V,L; X27 is Y,A; X28 is K,Q; X29 is A,S,V; X30 is R,V,K; X31 is V,L; X32 is A,G,Q; X33 is A,S; X34 is S,T; X35 is G,S,K,I,T; X36 is R,L; X37 is A,P,Q; X38 is S,T; X39 is I,V; X40 is E,S,D; X41 is G,N; X42 is N,T; X43 is D,T; X44 is A,F; X45 is R,G,S; X46 is L,T; X47 is E,Q; X48 is A,P; X49 is E,M; X50 is E,F;; X51 is D,V,T; X52 is A,Q; X53 is Y,S,A,T; X54 is G,E,Y,H,Q; X55 is A,S; X56 is S,T,F; X57 is P,T; X58 is R,A,L,F; X59 is V,T; X60 is P,C; X61 is V,L; X62 is E,T; X63 is I,V; and X64 is L,K (wherein all aa per position are meant to be in the alternative “or” even if not explicitly stated).

[48] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or the third (target) binding domain comprise a VH region comprising CDR-H 1, CDR-H2 and CDR-H3 selected from SEQ ID NO: 77 to 79, 86 to 88, 95 to 97, 103 to 105, 111 to 113, 119 to 121, 127 to 129, 135 to 137, 143 to 145, 151 to 153, 159 to 161,

168 to 170, 177 to 179, 185 to 187, 194 to 196, 203 to 205, 212 to 214, 221 to 223, 230 to 232, 238 to 240, 334 to 336, 356 to 358, 365 to 367, 376 to 378, 385 to 387, and 194, 432 and 196, or any combination of CDR-H 1, CDR-H2 and CDR-H3 as disclosed together in the sequence table Tab. 6, preferably 86 to 88, 194, to 196 or 1227 to 1229 and 1237 to 1239.

5 [49] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or third (target) binding domain comprise a VL region comprising CDR-L1, CDR-L2 and CDR-L3 selected from SEQ ID NO: 80 to 82, 89 to 91, 98 to 100, 106 to 108, 114 to 116, 122 to 124, 130 to 132, 138 to 140, 146 to 148, 154 to 156, 162 to 164, 171 to 173, 180 to 182, 188 to 190, 197 to 199, 206 to 208, 215 to 217, 224 to 226, 233 to 235, 241 to 243,
10 337 to 339, 359 to 361, 368 to 370, 379 to 381, 388 to 390, or any combination of CDR-H 1, CDR-H2 and CDR-H3 as disclosed together in the sequence table Tab. 6, preferably 89 to 91 and 197 to 199 or 1230 to 1232 and 1240 1242.

[50] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or third (target) binding domain comprise a VH region
15 selected from SEQ ID NO: 83, 92, 101, 109, 117, 125, 133, 141, 149, 157, 165, 174, 183, 191, 200, 209, 218, 227, 236, 244, 340, 362, 371, 382, 391 and 433, preferably 433 and 92 or 1233 + 1235 and 1243 + 1245 (VH and CH1 in Fab) for the first and third binding domain, respectively.

[51] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or third (target) binding domain comprises a VL
20 region selected from SEQ ID NO: 84, 93, 102, 110, 118, 126, 134, 142, 150, 158, 166, 175, 184, 192, 201, 210, 219, 228, 237, 245, 341, 363, 372, 383, 392, preferably 200 and 93 or 1234 + 1236 and 1244 + 1246 (VL and CL in Fab) for the first and third binding domain, respectively.

[52] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, wherein the first and/or third (target) binding domain comprises a VL
25 region of increased stability by a single amino acid exchange (E to I), selected from SEQ ID NO: 85, 94, 193, 202, 211, 220, 229, 364, 384, 393, preferably 94 and 202.

[53] Within said aspect, it is also envisaged in the context of the present invention to provide an antigen-binding molecule, which comprises a combination of amino acid sequences selected from the
30 group consisting of SEQ ID NOs: 1259 and 1251, 1247 and 1248, 1249 and 1250, 1254, 1255 and 1253, 1252, 1257, 1253 and 1256, and 1254, 1258, 1253 and 1256.

[54] In a second aspect, it is further envisaged in the context of the present invention to provide a polynucleotide encoding an antigen-binding molecule of the present invention, preferably selected from SEQ ID NO: 1070 to 1072 and 1074.

[55] In a third aspect, it is also envisaged in the context of the present invention to provide a vector comprising a polynucleotide of the present invention.

[56] In a fourth aspect, it is further envisaged in the context of the present invention to provide a host cell transformed or transfected with the polynucleotide or with the vector of the present invention.

5 [57] In a fifth aspect, it is also envisaged in the context of the present invention to provide a process for the production of an antigen-binding molecule of the present invention, said process comprising culturing a host cell of the present invention under conditions allowing the expression of the antigen-binding molecule and recovering the produced antigen-binding molecule from the culture.

10 [58] In a sixth aspect, it is further envisaged in the context of the present invention to provide a pharmaceutical composition comprising an antigen-binding molecule of the present invention or produced according to the process of the present invention.

[59] Within said aspect, is also envisaged in the context of the present invention that the pharmaceutical composition is stable for at least four weeks at about -20°C.

15 [60] It is further envisaged in the context of the present invention to provide the antigen-binding molecule of the present invention, or produced according to the process of the present invention, for use in the prevention, treatment or amelioration of a disease selected from a proliferative disease, a tumorous disease, cancer or an immunological disorder.

20 [61] Within said aspect, it is also envisaged in the context of the present invention that the disease preferably is acute myeloid leukemia (AML), Non-Hodgkin lymphoma (NHL), Non-small-cell lung carcinoma (NSCLC), pancreatic cancer and Colorectal cancer (CRC)]. In a seventh aspect, it is further envisaged in the context of the present invention to provide a method for the treatment or amelioration of a proliferative disease, the method comprising administering to a subject in need thereof a molecule comprising at least one polypeptide chain, wherein the molecule comprises

25 (i.) a first binding domain which preferably comprises a paratope which specifically binds to a first target cell surface antigen (e.g. TAA1),

(ii.) a second binding domain which preferably comprises a paratope which specifically binds to an extracellular epitope of the human - and preferably the Macaca- CD3ε chain,

(iii.) a third binding domain which preferably comprises a paratope which specifically binds to a second target cell surface antigen (e.g. TAA2), and

30 (iv.) a fourth binding domain which preferably comprises a paratope which specifically binds to an extracellular epitope of the human -and preferably the Macaca- CD3ε chain,

wherein the first binding domain and the second binding domain form a first bispecific entity and the third and the fourth binding domain form a second bispecific entity, and

wherein the molecule comprises a spacer entity having a molecular weight of at least about larger than about 5 kDa and/or having a length of more than 50 amino acids, wherein the spacer entity spaces apart the first and the second bispecific entity by at least about 50 Å (distance between centers of mass of the first and the second bispecific entity), and which spacer entity is positioned between the first and the second bispecific entity.

[62] Within said aspect, it also envisaged in the context of the present invention also provides a method to address a disease-associated target being significantly co-expressed on a pathophysiological and one or more physiological tissues by providing a multichain multitargeting bispecific antigen-binding molecule of the format described herein, wherein the molecule addresses (i.) the target expressed both on the disease-associated and the physiological tissue and (ii.) a further target which is disease associated but not expressed on the physiological tissue under (i.), wherein the method preferably avoids the formation of intra-abdominal adhesions and/or fibrosis where such target is MSLN.

[63] Within said aspect, it is also envisaged in the context of the present invention that the disease preferably is a tumorous disease, cancer, or an immunological disorder, comprising the step of administering to a subject in need thereof the antigen-binding molecule of the present invention, or produced according to the process of the present invention, wherein the disease preferably is acute myeloid leukemia, Non-Hodgkin lymphoma, Non-small-cell lung carcinoma, pancreatic cancer and/or Colorectal cancer.

[64] Within said aspect, it is also envisaged in the context of the present invention that TAA1 and TAA2 are preferably selected from EpCAM and MSLN, MSLN and EpCAM, MSLN and CDH3, CDH3 and MSLN, FLT3 and CLL1, and CLL1 and FLT3.

[65] In an eighth aspect, it is also envisaged in the context of the present invention to provide a kit comprising an antigen-binding molecule of the present invention, or produced according to the process of the present invention, a polynucleotide of the present invention, a vector of the present invention, and/or a host cell of the present invention.

DESCRIPTION OF THE FIGURES

[66] **Figure 1:** Overview of multichain multitargeting bispecific antigen-binding molecules disclosed in the invention comprising a dimerizing domain as spacer. Black domains are Anti-CD3 domains, and striped and dotted domains are target binding domains (see in-figure legend of Figure 2). Domain arrangement in each molecule as follows: A: target binding domains and CD3 binding domains in the format of scFv, both N- and C-terminally of dimerizing spacer domain, respectively (“cis” orientation, i.e. the at least two target binding domains and at least two CD3 binding domains

are connected to the same monomer of the dimerized spacer, respectively, or in other words, are on the same side of a vertical plane defined by the dimerizing spacer cutting the dimerizing spacer); B: first target binding domain is a Fab, CD3 binding domains and other target binding domain are in the form of scFv, both N- and C-terminally of spacer dimerizing domain, respectively (trans orientation, i.e. the at least two target binding domains and at least two CD3 binding domains are connected to the opposite monomer of the dimerized spacer, respectively, or in other words, are on the opposite side of a vertical plane defined by the dimerizing spacer cutting the dimerizing spacer); C: both at least two target binding domains are Fabs, and CD3 binding domains are in the form of scFvs, both N- and C-terminally of spacer dimerizing domain, respectively (trans); D: both at least two target binding domains are Fabs N-terminally of dimerizing spacer domain, both and CD3 binding domains are in the form of scFv ; E: target binding domains are Fabs N-terminally of spacer dimerizing domain and CD3 binding domains are Fabs C-terminally of spacer dimerizing domain

[67] **Figure 2:** FIG. 2 A to M shows examples of multichain multitargeting bispecific antigen-binding molecules of the invention, wherein the spacer is a dimerizing heteroFc, and wherein at least three of the four N- and C-termini, respectively, are linked to a target and/or CD3 binding domain.

[68] **Figure 3:** FIG.3 A to H shows examples of multichain multitargeting bispecific antigen-binding molecules of the invention, wherein the spacer is a dimerizing heteroFc, and wherein one N- and one C-terminus out of the four N- and C-termini of the heteroFc, respectively, are linked to a target and/or CD3 binding domain (A, C, E to G), or wherein the spacer is a scFv (B, D and H)..

[69] **Figure 4:** FIG. 4 A to I shows cytotoxicity curves of CDH3 T-cell engager molecule 1, MSLN T-cell engager molecule 1 and MSLN-CDH3 T-cell engager molecules 1-7 on parental double positive HCT116 WT cells versus target-knockout HCT116 cells. Effector cells were unstimulated Pan T-cells..

[70] **Figure 5:** FIG. 5 A to I shows cytotoxicity curves of CDH3 T-cell engager molecule 1, MSLN T-cell engager molecule 1 and MSLN-CDH3 T-cell engager molecules 1-7 on parental double positive GSU WT cells versus target-knockout GSU cells. Effector cells were unstimulated Pan T-cells.

[71] **Figure 6:** FIG 6. A to C shows cytotoxicity curves of mono CDH3 T-cell engager molecule 1 (A), MSLN T-cell engager molecule 1 (B) and MSLN-CDH3 T-cell engager molecule 24 comprising both target binder in Fab format on parental double positive GSU WT cells versus target-knockout GSU cells. Effector cells were unstimulated Pan T-cells.

Detailed Description

[72] In the context of the present invention, a multichain multitargeting bispecific molecule is provided comprising at least five distinctive structural entities, i.e. (i.) a first domain binding to a target cell surface antigen (e.g. a first tumor associated antigen, TAA), (ii.) a second domain binding to an extracellular epitope of the human - and preferably non-human, e.g. Macaca- CD3ε chain,

wherein the first binding domain and the second binding domain together form a first bispecific entity, (iii.) a spacer which connects but spaces apart the first bispecific entity from a second bispecific entity comprising (iv.) a third domain binding to the same or preferably a different target cell surface antigen (e.g. a second TAA), and (v.) a fourth domain binding to an extracellular epitope of the human -and
5 preferably non-human, e.g. Macaca- CD3 ϵ chain. Molecules of the format of the present invention typically exhibit the advantage to be characterized by avidity-driven potency and specificity from two targets being co-expressed on the target cell, which typically leads to a reduction of undesired cytokine release (and associated clinically relevant side effects such as CRS) while at the same time ensuring effective antitumor activity, preferably also in solid tumors such as colorectal cancer, non-small-cell
10 lung carcinoma and pancreatic cancer.

[73] It is a surprising finding in the context of the present invention that bispecific (T-cell engaging) multichain multitargeting (antigen-binding) molecules according to the present invention provides a double avidity effect, both on the target cell binder and the effector cell binder side due to their specific format which leads to an efficient each other complementing target cell kill. This effect
15 is facilitated by the molecule format specifically targeting two (different) antigens on one target cell, such as a cancer cell, and in contrast, by significantly less targeting non-target cells while mediating a potent T-cell response against said target cell at the same time. By being capable to address two target antigens at the same time, the likeliness of targeting a target cell associated with a disease instead of a physiologic cell is greatly increased when two TAAs are chosen which are typically associated with a
20 target cell associated with a disease. Hence, a T-cell engaging multichain multitargeting molecule according to the present invention, both provides improved efficacy and safety with regard to existing bispecific antibodies or antibody-derived constructs which are T-cell engaging. Said advantageous properties are preferably achieved by the fact that the multichain multitargeting bispecific molecules of the present invention comprise two bispecific entities comprising each a target binding domain and
25 an effector (CD3) binding domains which can act in a pathophysiologic environment without (e.g. sterically) hindering each other while complementing each other at the same time. Said action of the two bispecific entities within the one multichain multitargeting bispecific molecule of the present invention from each other means that the target binding domain (e.g. the first domain) and the effector CD3 binding domain (e.g. the second domain) of the first bispecific entity can interact with their
30 respective binding partners to form a cytolytic synapse between target cell and T-cell, without disturbing interaction with the target binding domain (e.g. the third domain) and the effector domain (e.g. the forth domain) of the second bispecific entity. However, in order to provide the desired action and, in consequence, therapeutic function, preferably both target binding domains of both the first and the second bispecific entity must engage their respective target in order to involve the effector CD3
35 binding domains of the first and second bispecific entity completely. Further, it was a surprising finding that the two respective bispecific entities must be functionally preserved by structural separation in the molecule format in a specific manner in order to benefit from the double avidity

effect required to achieve the extraordinary efficacy described and safety implied herein. It was especially surprising that the two bispecific entities comprising a target binding and a CD3 binding domain, respectively, do not need to be on one chain N- and C-terminally of the (central) spacer in order to be structurally positioned to act as described herein. Both the target and/or CD3 binder of one or both bispecific entities can be a Fab, i.e. comprise two chains respectively. What is more surprising is the fact that also the spacer may be two-chained, preferably in the form of a heteroFc. In such a case, it is surprising that also bispecific entities when two domains are not on the same chain but kept close together by means of the four-moiety spacer (e.g. hetero Fc) which at the same time keeps the two domains of each bispecific entity in place to act together and separates the two bispecific entities from each other to act without interfering with each other. This arrangement of spacer and domains and the employment of two different TAA binding domains are two CD3 binding domains of preferably low affinity -preferably binding to CD3 ϵ - leads to the surprising technical effect of increased target cell selectivity and the reduced risk of the key side effect, i.e. undesired cytokine release. At the same time, multichain molecules of the invention can be produced well both regarding yield and purity.

[74] As a secondary effect in addition or alternatively to the herein described increased specificity, and therefore safety, the likeliness of targeting a target cell such as a cancer cell by a multichain multitargeting antigen-binding molecule versus a monotargeting molecule is greatly increased once such target cell has undergone antigen loss and, thus, is prone to tumor escape from effective anti-tumor therapy because one valid antigen to target remains on the cell which has undergone antigen escape. Said effect in terms of increased activity compared to molecules comprising only one CD3 binder and/or target binder and do not comprise the two linked but spaced apart bispecific entities is preferably achieved when both CD3 binders are of low affinity, such as a CD3 binding domain comprising a VH and VL of, for example, SEQ ID NOs 67 and 68, respectively, linked by a linker of SEQ ID NO 1 or 3.

[75] The above-specified finding underlying the present invention is surprising in view of the teaching of the prior art. For example, antigen-binding formats comprising more than one target binding domain and effector binding domain, respectively, are known in the art, e.g. the Adaptir™ format. However, such formats do not provide two bispecific entities which can individually interact with their respective target and effector and work together at the same time and, consequently, cannot achieve the effect of double avidity on both the target binder and the effector binder side to the extent of effectively providing a large selectivity gap to the advantage of the multichain multitargeting molecule. According to the present invention, the two bispecific entities must be spaced apart from each other by a certain distance, preferably of at least 50 Å, more preferably at least 60, 70, 80, 90 or at least 100 Å. The indicated distance [Å] between the two bispecific entities is typically understood in the context of the present invention as the distance between the centers of mass of the two bispecific entities, respectively. In general, the center of mass (COM) of a distribution of mass (here, a bispecific

entity comprising a binding domain which binds to a target cell surface antigen and a binding domain which binds to an extracellular epitope of the human -and preferably the Macaca- CD3 ϵ chain, both binding domains preferably in a Fab or a single domain format, preferably selected from scFv and scFab format and linked by a peptide linker) in space is understood as the unique point where the weighted relative position of the distributed mass sums to zero. The distance is typically determined by molecular modeling making use of generally accepted modeling programs (MD/visualization software) which can identify COMs given input structures and such as PyMOL (The PyMOL Molecular Graphics System, Version 2.3.3, Schrödinger, LLC.) which is typically based on ensembles of snapshot structures from MD simulations. The mass of each atom is typically part of an underlying “force field” as generally known in the art. Alternatively and/or additionally, distances can be determined by crystallography, cryo electron microscopy, or nuclear magnetic resonance analytic technology.

[76] A typical approach of obtaining distances through molecular modeling as given in the present invention is as follows:

- 1) Obtaining an atomistic structure of the complete bispecific antigen-binding molecule. Structure sources may be selected from the group consisting of:
 - a. Protein X-ray crystallography with resolution preferably below 5 Å enabling visibility of amino acid backbones and side-chains;
 - b. Cryogenic electron microscopy (cryo-EM) with resolution preferably below 5 Å enabling visibility of amino acid backbones and side-chains;
 - c. In silico homology modeling of the entire molecule based on a single, highly-homologous crystal and/or cryo-EM structure (preferably above 60% sequence identity);
 - d. In silico homology modeling involving linking 2 or more experimental structures. The structures are preferably identical or highly homologous (preferably above 60% sequence identity) to domains found in the complete bispecific antigen-binding molecule. In case of lack of experimental linker conformations, the model is preferably refined in an explicit-solvent Molecular Dynamics (MD) simulation (simulation length of preferably at least 100 ns unless energy convergence is obtained faster). The simulation is carried out with a state-of-the-art software (e.g. Schrodinger, Amber, Gromacs, NAMD or equivalent) with parameters corresponding to room temperature and pressure. No artificial forces are applied during the simulation (i.e. preferably excludes methods such as metadynamics or steered molecular dynamics). Similarly, preferably no artificial geometrical restraints are imposed on the molecule.

2) Identifying centers of mass (COM) of the relevant molecule domains. This is typically performed with the used MD software or with visualization tools such as PyMOL or VMD. The centers of mass can be defined as a pseudo-atoms or non-hydrogen atoms closest to the true COM. Inter-domain linkers are typically not considered as part of the domain.

5 3) Using the same software, report the distance (in Angstrom, Å) between the two COMs. If an MD simulation was used to refine a homology model (as described in 1d), the median distance over multiple simulation snapshots is reported. To further diminish potential inaccuracy of the initial model, at least the first 10% of the simulation, preferably up to 50% if the signal significantly changes, are omitted when calculating the median distance between COMs and when extracting the snapshots
10 for visualizing the MD simulation.

[77] If not indicated otherwise, distances [Å] in the context of the present invention are median distances as determined by MD simulations.

[78] The preferred distance between the first and the second bispecific entity as disclosed herein is facilitated by a spacer entity (in short spacer) between the two bispecific entities which spaces the two
15 bispecific entities apart and keeps them in a separated position. The spacer is of a certain size, preferably at least more than 5 kDa, more preferably at least about 10, 15, 20, 25, 30, 35, 40, 45 or even at least 50 kDa and hereby prevents an undesired interaction of the two separated bispecific entities. The preferred range in molecular size of the spacer is about 15 to 200 kDa, preferably about 15 to 150 kDa, in order to facilitate the separation of the two bispecific entities according to the
20 present invention and to maintain a high overall activity of the molecule. Typically, too large spacers, e.g. larger than about 200 kDa, may impact the ability of the two bispecific entities to bind to two target surface structures on the same target cell which in turn may reduce the overall activity of the molecule against the target cell. Hence, the typical maximum preferred size in terms of molecular weight of the spacer is about 200 kDa, preferably about 150 or 120 kDa and even more preferably
25 about 100 kDa. A typical spacer of maximum preferred size is a double scFc domain as disclosed herein (two scFc linked to each other forming one larger single chain spacer) of about 105.7 kDa. Example sizes of spacers which typically sufficiently separate the two bispecific entities are PSI domain of Met-receptor of about 5.3 kDa, ubiquitin of about 8.6 kDa, fibronectin type III domain from tenascin of about 10.1 kDa, SAND domain of about 11 kDa, neta-2-microglobulin of about 11.9 kDa,
30 Tim-3 (aa 24-130) of about 12.2 kDa, MiniSOG of about 13.3 kDa, SpyCatcher of about 12.1 kDa associated with SpyTag of about 1.7 kDa linked together preferably via isopeptide bond formation to form a two-chain-spacer of about 13.8 kDa, VHH antibody lama domain of about 14 kDa, PD-1 binding domain from human programmed cell death 1 ligand (PDL1) of about 14.4 kDa, granulocyte-macrophage colony stimulating factor (GM-CSF) of about 14.5 kDa, intrleukin-4 of about 15 kDa,
35 interleukin-2 of about 15.4 kDa, CD137L (4-1BBL; TNFSF9) ectodomain of about 17.7 kDa, programmed cell death protein 1 (PD-1) of about 16.6 kDa, green fluorescent protein (GFP) of about

26.3 kDa, single chain Fc region (scFc) as described herein of about 52.8 kDa (about 54.6 kDa with N- and C-terminal linkers (G₄S)₃, respectively), human serum albumin (HSA) of about 66.5 kDa (about 68.3 kDa with N- and C-terminal linkers (G₄S)₃, respectively) and double scFc (two scFc linked to each other forming one larger single chain spacer) of about 105.7 kDa (about 107.5 kDa with N- and C-terminal linkers (G₄S)₃, respectively). In general, the more rigid the spacer is, the less is the median distance required which otherwise has to include a safety margin for flexible spacers.

[79] Also, a preferred spacer in the context of the present invention, such as a globular domain, typically has a N- and a C-terminus which are spatially not too close to each other in order to efficiently space apart the two bispecific entities according to the invention. In this regard, spacers typically show a distance between the N- and the C-terminus which is significantly larger than 10 Å. A distance between N- and C-terminus of the spacer which is lower or about 10 Å is considered “close”. Hence, a spacer in the context of the present invention preferably has a distance between the alpha-carbon atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus of at least 20 Å, more preferably at least 30 Å, even more preferably at least 50 Å, which distance typically ensures to space the first and the second bispecific entity apart by at least 50 Å as described herein. Alpha-carbon (α -carbon) is understood herein as a term that applies to proteins and amino acids. It is the backbone carbon before the carbonyl carbon atom in the molecule. Therefore, reading along the backbone of a typical protein would give a sequence of $-\text{[N—C}\alpha\text{—carbonyl C]}_n\text{—}$ etc. (when reading in the N to C direction). The α -carbon is where the different substituents attach to each different amino acid. That is, the groups hanging off the chain at the α -carbon are what give amino acids their diversity. Hence, in the context of the present invention, a spacer is less preferred, even if it has a size of at least 5 kDa and a length of more than 50 aa if the distance between the alpha-carbon atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus is too close, i.e. if it is only, e.g., about 10 Å. For example, preferred spacers show typical distances between the alpha-carbon atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus as follows: scFc (based on 5G4S crystal structure) 89 Å, HSA (based on 5VNW crystal structure): 77 Å, ubiquitin (based on 1UBQ crystal structure): 37 Å and SAND (based on 1OQJ crystal structure): 32 Å. In contrast, HSP70-1 (based on 3JXU crystal structure) shows only a distance of 9 Å between the alpha-carbon atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus. At the same time, HSP70-1 provides only a median distance between the COMs of first and the second bispecific entity in the context of the present invention of about 48 Å which is below the threshold of 50 Å median distance, and significantly below the typically about 60 – 100 Å median distance between the COMs of the two bispecific entities as facilitates by preferred spacers such as scFc, HSA, ubiquitin and SAND. Thereof, scFc (SEQ IN NO: 25) is preferred.

[80] Alternatively, a non-globular but rigid linker may serve as a spacer in the context of the present invention which spaces apart the two bispecific entities. Such linkers comprise (PA)₂₅P (SEQ ID NO: 1097) and A(EAAAK)₄LEA(EAAAK)₄A (SEQ ID NO: 1096), even if the Mw is below 5 kDa (here 4.3 kDa) and the amino acid length is only about or below 50 (51 and 46 aa, respectively).

5 However, such spacers are typically less preferred than globular domains which preferably additionally increase half-life.

[81] As it is also contemplated within the context of the present invention, the spacer between the two bispecific entities is a polypeptide which typically comprises more than 50 amino acids, preferably at least about 75, 100, 150, 200, 250, 300, 350, 400, 450 or at least 500 amino acids. The preferred range in amino acid length of the spacer is about 100 to 1500 amino acids, preferably about 100 to 1000 amino acids, more preferably about 250 to 650 amino acids in order to facilitate the separation of the two bispecific entities according to the present invention. This is to preferably maintain a high overall activity of the entire molecule according to the present invention (not necessarily of the individual and spaced-apart bispecific entities, which may have low affinities (and low activities) individually in order to increase specificity for double positive target cells) which is typically below 20 pM, preferably below 5 pM, more preferably below 1 pM. Typically, too large spacers, e.g. longer than about 1500 amino acids, may impact the ability of the two bispecific entities to bind to two target surface structures on the same target cell which in turn may reduce the overall activity of the molecule against the target cell. Hence, the typical maximum preferred length of the spacer is about 1500 amino acids, more preferably about 1000 amino acids. Example amino acid lengths of spacers which sufficiently separate the two bispecific entities are PD-1 of about (ECD 25-167) 143 aa, scFc as described herein of about 484 aa (about 514 aa with N- and C-terminal linkers (G₄S)₃, respectively), HSA of about 585 aa (about 615 aa with N- and C-terminal linkers (G₄S)₃, respectively), and double scFc of about 968 aa (about 998 aa with N- and C-terminal linkers (G₄S)₃, respectively). Further spacers include, ubiquitin of about 76 aa, fibronectin type III domain from tenascin of about 90 aa, SAND domain of about 90 or 97 aa, beta-2-microglobulin of about 100 aa, Tim-3 (aa 24-130) of about 108 aa, MiniSOG of about 115 aa, SpyCatcher of about 113 aa associated with SpyTag of about 14 aa linked together preferably via isopeptide bond formation to form a two-chain-spacer of about 127, VHH antibody lama domain of about 129 aa, PD-1 binding domain from human programmed cell death 1 ligand (PDL1) of about 126 aa, granulocyte-macrophage colony stimulating factor (GM-CSF) of about 127 aa, interleukin-4 of about 129 aa, interleukin-2 of about 133 aa, CD137L (4-1BBL; TNFSF9) ectodomain of about 167 aa, and green fluorescent protein (GFP) of about 238 aa.

[82] The composition and arrangement of the preferred spacer amino acid sequences preferably confer a certain rigidity and are not characterized by high flexibility. Rigidity in the context of the present invention is typically present when a spacer of more than 50 aa and/or a molecular weight over

5 kDa facilitates a maximum distance between the centers of mass of the two bispecific entities in a molecule according to the present invention which is smaller than 200% (or 2-fold) the median distance. Accordingly, a preferred rigid spacer in the context of the present invention does not extend further than about 100% of its median length, more preferably not more than about 80% (each calculated as distance between centers of mass of the two bispecific entities). Hence, a preferred rigid spacer in the context of the present invention which spaces apart the two bispecific entities by about 100 Å (median distance) does not extend further than to 200 Å (maximum distance). For example, a typical median distance between centers of mass of the bispecific entities of a molecule having the format of the present invention comprising a scFc (such as SEQ ID NO: 25) as spacer is about 101 Å. However, a maximum distance in such a case is typically about 182 Å, i.e. not more than about 100% or even only about 80% with respect to the median distance. Such a spacer is considered rigid in the context of the present invention. In contrast, a molecule comprising a (G₄S)₁₀ (SEQ ID NO: 8) as spacer, which is a linear polypeptide without a e.g. globular structure, shows a typical a median distance of about 48 Å and a maximum distance of about 179 Å. Hence, such a spacer as (G₄S)₁₀ shows a high flexibility and not the rigidity of a preferred spacer as advantageous feature according to the present invention. In this regard, spacer amino acid sequences may typically be rich in proline and less rich in serine and glycine. Especially envisaged are spacers which are folded polypeptides e.g. of secondary order (e.g. helical structures) or of ternary order forming e.g. three dimensional protein domains structures which in turn ensure a certain rigidity by their constitution and preferably confer further advantageous effects such as in vivo half-life extension of the multichain multitargeting bispecific molecule as a therapeutic agent. Typical domain structures comprise hydrophobic cores with hydrophilic surfaces. In the context of the present invention, proteins having a structure of a globular protein are preferred as spacers. Globular proteins are understood in the context of the present invention to be spherical ("globe-like") proteins and are one of the common protein types. Globular proteins in the context of the present invention may be characterized by a globin fold. Spacers comprising an Fc domain or parts or a multiple thereof, a PD-1 or an HSA domain are in particular envisaged. Also envisaged are spacers which comprise combinations of different globular proteins or parts thereof, which even more preferably comprise a Fc receptor binding function in order to increase the half-life of the molecule according to the present invention. The format described herein with the separation of the two bispecific entities has distinctive advantages. If only one target is present which is addressed by the first binding domain, then the first domain "uses" only the second domain to engage a T cell but not the fourth domain, or alternatively, the third domain uses the fourth but not the second (or to a much lesser extend due to the spacer). If only one target is present, the K_d of preferably low affinity CD3 binder as disclosed herein prevents efficient T-cell engagement. Thus, selectivity is increased with respect to other (dual) targeting molecules.

[83] If both targets are present, the multichain multitargeting bispecific T cell engager of the invention binds more firmly to the target cell (by avidity gain) and both low affinity CD3 binding

domains of the invention such as I2L can be used to engage T cells (also by avidity gain), for example the second domain binding to a CD3 domain on an effector T cell and the third domain binding to a target antigen are less likely to form a cytolytic synapse and therefore do not act together as a bispecific entity which would otherwise lead to less beneficial cytotoxic activity profile. This has the advantage that the first and the fourth domain are not left “useless” which would mean that the full effect of the double avidity by double binding of a target and an effector binding domain, respectively, could not be made use of. Likewise, the first domain binding to a target antigen and the fourth domain binding to a CD3 domain on an effector T cell are prevented from theoretical interaction which would eventually render the second and the third domain useless for forming a cytolytic synapse with their intended “partner domains” in their respective bispecific entities.

[84] Typically, the advantageous avidity effect conferred by a multichain multitargeting bispecific molecule according to the present invention is indicated by a differential activity factor or “selectivity gap” between the activity of the molecule on double positive cells, i.e. a target cell which carries (i.) two different targets which combination is overexpressed on the cell type to be targeted and being associated with a particular disease and/or (ii.) one target at overexpressed levels. In either case, a molecule according to the present invention targeting two (preferably different) targets at the same time, will preferably bind to such a target cell in comparison to a non-target cell expressing either only one of two targets or the one target at lower expression levels and, in consequence, will induce a more pronounced T cell response. As it preferred for a multichain multitargeting bispecific molecule of the present invention, the activity in terms of increased cytotoxicity as determined, for example, by lower EC_{50} values, is at least 5 times, preferably 10, more preferably 30, 50, 80 or even 100 times larger on target cells (e.g. characterized by expressing both different targets or the one target at high levels) than on non-target cells (e.g. characterized by expressing only one of two targets or the one target only at low levels). Said selectivity gap in the context of the present invention is preferably larger than 100 times. It is envisaged in the context of the present invention that the selectivity gap (which can also be defined as activity gap) is at least 250, 500, 750 or even 1000 times which greatly improves efficacy and safety of the present multichain multitargeting bispecific molecule in comparison to monotargeting bispecific molecules of various formats.

[85] A further aspect envisaged in the context of the present invention is the further support of the double avidity effect conferred by the format of the multichain multitargeting antigen-binding molecule by means of a low affinity, preferably both of the target antigen binders and the CD3 effector binders. In the context of the present invention, a CD3 binder with an affinity below $KD 1.2 \times 10^{-8} M$ is preferred. Especially preferred are CD3 binders which have an activity which is 10 times lower, more preferably 50 times lower or even more preferred 100 times lower than that of a CD3 binder having a $KD 1.2 \times 10^{-8}$. Without wanting to be bound by theory, the avidity effect is contemplated to be more pronounced when two binders with relatively balanced, i.e. typically two low to medium

high, preferably low affinity binders bind to two targets on the same target cell compared to binders with mixed or, typically, higher affinity which would trigger cytolytic activity also if only one target on a cell was bound which could, for example, be a physiologic non-target cell which should not be targeted in order to avoid off-target toxicity and related side effects.

5 [86] Accordingly, the multichain multitargeting bispecific antigen-binding molecules according to the present invention which bind to two (preferably different) targets on a target cell in order to show significant cytotoxic activity preferably do show less side effects than monotargeting bispecific antigen-binding molecules which bring together effector T cell and target cell. This is demonstrated, for example, by a significant reduction in release of key cytokines IL-2, IL-6, IL-10, TNF α and IFN γ
10 which are an indicator for side effects on a clinical stage. For example, release of IL-6 is typically reduced upon use of a multichain multitargeting bispecific antigen-binding molecule according to the present invention with respect to a corresponding monotargeting bispecific molecule. As it is known in the art, interleukin 6 (IL-6) seems to hold a key role in CRS pathophysiology since highly elevated IL-6 levels are seen in patients with CRS (Shimabukuro-Vornhagen et al. Journal for ImmunoTherapy of
15 Cancer (2018) 6:56). As CRS is a serious side effect in immunotherapies, such reduction is an indication for less CRS in the clinical stage.

[87] Further, the multichain multitargeting bispecific antigen-binding molecules according to the present invention which bind to two (preferably different) targets on a target cell in order to show significant cytotoxic activity preferably do show less side effects than monotargeting bispecific
20 antigen-binding molecules in terms of toxicity tissue damage. It has been a surprising finding that a multispecific molecule of the format as described herein shows higher tolerability, i.e. higher doses can be administered than corresponding monotargeting bispecific molecules without clinical findings such as tissue damage examined by histopathological examination. For example, a dose of 1.5 $\mu\text{g}/\text{kg}$ of a MSLN monotargeting bispecific antigen-binding molecule (SEQ ID NO: 1183) was not tolerated and resulted in mortality whereas a dose of 0.1 $\mu\text{g}/\text{kg}$ was tolerated. Conversely, a multichain
25 multitargeting CDH3-MSLN bispecific molecule (SEQ ID NO: 251) according to the present invention was tolerated at doses of up to 1000 $\mu\text{g}/\text{kg}$. Histopathological changes seen with the monotargeting molecule were generally more severe at doses of 1.5 $\mu\text{g}/\text{kg}$ than those with the multichain multitargeting molecule at 1000 $\mu\text{g}/\text{kg}$, respectively. Adhesions or irreversible fibrotic changes as induced by the monotargeting molecule were absent after treatment with the multichain
30 multitargeting molecule. Therefore, the tolerability of a multichain multitargeting molecule according to the present invention is, e.g., 600 (histopathology) to, e.g., 10.000 (tolerated dose) times higher than for a corresponding monotargeting molecule despite equivalent in vitro potency against tumor cells. Hence, the multichain multitargeting molecules of the present invention are particularly suitable in
35 therapeutic settings, where targets are addressed which are significantly present not only on disease-associated (pathophysiological) but also or even predominately on physiological tissues which should, however, not be targeted by a cytotoxic immunotherapy. This is the case, e.g., for MSLN which is

typically expressed in mesothelial cells which form the lining of several body cavities: the pleura (pleural cavity around the lungs), peritoneum (abdominopelvic cavity including the mesentery, omenta, falciform ligament and the perimetrium) and pericardium (around the heart). Addressing targets like MSLN by cytotoxic immunotherapy bears the risk of severe side effects such as intra-abdominal adhesions and/or fibrosis. Intra-abdominal adhesions are understood herein as pathologic scars formed between intra-abdominal organs. Adhesions can occur in the presence of intraperitoneal inflammation and cause peritoneal surfaces to adhere to each other. Adhesions can cause problems if the scarring limits the free movement of organs (Mutsaers S.E., Prele C.M., Pengelly, S., Herrick, S.E. Mesothelial cells and peritoneal homeostasis. *Fertil Steril* 2016, 106(5) 1018-1024). Fibrosis is understood herein as a common pathological outcome of several etiological conditions resulting in chronic tissue injury and is usually defined as an excessive deposition of extracellular matrix (ECM) components, leading with time to scar tissue formation and eventually organ dysfunction and failure (Maurizio Parola, Massimo Pinzani, *Pathophysiology of Organ and Tissue Fibrosis, Molecular Aspects of Medicine* 2019, (65) 1). Hence, the present invention also provides a method to address a disease-associated target being significantly co-expressed on a pathophysiological and one or more physiological tissues by providing a multichain multitargeting bispecific antigen-binding molecule of the format described herein, wherein the molecule addresses (i.) the target expressed both on the disease-associated and the physiological tissue and (ii.) a further target which is disease associated but not expressed on the physiological tissue under (i.), wherein the method preferably avoids the formation of intra-abdominal adhesions and/or fibrosis where such target is MSLN.

[88] It is envisaged that the bispecific antigen-binding molecules according to the present invention have cross-reactivity to, for example, cynomolgus monkey tumor-associated antigens such as CDH3, MSLN, CD20, CD22, FLT3, CLL1, and EpCAM. It is in particular envisaged in the context of the present invention that two targets can be addressed by one multichain multitargeting bispecific antigen-binding molecule simultaneously.

[89] Alternatively and besides the major advantage of increasing selectivity as described herein, dual targeting can mitigate lack of accessibility of one target when targeting the remaining target can trigger a sufficient residual effect. Examples are (i) the presence of soluble target which would “mask” the target on the target cell by binding the antigen-binding molecule without allowing the remaining molecule any therapeutic effect and (ii) antigen loss (lowering target expression on target cell) as the driving factor for tumor escape..

[90] For example, a multichain multitargeting antigen-binding molecule according to the present invention such as a construct directed against MSLN as TAA1 and CDH3 as TAA2 is suitable for use in the treatment, amelioration or prevention of cancer, in particular cancer selected from the group consisting of, lung carcinoma, head and neck carcinoma, a primary or secondary CNS tumor, a

primary or secondary brain tumor, primary CNS lymphoma, spinal axis tumors, brain stem glioma, pituitary adenoma, adrenocortical cancer, esophagus carcinoma, colon cancer, breast cancer, ovarian cancer, NSCLC (non-small cell lung cancer), SCLC (small cell lung cancer), endometrial cancer, cervical cancer, uterine cancer, transitional cell carcinoma, bone cancer, pancreatic cancer, skin cancer, cutaneous or intraocular melanoma, hepatic cancer, biliary duct cancer, gall bladder cancer, kidney cancer, rectal cancer, cancer of the anal region, stomach cancer, gastrointestinal (gastric, colorectal, and duodenal) cancer, cancer of the small intestine, biliary tract cancer, cancer of the urethra, renal cell carcinoma, carcinoma of the endometrium, thyroid cancer, testicular cancer, cutaneous squamous cell cancer, melanoma, stomach cancer, prostate cancer, bladder cancer, osteosarcoma, mesothelioma, Hodgkin's Disease, non Hodgkins's lymphoma, chronic or acute leukemia, chronic myeloid leukemia, lymphocytic lymphomas, multiple myeloma, fibrosarcoma, neuroblastoma, retinoblastoma, and soft tissue sarcoma..

[91] It is especially envisaged in the context of the present invention that a multichain multitargeting antigen-binding molecule which preferably addresses two different target cell surface antigens thereby is very specific for its target cell and, therefore, preferably safe in its therapeutic use. Efficacy in terms of tumor growth inhibition has been demonstrated in vivo in a mouse model.

[92] Preferred target cell surface antigens in the context of the present invention are, MSLN, CDH3, FLT3, CLL1, EpCAM, CD20, and CD22. Typically, target cell surface antigens in the context of the present invention are tumor associated antigens (TAA). B-lymphocyte antigen CD20 or CD20 is expressed on the surface of all B-cells beginning at the pro-B phase (CD45R+, CD117+) and progressively increasing in concentration until maturity. CD22, or cluster of differentiation-22, is a molecule belonging to the SIGLEC family of lectins. It is found on the surface of mature B cells and to a lesser extent on some immature B cells. Fms like tyrosine kinase 3 (FLT3) is also known as Cluster of differentiation antigen 135 (CD135), receptor-type tyrosine-protein kinase FLT3, or fetal liver kinase-2 (Flk2). FLT3 is a cytokine receptor which belongs to the receptor tyrosine kinase class III. CD135 is the receptor for the cytokine Flt3 ligand (FLT3L). The FLT3 gene is frequently mutated in acute myeloid leukemia (AML). C-type lectin-like receptor (CLL1), also known as CLEC12A, or as MICL. It contains an ITIM motif in cytoplasmic tail that can associate with signaling phosphatases SHP-1 and SHP-2. Human MICL is expressed as a monomer primarily on myeloid cells, including granulocytes, monocytes, macrophages and dendritic cells and is associated with AML. Mesothelin (MSLN) is a 40 kDa protein that is expressed in mesothelial cells and overexpressed in several human tumors. Cadherin-3 (CDH3), also known as P-Cadherin, is a calcium-dependent cell-cell adhesion glycoprotein composed of five extracellular cadherin repeats, a transmembrane region and a highly conserved cytoplasmic tail. It is associated with some types of tumors. Epithelial cell adhesion molecule (EpCAM) is a transmembrane glycoprotein mediating Ca²⁺-independent homotypic cell-cell adhesion in epithelia. EpCAM has oncogenic potential and appears to play a role in tumorigenesis and metastasis of carcinomas.

[93] Further, it is envisaged as optionally but advantageously in the context of the present invention that the multichain multitargeting antigen-binding molecule is provided with a spacer, preferably a globular protein structure such as a scFc domain or dimerized Fc domain such as heteroFc, which also increases the molecule's half-life and enables intravenous dosing that is administrated only once every 5 week, once every two weeks, once every three weeks or even once every four weeks, or less frequently.

[94] In order to determine the epitope(s) of preferred multichain multitargeting antigen-binding molecules according to the present invention directed, e.g. to the CDH3, MSLN or CD20 epitope, mapping was conducted as described herein. Preferred bispecific antigen-binding molecules having a 10 target binder for CD20 are directed to all the of the epitope cluster E1A, E2B and E2C. An epitope cluster is understood herein as a stretch of amino acids (as disclosed herein and defined by their position according to the Kabat) within a target (as disclosed herein and defined by their position according to the Kabat) to which target a the whole the a target binder of a multichain multitargeting bispecific antigen-binding molecule as described herein does essentially no longer bind, if said stretch 15 of amino acid of the human target is replaced by a corresponding stretch of amino acids of the murine target. Therefore, said method of epitope clusters is understood herein as murine chimere sequence analysis. The method has been described, e.g. by Münz et al. Cancer Cell International 2010, 10:44 and was applied as described in detail in the examples with respect to CDH3 and MSLN.

[95] The preferred epitope cluster is D4B for CDH3 as described herein and E1 for MSLN as 20 described herein. As exemplified in the examples, selectivity gaps of multichain multitargeting bispecific antigen-binding molecules of the present invention (with respect to comparable monotargeting bispecific antigen-binding molecules) are typically even larger and, hence, more preferably, if the MSLN target binder addresses the E1 epitope cluster and if the CDH3 target binder addresses the D4B epitope cluster. While addressing other epitope clusters also leads to surprisingly 25 high selectivity gaps and the associated advantages in terms of efficacy and tolerability/safety, selectivity gaps are especially high and, thus preferred for molecules which comprise target binders which address E1 and D4B. Such molecules comprise, for example, a molecule with a MSLN target binder comprising CDR H1-H3 of SEQ ID NO 774 to 776 and CDR L1-L3 of 777 to 779 (and corresponding VH and VL of 780 and 781), CDR H1-H3 of SEQ ID NO 782 to 784 and CDR L1-L3 30 of 785 to 787 (and corresponding VH and VL of 788 and 789), CDR H1-H3 of SEQ ID NO 806 to 808 and CDR L1-L3 of 809 to 811 (and corresponding VH and VL of 812 and 813), CDR H1-H3 of SEQ ID NO 838 to 840 and CDR L1-L3 of 841 to 843 (and corresponding VH and VL of 844 and 845), CDR H1-H3 of SEQ ID NO 862 to 864 and CDR L1-L3 of 865 to 867 (and corresponding VH and VL of 868 and 869), CDR H1-H3 of SEQ ID NO 894 to 896 and CDR L1-L3 of 897 to 899 (and 35 corresponding VH and VL of 900 and 901), CDR H1-H3 of SEQ ID NO 950 to 952 and CDR L1-L3 of 953 to 955 (and corresponding VH and VL of 956 and 957), CDR H1-H3 of SEQ ID NO 1030 to

1032 and CDR L1-L3 of 1033 to 1035 (and corresponding VH and VL of 1036 and 1037), or CDR H1-H3 of SEQ ID NO 86 to 88 and CDR L1-L3 of 89 to 91 (and corresponding VH of 92 and VL 93 or 94). A preferred example for a CDH3 binder binding to the preferred DB4 epitope cluster comprises CDR H1-H3 of SEQ ID NO 194, 432 and 196 and CDR L1-L3 of 197 to 199 (and
5 corresponding VH and VL of 433 and 200). Further target binder which preferably bind to the preferred epitope cluster of D4B are, e.g., identified herein as CH3 15-E11 CC and CH3 24-D7 CC.

[96] It is particular surprising that a multichain multitargeting antigen-binding molecule according to the present invention is capable, to bind, preferably simultaneously to two different targets. Simultaneous binding has been demonstrated herein for several targets. However, this is surprising
10 given the typically typical distance between the targets. For example, CD20 comprises two small extra cellular domains of only 6 aa and 47 aa. In contrast, CD22 comprises a 7 Ig domain long extracellular domain with 676 aa. However, despite the significantly different extracellular size and setup, a multichain multitargeting antigen-binding molecule according to the present invention may successfully address both TAAs CD20 and CD22 at the same time for the benefit of increased efficacy
15 and less toxicity.

[97] It is envisaged in the context of the present invention, that preferred multichain multitargeting antigen-binding molecules do not only show a favorable ratio of cytotoxicity to affinity, but additionally show sufficient stability characteristics in order to facilitate practical handling in
20 formulating, storing and administrating said constructs. Sufficient stability is, for example, characterized by a high monomer content (i.e. non-aggregated and/or non-associated, native molecule) after standard preparation, such as at least 65% as determined by preparative size exclusion chromatography (SEC), more preferably at least 70% and even more preferably at least 75%. Also, the turbidity measured, e.g., at 340 nm as optical absorption at a concentration of 2.5 mg/ml should,
25 preferably, be equal to or lower than 0.025, more preferably 0.020, e.g., in order to conclude to the essential absence of undesired aggregates. Advantageously, high monomer content is maintained after incubation in stress conditions such as freeze/thaw or incubation at 37 or 40°C. Even more, multichain multitargeting antigen-binding molecules according to the present invention typically have a thermal stability which is at least comparable or even higher than that of bispecific antigen-binding molecules
30 which have only one target binding domain but otherwise comprise a CD3 binding domain and, a half-life extending scFc domain, i.e. which are structurally less complex. The skilled person would expect that a structurally more complex protein-based molecule was more prone to thermal and other degradation, i.e. be less thermal stable. However, surprisingly the contrary is the case, a multichain multitargeting bispecific antigen-binding molecule according to the present invention shows at least
35 comparable or even better thermal stability than single chain molecules. The molecules of the invention when tested also regarding long-term storage stability and freeze-thaw stability

advantageously exhibit at least comparable or even better characteristics as single chain molecules with the same binding domains. Preferably, molecules of the invention also show less monomer decrease after storage, and higher protein homogeneity than a respective single chain bispecific antigen-binding molecule, i.e. comprising the same target and CD3 binders, e.g. as disclosed herein.

5 [98] In an embodiment, the present invention provides a multichain multitargeting bispecific antigen-binding molecule comprising all four such domains. In a preferred embodiment, the domains under (i.), (ii.), (iii.) and (iv.) are arranged as described in Figures 1, 2 and 3.

10 [99] The term “polypeptide” is understood herein as an organic polymer which comprises at least one continuous, unbranched amino acid chain. In the context of the present invention, a polypeptide comprising more than one amino acid chain is likewise envisaged. An amino acid chain of a polypeptide typically comprises at least 50 amino acids, preferably at least 100, 200, 300, 400 or 500 amino acids. It is also envisaged in the context of the present invention that an amino acid chain of a polymer is linked to an entity which is not composed of amino acids.

15 [100] The term “antigen-binding polypeptide” according to the present invention is preferably a polypeptide which immuno-specifically binds to its target or antigen. It typically comprises the heavy chain variable region (VH) and/or the light chain variable region (VL) of an antibody, or comprises domains derived therefrom. A polypeptide according to the invention comprises the minimum structural requirements of an antibody which allow for immuno-specific target binding. This minimum requirement may e.g. be defined by the presence of at least three light chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VL region) and/or three heavy chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VH region), preferably of all six CDRs. An antigen-binding molecule of the present invention is preferably a T-cell engaging polypeptide which may hence be characterized by the presence of three or six CDRs in either one or both binding domains, and the skilled person knows where (in which order) those CDRs are located within the binding domain. Preferably, an “antigen-binding molecule” is understood as an “antigen-binding polypeptide” in the context of the present invention. In an alternative 25 embodiment, an antigen-binding polypeptide of the present invention may be an aptamer.

30 [101] Alternatively, a molecule in the context of the present invention, is an antigen-binding polypeptide which corresponds to an “antibody construct” which typically refers to a molecule in which the structure and/or function is/are based on the structure and/or function of an antibody, e.g., of a full-length or whole immunoglobulin molecule. An antigen-binding molecule is hence capable of binding to its specific target or antigen and/or is/are drawn from the variable heavy chain (VH) and/or variable light chain (VL) domains of an antibody or fragment thereof. Furthermore, the domain which binds to its binding partner according to the present invention is understood herein as a binding domain of an antigen-binding molecule according to the invention. Typically, a binding domain 35 according to the present invention comprises the minimum structural requirements of an antibody

which allow for the target binding. This minimum requirement may *e.g.* be defined by the presence of at least the three light chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VL region) and/or the three heavy chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VH region), preferably of all six CDRs. An alternative approach to define the minimal structure requirements of an antibody is the definition of the epitope of the antibody within the structure of the specific target, respectively, the protein domain of the target protein composing the epitope region (epitope cluster) or by reference to a specific antibody competing with the epitope of the defined antibody. The antibodies on which the constructs according to the invention are based include for example monoclonal, recombinant, chimeric, deimmunized, humanized and human antibodies.

5 [102] In the context of the present invention, a polypeptide of the present invention binds to its respective target structure in a particular manner. Preferably, a polypeptide according to the present invention comprises one paratope per binding domain which specifically or immuno-specifically binds to”, “(specifically or immuno-specifically) recognizes”, or “(specifically or immuno-specifically) reacts with” its respective target structure. This means in accordance with this invention that a polypeptide or a binding domain thereof interacts or (immuno-)specifically interacts with a given epitope on the target molecule (antigen) and CD3, respectively. This interaction or association occurs more frequently, more rapidly, with greater duration, with greater affinity, or with some combination of these parameters, to an epitope on the specific target than to alternative substances (non-target molecules). Because of the sequence similarity between homologous proteins in different species, a binding domain that (immuno-) specifically binds to its target (such as a human target) may, however, cross-react with homologous target molecules from different species (such as, from non-human primates). The term “specific / immuno-specific binding” can hence include the binding of a binding domain to epitopes and/or structurally related epitopes in more than one species. The term “(immuno-) selectively binds” does exclude the binding to structurally related epitopes.

25 [103] The binding domain of an antigen-binding molecule according to the invention may *e.g.* comprise the above referred groups of CDRs. Preferably, those CDRs are comprised in the framework of an antibody light chain variable region (VL) and an antibody heavy chain variable region (VH); however, it does not have to comprise both. Fd fragments, for example, have two VH regions and often retain some antigen-binding function of the intact antigen-binding domain.

30 Additional examples for the format of antibody fragments, antibody variants or binding domains include (1) a Fab fragment, a monovalent fragment having the VL, VH, CL and CH1 domains; (2) a F(ab')₂ fragment, a bivalent fragment having two Fab fragments linked by a disulfide bridge at the hinge region; (3) an Fd fragment having the two VH and CH1 domains; (4) an Fv fragment having the VL and VH domains of a single arm of an antibody, (5) a dAb fragment (Ward et al., (1989) Nature 341 :544-546), which has a VH domain; (6) an isolated complementarity determining region (CDR), and (7) a single chain Fv (scFv) , the latter being preferred (for example, derived from an

scFv-library). Examples for embodiments of antigen-binding molecules according to the invention are e.g. described in WO 00/006605, WO 2005/040220, WO 2008/119567, WO 2010/037838, WO 2013/026837, WO 2013/026833, US 2014/0308285, US 2014/0302037, WO 2014/144722, WO 2014/151910, and WO 2015/048272.

5 [104] Also, within the definition of “binding domain” or “domain which binds” are fragments of full-length antibodies, such as VH, VHH, VL, (s)dAb, Fv, Fd, Fab, Fab’, F(ab’)2 or “r IgG” (“half antibody”). Antigen-binding molecules according to the invention may also comprise modified fragments of antibodies, also called antibody variants, such as scFv, di-scFv or bi(s)-scFv, scFv-Fc, scFv-zipper, scFab, Fab₂, Fab₃, diabodies, single chain diabodies, tandem diabodies (Tandab’s),
10 tandem di-scFv, tandem tri-scFv, “multibodies” such as triabodies or tetrabodies, and single domain antibodies such as nanobodies or single variable domain antibodies comprising merely one variable domain, which may be VHH, VH or VL, that specifically bind an antigen or epitope independently of other V regions or domains. Typically, a binding domain of the present invention comprises a paratope which facilitates the binding to its binding partner.

15 [105] As used herein, the terms "single-chain Fv," "single-chain antibodies" or "scFv" refer to single polypeptide chain antibody fragments that comprise the variable regions from both the heavy and light chains, but lack the constant regions. Generally, a single-chain antibody further comprises a polypeptide linker between the VH and VL domains which enables it to form the desired structure which would allow for antigen binding. Single chain antibodies are discussed in detail by Pluckthun
20 in *The Pharmacology of Monoclonal Antibodies*, vol. 113, Rosenberg and Moore eds. Springer-Verlag, New York, pp. 269-315 (1994). Various methods of generating single chain antibodies are known, including those described in U.S. Pat. Nos. 4,694,778 and 5,260,203; International Patent Application Publication No. WO 88/01649; Bird (1988) *Science* 242:423-442; Huston *et al.* (1988) *Proc. Natl. Acad. Sci. USA* 85:5879-5883; Ward *et al.* (1989) *Nature* 334:54454; Skerra *et al.* (1988)
25 *Science* 242:1038-1041. In specific embodiments, single-chain antibodies can also be bispecific, multispecific, human, and/or humanized and/or synthetic.

[106] In the context of the present invention, a paratope is understood as an antigen-binding site which is a part of a polypeptide as described herein and which recognizes and binds to an antigen. A paratope is typically a small region of about at least 5 amino acids. A paratope as understood herein
30 typically comprises parts of antibody-derived heavy (VH) and light chain (VL) sequences. Each binding domain of a molecule according to the present invention is provided with a paratope comprising a set of 6 complementarity-determining regions (CDR loops) with three of each being comprised within the antibody-derived VH and VL sequence, respectively.

[107] Furthermore, the definition of the term “antigen-binding molecule” includes preferably
35 polyvalent / multivalent constructs and, thus, bispecific molecules, wherein bispecific means that

they specifically bind to two cell types comprising distinctive antigenic structures, i.e. target cell(s) and effector cell(s). As the antigen-binding molecules of the present invention are preferably multichain multitargeting, they are typically as well as polyvalent / multivalent molecules, i.e. they specifically bind more than two antigenic structures, preferably four distinct binding domains in the context of the present invention which are two target binding domains and two CD3 binding domains. The term “multichain multitargeting bispecific antigen-binding molecule” comprises the terms “multichain multitargeting bispecific T-cell engager molecule” and “multichain multitargeting bispecific T-cell engager polypeptide (MMBiTEP)”. A preferred “multichain multitargeting bispecific antigen-binding molecule” is a “multichain multitargeting bispecific T-cell engager molecule” or a “multichain multitargeting bispecific T-cell engager polypeptide (MMBiTEP)”. The term “multichain multitargeting bispecific T-cell engager molecule” is understood to comprise the term “multichain multitargeting bispecific T-cell engager polypeptide. Moreover, the definition of the term “antigen-binding molecule” includes molecules comprising only one polypeptide chain as well as molecules consisting of more than one polypeptide chain, which chains can be either identical (homodimers, homotrimers or homo oligomers) or different (heterodimer, heterotrimer or heterooligomer). Such molecules comprising more than one polypeptide chain, i.e. typically two chains, have these chains typically attached to each other as heterodimers via charged pair binding, e.g. within a heteroFc entity which serves as a spacer and half-life extending moiety in between the two bispecific entities as described herein. Examples for the above identified antigen-binding molecules, e.g. antibody-based molecules and variants or derivatives thereof are described *inter alia* in Harlow and Lane, *Antibodies a laboratory manual*, CSHL Press (1988) and *Using Antibodies: a laboratory manual*, CSHL Press (1999), Kontermann and Dübel, *Antibody Engineering*, Springer, 2nd ed. 2010 and Little, *Recombinant Antibodies for Immunotherapy*, Cambridge University Press 2009.

[108] The term “bispecific” as used herein refers to an antigen-binding molecule which is “at least bispecific”, *i.e.*, it addresses two different cell types, i.e. target and effector cells, and comprises at least a first and third binding domain and a second and fourth binding domain, wherein at least two binding domains bind to two antigens or targets selected preferably from CD20, CD22, FLT3, MSLN, CDH3, CLL1 and EpCAM, and the other two binding domains of the same molecule bind to another antigen (here: CD3) on an effector cell, typically on a T cell. Accordingly, antigen-binding molecules according to the invention comprise specificities for at least two different antigens or targets. For example, two domains do preferably not bind to an extracellular epitope of CD3 ϵ of one or more of the species as described herein.

[109] The term “target cell surface antigen” refers to an antigenic structure expressed by a cell and which is present at the cell surface such that it is accessible for an antigen-binding molecule as described herein. A preferred target cell surface antigen in the context of the present invention is a

tumor associated antigen (TAA). It may be a protein, preferably the extracellular portion of a protein, or a carbohydrate structure, preferably a carbohydrate structure of a protein, such as a glycoprotein. It is preferably a tumor antigen. The term “bispecific antigen-binding molecule” of the invention also encompasses bispecific multichain multitargeting antigen-binding molecules such as tritargeting antigen-binding molecules, the latter ones including three binding domains, or constructs having more than three (e.g. four, five...) specificities.

[110] Preferred in the context of the present invention is a molecule which is “multitargeting”, which is understood herein to be “at least targeting two targets (e.g. TAAs) per molecule of the invention typically per target cell”. In this regard, a multitargeting molecule such as an antigen-binding molecule is specific for two – typically identical- effector structures on an effector cell such as CD3, more preferably CD3 ϵ (CD3 ϵ , which is comprised whenever reference is made to the “CD3” in the present invention), and at least two target cell surface antigens. Said specificity is conferred by respective binding domains as defined herein. Typically, “multitargeting” refers to a molecule which is specific for at least two (preferably different) target cell surface antigens (e.g. TAAs) which confers preferred properties of a multitargeting antigen-binding molecule according to the present invention, namely mitigation of antigen loss and increase of selectivity, i.e. selectivity for killing target cells which co-express the targets for which the molecule of the invention has binding domains and which target cells are associated with a disease. Thereby, the therapeutic window of the molecule of the invention is increased with respect to monotargeting bispecific molecules which typically leads to higher drug tolerability as demonstrated herein.

[111] A T-cell engaging antigen-binding molecule, e.g. a multichain polypeptide, according to the present invention is preferably bispecific which is understood herein to typically comprise one domain binding to at least one target antigen and another domain binding to CD3. Hence, it does not occur naturally, and it is markedly different in its function from naturally occurring products. A polypeptide in accordance with the invention is hence an artificial “hybrid” polypeptide comprising at least two distinct binding domains with different specificities and is, thus, bispecific. Bispecific antigen-binding molecules can be produced by a variety of methods including fusion of hybridomas or linking of Fab' fragments. See, e.g., Songsivilai & Lachmann, Clin. Exp. Immunol. 79:315-321 (1990).

[112] The at least four binding domains and the variable domains (VH / VL) of the antigen-binding molecule of the present invention typically comprise peptide linkers (spacer peptides). The term “peptide linker” comprises in accordance with the present invention an amino acid sequence by which the amino acid sequences of one (variable and/or binding) domain and another (variable and/or binding) domain of the antigen-binding molecule of the invention are linked with each other. The peptide linker between the first and the second binding domain and the third and the fourth domain, wherein the first and the third domain are preferably capable to bind simultaneously to two

targets, which are preferably different targets (e.g. TAA1 and TAA2) preferably on the same cell, are preferably flexible and of limited length, e.g. of 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17 or 18 amino acids. The peptide linkers can also be used to fuse the spacer to the other domains of the antigen-binding molecule of the invention. An essential technical feature of such peptide linker is that it does not comprise any polymerization activity. Among the suitable peptide linkers are those described in U.S. Patents 4,751,180 and 4,935,233 or WO 88/09344. The peptide linkers can also be used to attach other domains or modules or regions (such as half-life extending domains) to the antigen-binding molecule of the invention. However, typically the linker between the first and the second target binding domain differs from the intra-binder linker which links the VH and VL within the target binding domain. Said difference is the linker between the first and the second binding domain having one amino acid more than intra-binder linkers, e.g. six and five amino acids, respectively, such as SGGGS versus GGGGS. This confers surprisingly flexibility and stability at the same time in the specific antigen-binding molecule format as described herein. The spacer (or synonymously spacer entity) between the two bispecific entities as described herein is a specific embodiment of a linker because a spacer also functions as a linker because it contributes to linking the two bispecific entities to preferably build at least one continuous polypeptide chain comprising the four binding domains or parts thereof. However, in addition, the spacer functions as an entity which spaces the two bispecific entities sterically apart. Accordingly, a spacer in the context of the present invention is a specific embodiment of a linker which -together with two further short and flexible linkers on each end- contributes to linking the two binding domains (of two different bispecific entities) but first and foremost spaces them apart in such a way that the two bispecific entities can advantageously act as described herein, e.g. show a surprisingly high selectivity gap.

[113] The antigen-binding molecules of the present invention are preferably “*in vitro* generated antigen-binding molecules”. This term refers to an antigen-binding molecule according to the above definition where all or part of the variable region (e.g., at least one CDR) is generated in a non-immune cell selection, e.g., an *in vitro* phage display, protein chip or any other method in which candidate sequences can be tested for their ability to bind to an antigen. This term thus preferably excludes sequences generated solely by genomic rearrangement in an immune cell in an animal. A “recombinant antibody” is an antibody made through the use of recombinant DNA technology or genetic engineering.

[114] The term “monoclonal antibody” (mAb) or monoclonal antibody from which an antigen-binding molecule as used herein is derived refers to an antibody obtained from a population of substantially homogeneous antibodies, *i.e.*, the individual antibodies comprising the population are identical except for possible naturally occurring mutations and/or post-translation modifications (e.g., isomerizations, amidations) that may be present in minor amounts. Monoclonal antibodies are highly specific, being directed against a single antigenic side or determinant on the antigen, in

contrast to conventional (polyclonal) antibody preparations which typically include different antibodies directed against different determinants (or epitopes). In addition to their specificity, the monoclonal antibodies are advantageous in that they are synthesized by the hybridoma culture, hence uncontaminated by other immunoglobulins. The modifier “monoclonal” indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies, and is not to be construed as requiring production of the antibody by any particular method.

[115] For the preparation of monoclonal antibodies, any technique providing antibodies produced by continuous cell line cultures can be used. For example, monoclonal antibodies to be used may be made by the hybridoma method first described by Koehler *et al.*, Nature, 256: 495 (1975), or may be made by recombinant DNA methods (see, e.g., U.S. Patent No. 4,816,567). Examples for further techniques to produce human monoclonal antibodies include the trioma technique, the human B-cell hybridoma technique (Kozbor, Immunology Today 4 (1983), 72) and the EBV-hybridoma technique (Cole *et al.*, Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc. (1985), 77-96).

[116] Hybridomas can then be screened using standard methods, such as enzyme-linked immunosorbent assay (ELISA) and surface plasmon resonance analysis, e.g. Biacore™ to identify one or more hybridomas that produce an antibody that specifically binds with a specified antigen. Any form of the relevant antigen may be used as the immunogen, e.g., recombinant antigen, naturally occurring forms, any variants or fragments thereof, as well as an antigenic peptide thereof. Surface plasmon resonance as employed in the Biacore system can be used to increase the efficiency of phage antibodies which bind to an epitope of a target cell surface antigen (Schier, Human Antibodies Hybridomas 7 (1996), 97-105; Malmberg, J. Immunol. Methods 183 (1995), 7-13).

[117] Another exemplary method of making monoclonal antibodies includes screening protein expression libraries, e.g., phage display or ribosome display libraries. Phage display is described, for example, in Ladner *et al.*, U.S. Patent No. 5,223,409; Smith (1985) Science 228:1315-1317, Clackson *et al.*, Nature, 352: 624-628 (1991) and Marks *et al.*, J. Mol. Biol., 222: 581-597 (1991).

[118] In addition to the use of display libraries, the relevant antigen can be used to immunize a non-human animal, e.g., a rodent (such as a mouse, hamster, rabbit or rat). In one embodiment, the non-human animal includes at least a part of a human immunoglobulin gene. For example, it is possible to engineer mouse strains deficient in mouse antibody production with large fragments of the human Ig (immunoglobulin) loci. Using the hybridoma technology, antigen-specific monoclonal antibodies derived from the genes with the desired specificity may be produced and selected. See, e.g., XENOMOUSE™, Green *et al.* (1994) Nature Genetics 7:13-21, US 2003-0070185, WO 96/34096, and WO 96/33735.

[119] A monoclonal antibody can also be obtained from a non-human animal, and then modified, e.g., humanized, deimmunized, rendered chimeric etc., using recombinant DNA techniques known in the art. Examples of modified antigen-binding molecules include humanized variants of non-human antibodies, "affinity matured" antibodies (see, e.g. Hawkins et al. J. Mol. Biol. 254, 889-896 (1992) and Lowman *et al.*, Biochemistry 30, 10832- 10837 (1991)) and antibody mutants with altered effector function(s) (see, e.g., US Patent 5,648,260, Kontermann and Dübel (2010), *loc. cit.* and Little (2009), *loc. cit.*).

[120] In immunology, affinity maturation is the process by which B cells produce antibodies with increased affinity for antigen during the course of an immune response. With repeated exposures to the same antigen, a host will produce antibodies of successively greater affinities. Like the natural prototype, the *in vitro* affinity maturation is based on the principles of mutation and selection. The *in vitro* affinity maturation has successfully been used to optimize antibodies, antigen-binding molecules, and antibody fragments. Random mutations inside the CDRs are introduced using radiation, chemical mutagens or error-prone PCR. In addition, the genetic diversity can be increased by chain shuffling. Two or three rounds of mutation and selection using display methods like phage display usually results in antibody fragments with affinities in the low nanomolar range.

[121] A preferred type of an amino acid substitutional variation of the antigen-binding molecules involves substituting one or more hypervariable region residues of a parent antibody (e. g. a humanized or human antibody). Generally, the resulting variant(s) selected for further development will have improved biological properties relative to the parent antibody from which they are generated. A convenient way for generating such substitutional variants involves affinity maturation using phage display. Briefly, several hypervariable region sides (e. g. 6-7 sides) are mutated to generate all possible amino acid substitutions at each side. The antibody variants thus generated are displayed in a monovalent fashion from filamentous phage particles as fusions to the gene III product of M13 packaged within each particle. The phage-displayed variants are then screened for their biological activity (e. g. binding affinity) as herein disclosed. In order to identify candidate hypervariable region sides for modification, alanine scanning mutagenesis can be performed to identify hypervariable region residues contributing significantly to antigen binding. Alternatively, or additionally, it may be beneficial to analyze a crystal structure of the antigen-antibody complex to identify contact points between the binding domain and, e.g., human CS1, BCMA, CD20, CD22, FLT3, CD123, CDH3, MSLN, CLL1 or EpCAM. Such contact residues and neighbouring residues are candidates for substitution according to the techniques elaborated herein. Once such variants are generated, the panel of variants is subjected to screening as described herein and antibodies with superior properties in one or more relevant assays may be selected for further development.

[122] The monoclonal antibodies and antigen-binding molecules of the present invention specifically include "chimeric" antibodies (immunoglobulins) in which a portion of the heavy and/or

light chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a particular antibody class or subclass, while the remainder of the chain(s) is/are identical with or homologous to corresponding sequences in antibodies derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies, so long as they exhibit the desired biological activity (U.S. Patent No. 4,816,567; Morrison *et al.*, Proc. Natl. Acad. Sci. USA, 81: 6851-6855 (1984)). Chimeric antibodies of interest herein include “primitized” antibodies comprising variable domain antigen-binding sequences derived from a non-human primate (e.g., Old World Monkey, Ape etc.) and human constant region sequences. A variety of approaches for making chimeric antibodies have been described. See *e.g.*, Morrison *et al.*, Proc. Natl. Acad. Sci. U.S.A. 81:6851, 1985; Takeda *et al.*, Nature 314:452, 1985, Cabilly *et al.*, U.S. Patent No. 4,816,567; Boss *et al.*, U.S. Patent No. 4,816,397; Tanaguchi *et al.*, EP 0171496; EP 0173494; and GB 2177096.

[123] An antibody, antigen-binding molecule, antibody fragment or antibody variant may also be modified by specific deletion of human T cell epitopes (a method called “deimmunization”) by the methods disclosed for example in WO 98/52976 or WO 00/34317. Briefly, the heavy and light chain variable domains of an antibody can be analyzed for peptides that bind to MHC class II; these peptides represent potential T cell epitopes (as defined in WO 98/52976 and WO 00/34317). For detection of potential T cell epitopes, a computer modeling approach termed “peptide threading” can be applied, and in addition a database of human MHC class II binding peptides can be searched for motifs present in the VH and VL sequences, as described in WO 98/52976 and WO 00/34317. These motifs bind to any of the 18 major MHC class II DR allotypes, and thus constitute potential T cell epitopes. Potential T cell epitopes detected can be eliminated by substituting small numbers of amino acid residues in the variable domains, or preferably, by single amino acid substitutions. Typically, conservative substitutions are made. Often, but not exclusively, an amino acid common to a position in human germline antibody sequences may be used. Human germline sequences are disclosed *e.g.* in Tomlinson, *et al.* (1992) J. Mol. Biol. 227:776-798; Cook, G.P. *et al.* (1995) Immunol. Today Vol. 16 (5): 237-242; and Tomlinson *et al.* (1995) EMBO J. 14: 14:4628-4638. The VBASE directory provides a comprehensive directory of human immunoglobulin variable region sequences (compiled by Tomlinson, L.A. *et al.* MRC Centre for Protein Engineering, Cambridge, UK). These sequences can be used as a source of human sequence, *e.g.*, for framework regions and CDRs. Consensus human framework regions can also be used, for example as described in US Patent No. 6,300,064.

[124] “Humanized” antibodies, antigen-binding molecules, variants or fragments thereof (such as Fv, Fab, Fab', F(ab')₂ or other antigen-binding subsequences of antibodies) are antibodies or immunoglobulins of mostly human sequences, which contain (a) minimal sequence(s) derived from non-human immunoglobulin. For the most part, humanized antibodies are human immunoglobulins (recipient antibody) in which residues from a hypervariable region (also CDR) of the recipient are

replaced by residues from a hypervariable region of a non-human (*e.g.*, rodent) species (donor antibody) such as mouse, rat, hamster or rabbit having the desired specificity, affinity, and capacity. In some instances, Fv framework region (FR) residues of the human immunoglobulin are replaced by corresponding non-human residues. Furthermore, “humanized antibodies” as used herein may also
5 comprise residues which are found neither in the recipient antibody nor the donor antibody. These modifications are made to further refine and optimize antibody performance. The humanized antibody may also comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. For further details, see Jones *et al.*, *Nature*, 321: 522-525 (1986); Reichmann *et al.*, *Nature*, 332: 323-329 (1988); and Presta, *Curr. Op. Struct. Biol.*, 2: 593-596
10 (1992).

[125] Humanized antibodies or fragments thereof can be generated by replacing sequences of the Fv variable domain that are not directly involved in antigen binding with equivalent sequences from human Fv variable domains. Exemplary methods for generating humanized antibodies or fragments thereof are provided by Morrison (1985) *Science* 229:1202-1207; by Oi *et al.* (1986) *BioTechniques*
15 4:214; and by US 5,585,089; US 5,693,761; US 5,693,762; US 5,859,205; and US 6,407,213. Those methods include isolating, manipulating, and expressing the nucleic acid sequences that encode all or part of immunoglobulin Fv variable domains from at least one of a heavy or light chain. Such nucleic acids may be obtained from a hybridoma producing an antibody against a predetermined target, as described above, as well as from other sources. The recombinant DNA encoding the humanized
20 antibody molecule can then be cloned into an appropriate expression vector.

[126] Humanized antibodies may also be produced using transgenic animals such as mice that express human heavy and light chain genes, but are incapable of expressing the endogenous mouse immunoglobulin heavy and light chain genes. Winter describes an exemplary CDR grafting method that may be used to prepare the humanized antibodies described herein (U.S. Patent No. 5,225,539).
25 All of the CDRs of a particular human antibody may be replaced with at least a portion of a non-human CDR, or only some of the CDRs may be replaced with non-human CDRs. It is only necessary to replace the number of CDRs required for binding of the humanized antibody to a predetermined antigen.

[127] A humanized antibody can be optimized by the introduction of conservative substitutions, consensus sequence substitutions, germline substitutions and/or back mutations. Such altered immunoglobulin molecules can be made by any of several techniques known in the art, (*e.g.*, Teng *et al.*, *Proc. Natl. Acad. Sci. U.S.A.*, 80: 7308-7312, 1983; Kozbor *et al.*, *Immunology Today*, 4: 7279, 1983; Olsson *et al.*, *Meth. Enzymol.*, 92: 3-16, 1982, and EP 239 400).
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[128] The term “human antibody”, “human antigen-binding molecule” and “human binding domain”
35 includes antibodies, antigen-binding molecules and binding domains having antibody regions such as

variable and constant regions or domains which correspond substantially to human germline immunoglobulin sequences known in the art, including, for example, those described by Kabat *et al.* (1991) (*loc. cit.*). The human antibodies, antigen-binding molecules or binding domains of the invention may include amino acid residues not encoded by human germline immunoglobulin sequences (e.g., mutations introduced by random or site-specific mutagenesis *in vitro* or by somatic mutation *in vivo*), for example in the CDRs, and in particular, in CDR3. The human antibodies, antigen-binding molecules or binding domains can have at least one, two, three, four, five, or more positions replaced with an amino acid residue that is not encoded by the human germline immunoglobulin sequence. The definition of human antibodies, antigen-binding molecules and binding domains as used herein also contemplates fully human antibodies, which include only non-artificially and/or genetically altered human sequences of antibodies as those can be derived by using technologies or systems such as the Xenomouse. Preferably, a “fully human antibody” does not include amino acid residues not encoded by human germline immunoglobulin sequences.

[129] In some embodiments, the antigen-binding molecules of the invention are “isolated” or “substantially pure” antigen-binding molecules. “Isolated” or “substantially pure”, when used to describe the antigen-binding molecules disclosed herein, means an antigen-binding molecule that has been identified, separated and/or recovered from a component of its production environment. Preferably, the antigen-binding molecule is free or substantially free of association with all other components from its production environment. Contaminant components of its production environment, such as that resulting from recombinant transfected cells, are materials that would typically interfere with diagnostic or therapeutic uses for the polypeptide, and may include enzymes, hormones, and other proteinaceous or non-proteinaceous solutes. The antigen-binding molecules may e.g. constitute at least about 5%, or at least about 50% by weight of the total protein in a given sample. It is understood that the isolated protein may constitute from 5% to 99.9% by weight of the total protein content, depending on the circumstances. The polypeptide may be made at a significantly higher concentration through the use of an inducible promoter or high expression promoter, such that it is made at increased concentration levels. The definition includes the production of an antigen-binding molecule in a wide variety of organisms and/or host cells that are known in the art. In preferred embodiments, the antigen-binding molecule will be purified (1) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (2) to homogeneity by SDS-PAGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Ordinarily, however, an isolated antigen-binding molecule will be prepared by at least one purification step.

[130] The term “binding domain” characterizes in connection with the present invention a domain which (specifically) binds to / interacts with / recognizes a given target epitope or a given target side on the target molecules (antigens), e.g. CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, MSLN, or

EpCAM, and CD3, respectively. The structure and function of the typically first and third or second and fourth binding domain (recognizing e.g. CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, MSLN, or EpCAM), and preferably also the structure and/or function of the effector binding domain (typically the second and fourth or first and third binding domain recognizing CD3), is/are based on the structure and/or function of an antibody, e.g. of a full-length or whole immunoglobulin molecule, and/or is/are drawn from the variable heavy chain (VH) and/or variable light chain (VL) domains of an antibody or fragment thereof. Preferably the target cell surface antigen(s) binding domain(s) is/are characterized by the presence of three light chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VL region) and/or three heavy chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VH region). The effector (typically CD3) binding domain preferably also comprises the minimum structural requirements of an antibody which allow for the target binding. More preferably, the second binding domain comprises at least three light chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VL region) and/or three heavy chain CDRs (i.e. CDR1, CDR2 and CDR3 of the VH region). It is envisaged that the first and/or second binding domain is produced by or obtainable by phage-display or library screening methods rather than by grafting CDR sequences from a pre-existing (monoclonal) antibody into a scaffold.

[131] According to the present invention, binding domains are in the form of one or more polypeptides. Such polypeptides may include proteinaceous parts and non-proteinaceous parts (e.g. chemical linkers or chemical cross-linking agents such as glutaraldehyde). Proteins (including fragments thereof, preferably biologically active fragments, and peptides, usually having less than 30 amino acids) comprise two or more amino acids coupled to each other via a covalent peptide bond (resulting in a chain of amino acids).

[132] The term "polypeptide" as used herein describes a group of molecules, which usually consist of more than 30 amino acids. Polypeptides may further form multimers such as dimers, trimers and higher oligomers, *i.e.*, consisting of more than one polypeptide molecule. Polypeptide molecules forming such dimers, trimers etc. may be identical or non-identical. The corresponding higher order structures of such multimers are, consequently, termed homo- or heterodimers, homo- or heterotrimers etc. An example for a heteromultimer is an antibody molecule, which, in its naturally occurring form, consists of two identical light polypeptide chains and two identical heavy polypeptide chains. The terms "peptide", "polypeptide" and "protein" also refer to naturally modified peptides / polypeptides / proteins wherein the modification is effected *e.g.* by post-translational modifications like glycosylation, acetylation, phosphorylation and the like. A "peptide", "polypeptide" or "protein" when referred to herein may also be chemically modified such as pegylated. Such modifications are well known in the art and described herein below.

[133] Preferably the binding domains which binds to any of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, and EpCAM, and/or the binding domains which binds to CD3 ϵ is/are

human binding domains. Antibodies and antigen-binding molecules comprising at least one human binding domain avoid some of the problems associated with antibodies or antigen-binding molecules that possess non-human such as rodent (*e.g.* murine, rat, hamster or rabbit) variable and/or constant regions. The presence of such rodent derived proteins can lead to the rapid clearance of the antibodies or antigen-binding molecules or can lead to the generation of an immune response against the antibody or antigen-binding molecule by a patient. In order to avoid the use of rodent derived antibodies or antigen-binding molecules, human or fully human antibodies / antigen-binding molecules can be generated through the introduction of human antibody function into a rodent so that the rodent produces fully human antibodies.

[134] The ability to clone and reconstruct megabase-sized human loci in yeast artificial chromosomes YACs and to introduce them into the mouse germline provides a powerful approach to elucidating the functional components of very large or crudely mapped loci as well as generating useful models of human disease. Furthermore, the use of such technology for substitution of mouse loci with their human equivalents could provide unique insights into the expression and regulation of human gene products during development, their communication with other systems, and their involvement in disease induction and progression.

[135] An important practical application of such a strategy is the “humanization” of the mouse humoral immune system. Introduction of human immunoglobulin (Ig) loci into mice in which the endogenous Ig genes have been inactivated offers the opportunity to study the mechanisms underlying programmed expression and assembly of antibodies as well as their role in B-cell development. Furthermore, such a strategy could provide an ideal source for production of fully human monoclonal antibodies (mAbs) – an important milestone towards fulfilling the promise of antibody therapy in human disease. Fully human antibodies or antigen-binding molecules are expected to minimize the immunogenic and allergic responses intrinsic to mouse or mouse-derivatized mAbs and thus to increase the efficacy and safety of the administered antibodies / antigen-binding molecules. The use of fully human antibodies or antigen-binding molecules can be expected to provide a substantial advantage in the treatment of chronic and recurring human diseases, such as inflammation, autoimmunity, and cancer, which require repeated compound administrations.

[136] One approach towards this goal was to engineer mouse strains deficient in mouse antibody production with large fragments of the human Ig loci in anticipation that such mice would produce a large repertoire of human antibodies in the absence of mouse antibodies. Large human Ig fragments would preserve the large variable gene diversity as well as the proper regulation of antibody production and expression. By exploiting the mouse machinery for antibody diversification and selection and the lack of immunological tolerance to human proteins, the reproduced human antibody repertoire in these mouse strains should yield high affinity antibodies against any antigen of

interest, including human antigens. Using the hybridoma technology, antigen-specific human mAbs with the desired specificity could be readily produced and selected. This general strategy was demonstrated in connection with the generation of the first Xenomouse mouse strains (see Green et al. *Nature Genetics* 7:13-21 (1994)). The Xenomouse strains were engineered with YACs containing 245 kb and 190 kb-sized germline configuration fragments of the human heavy chain locus and kappa light chain locus, respectively, which contained core variable and constant region sequences. The human Ig containing YACs proved to be compatible with the mouse system for both rearrangement and expression of antibodies and were capable of substituting for the inactivated mouse Ig genes. This was demonstrated by their ability to induce B cell development, to produce an adult-like human repertoire of fully human antibodies, and to generate antigen-specific human mAbs. These results also suggested that introduction of larger portions of the human Ig loci containing greater numbers of V genes, additional regulatory elements, and human Ig constant regions may recapitulate substantially the full repertoire that is characteristic of the human humoral response to infection and immunization. The work of Green et al. was recently extended to the introduction of greater than approximately 80% of the human antibody repertoire through introduction of megabase sized, germline configuration YAC fragments of the human heavy chain loci and kappa light chain loci, respectively. See Mendez *et al.* *Nature Genetics* 15:146-156 (1997) and U.S. patent application Ser. No. 08/759,620.

[137] The production of the Xenomouse animals is further discussed and delineated in U.S. patent applications Ser. No. 07/466,008, Ser. No. 07/610,515, Ser. No. 07/919,297, Ser. No. 07/922,649, Ser. No. 08/031,801, Ser. No. 08/112,848, Ser. No. 08/234,145, Ser. No. 08/376,279, Ser. No. 08/430,938, Ser. No. 08/464,584, Ser. No. 08/464,582, Ser. No. 08/463,191, Ser. No. 08/462,837, Ser. No. 08/486,853, Ser. No. 08/486,857, Ser. No. 08/486,859, Ser. No. 08/462,513, Ser. No. 08/724,752, and Ser. No. 08/759,620; and U.S. Pat. Nos. 6,162,963; 6,150,584; 6,114,598; 6,075,181, and 5,939,598 and Japanese Patent Nos. 3 068 180 B2, 3 068 506 B2, and 3 068 507 B2. See also Mendez *et al.* *Nature Genetics* 15:146-156 (1997) and Green and Jakobovits *J. Exp. Med.* 188:483-495 (1998), EP 0 463 151 B1, WO 94/02602, WO 96/34096, WO 98/24893, WO 00/76310, and WO 03/47336.

[138] In an alternative approach, others, including GenPharm International, Inc., have utilized a "minilocus" approach. In the minilocus approach, an exogenous Ig locus is mimicked through the inclusion of pieces (individual genes) from the Ig locus. Thus, one or more VH genes, one or more DH genes, one or more JH genes, a mu constant region, and a second constant region (preferably a gamma constant region) are formed into a construct for insertion into an animal. This approach is described in U.S. Pat. No. 5,545,807 to Surani *et al.* and U.S. Pat. Nos. 5,545,806; 5,625,825; 5,625,126; 5,633,425; 5,661,016; 5,770,429; 5,789,650; 5,814,318; 5,877,397; 5,874,299; and 6,255,458 each to Lonberg and Kay, U.S. Pat. Nos. 5,591,669 and 6,023,010 to Krimpenfort and

Berns, U.S. Pat. Nos. 5,612,205; 5,721,367; and 5,789,215 to Berns *et al.*, and U.S. Pat. No. 5,643,763 to Choi and Dunn, and GenPharm International U.S. patent application Ser. No. 07/574,748, Ser. No. 07/575,962, Ser. No. 07/810,279, Ser. No. 07/853,408, Ser. No. 07/904,068, Ser. No. 07/990,860, Ser. No. 08/053,131, Ser. No. 08/096,762, 5 Ser. No. 08/155,301, Ser. No. 08/161,739, Ser. No. 08/165,699, Ser. No. 08/209,741. See also EP 0 546 073 B1, WO 92/03918, WO 92/22645, WO 92/22647, WO 92/22670, WO 93/12227, WO 94/00569, WO 94/25585, WO 96/14436, WO 97/13852, and WO 98/24884 and U.S. Pat. No. 5,981,175. See further Taylor *et al.* (1992), Chen *et al.* (1993), Tuailon *et al.* (1993), Choi *et al.* (1993), Lonberg *et al.* (1994), Taylor *et al.* (1994), and Tuailon *et al.* (1995), Fishwild *et al.* (1996). 10

[139] Kirin has also demonstrated the generation of human antibodies from mice in which, through microcell fusion, large pieces of chromosomes, or entire chromosomes, have been introduced. See European Patent Application Nos. 773 288 and 843 961. Xenerex Biosciences is developing a technology for the potential generation of human antibodies. In this technology, SCID mice are reconstituted with human lymphatic cells, e.g., B and/or T cells. Mice are then immunized with an 15 antigen and can generate an immune response against the antigen. See U.S. Pat. Nos. 5,476,996; 5,698,767; and 5,958,765.

[140] Human anti-mouse antibody (HAMA) responses have led the industry to prepare chimeric or otherwise humanized antibodies. It is however expected that certain human anti-chimeric antibody 20 (HACA) responses will be observed, particularly in chronic or multi-dose utilizations of the antibody. Thus, it would be desirable to provide antigen-binding molecules comprising a human binding domain against CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM and a human binding domain against CD3 ϵ in order to vitiate concerns and/or effects of HAMA or HACA response.

[141] The terms “(specifically) binds to”, “(specifically) recognizes”, “is (specifically) directed to”, and “(specifically) reacts with” mean in accordance with this invention that a binding domain 25 interacts or specifically interacts with a given epitope or a given target side on the target molecules (antigens), here: CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM, and CD3 ϵ as effector, respectively.

[142] The term “epitope” refers to a side on an antigen to which a binding domain, such as an antibody or immunoglobulin, or a derivative, fragment or variant of an antibody or an immunoglobulin, specifically binds. An “epitope” is antigenic and thus the term epitope is 30 sometimes also referred to herein as “antigenic structure” or “antigenic determinant”. Thus, the binding domain is an “antigen interaction side”. Said binding/interaction is also understood to define a “specific recognition”. 35

[143] "Epitopes" can be formed both by contiguous amino acids or non-contiguous amino acids juxtaposed by tertiary folding of a protein. A "linear epitope" is an epitope where an amino acid primary sequence comprises the recognized epitope. A linear epitope typically includes at least 3 or at least 4, and more usually, at least 5 or at least 6 or at least 7, for example, about 8 to about 10 amino acids in a unique sequence.

[144] A "conformational epitope", in contrast to a linear epitope, is an epitope wherein the primary sequence of the amino acids comprising the epitope is not the sole defining component of the epitope recognized (*e.g.*, an epitope wherein the primary sequence of amino acids is not necessarily recognized by the binding domain). Typically, a conformational epitope comprises an increased number of amino acids relative to a linear epitope. With regard to recognition of conformational epitopes, the binding domain recognizes a three-dimensional structure of the antigen, preferably a peptide or protein or fragment thereof (in the context of the present invention, the antigenic structure for one of the binding domains is comprised within the target cell surface antigen protein). For example, when a protein molecule folds to form a three-dimensional structure, certain amino acids and/or the polypeptide backbone forming the conformational epitope become juxtaposed enabling the antibody to recognize the epitope. Methods of determining the conformation of epitopes include, but are not limited to, x-ray crystallography, two-dimensional nuclear magnetic resonance (2D-NMR) spectroscopy and site-directed spin labelling and electron paramagnetic resonance (EPR) spectroscopy.

[145] A method for epitope mapping is described in the following: When a region (a contiguous amino acid stretch) in the human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM protein is exchanged or replaced with its corresponding region of a non-human and non-primate CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM (*e.g.*, mouse CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM, but others like chicken, rat, hamster, rabbit etc. may also be conceivable), a decrease in the binding of the binding domain is expected to occur, unless the binding domain is cross-reactive for the non-human, non-primate CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM used. Said decrease is preferably at least 10%, 20%, 30%, 40%, or 50%; more preferably at least 60%, 70%, or 80%, and most preferably 90%, 95% or even 100% in comparison to the binding to the respective region in the human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM protein, whereby binding to the respective region in the human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM protein is set to be 100%. It is envisaged that the aforementioned human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM / non-human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM chimeras are expressed in CHO cells. It is also envisaged that the human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM / non-human CS1, BCMA, CD20, CD22, FLT3,

CD123, CLL1, CDH3, MSLN, or EpCAM chimeras are fused with a transmembrane domain and/or cytoplasmic domain of a different membrane-bound protein such as EpCAM.

[146] In an alternative or additional method for epitope mapping, several truncated versions of the human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM extracellular domain can be generated in order to determine a specific region that is recognized by a binding domain. In these truncated versions, the different extracellular CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM domains / sub-domains or regions are stepwise deleted, starting from the N-terminus. It is envisaged that the truncated CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM versions may be expressed in CHO cells. It is also envisaged that the truncated CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM versions may be fused with a transmembrane domain and/or cytoplasmic domain of a different membrane-bound protein such as EpCAM. It is also envisaged that the truncated CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM versions may encompass a signal peptide domain at their N-terminus, for example a signal peptide derived from mouse IgG heavy chain signal peptide. It is furthermore envisaged that the truncated CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM versions may encompass a v5 domain at their N-terminus (following the signal peptide) which allows verifying their correct expression on the cell surface. A decrease or a loss of binding is expected to occur with those truncated CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM versions which do not encompass any more the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM region that is recognized by the binding domain. The decrease of binding is preferably at least 10%, 20%, 30%, 40%, 50%; more preferably at least 60%, 70%, 80%, and most preferably 90%, 95% or even 100%, whereby binding to the entire human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM protein (or its extracellular region or domain) is set to be 100.

[147] A further method to determine the contribution of a specific residue of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM to the recognition by an antigen-binding molecule or binding domain is alanine scanning (see e.g. Morrison KL & Weiss GA. *Cur Opin Chem Biol.* 2001 Jun;5(3):302-7), where each residue to be analyzed is replaced by alanine, e.g. via site-directed mutagenesis. Alanine is used because of its non-bulky, chemically inert, methyl functional group that nevertheless mimics the secondary structure references that many of the other amino acids possess. Sometimes bulky amino acids such as valine or leucine can be used in cases where conservation of the size of mutated residues is desired. Alanine scanning is a mature technology which has been used for a long period of time.

[148] The interaction between the binding domain and the epitope or the region comprising the epitope implies that a binding domain exhibits appreciable affinity for the epitope / the region comprising the epitope on a particular protein or antigen (here:, CD20, CD22, FLT3, CD123, CLL1,

CDH3, MSLN, or EpCAM and CD3, respectively) and, generally, does not exhibit significant reactivity with proteins or antigens other than the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM or CD3. “Appreciable affinity” includes binding with an affinity of about 10^{-6} M (KD) or stronger. Preferably, binding is considered specific when the binding affinity is about 10^{-12} to 10^{-8} M, 10^{-12} to 10^{-9} M, 10^{-12} to 10^{-10} M, 10^{-11} to 10^{-8} M, preferably of about 10^{-11} to 10^{-9} M. Whether a binding domain specifically reacts with or binds to a target can be tested readily by, *inter alia*, comparing the reaction of said binding domain with a target protein or antigen with the reaction of said binding domain with proteins or antigens other than the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM or CD3. Preferably, a binding domain of the invention does not essentially or substantially bind to proteins or antigens other than CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM or CD3 (*i.e.*, the first binding domain is not capable of binding to proteins other than CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM and the second binding domain is not capable of binding to proteins other than CD3). It is an envisaged characteristic of the antigen-binding molecules according to the present invention to have superior affinity characteristics in comparison to other HLE formats. Such a superior affinity, in consequence, suggests a prolonged half-life in vivo. The longer half-life of the antigen-binding molecules according to the present invention may reduce the duration and frequency of administration which typically contributes to improved patient compliance. This is of particular importance as the antigen-binding molecules of the present invention are particularly beneficial for highly weakened or even multimorbid cancer patients.

[149] The term “does not essentially / substantially bind” or “is not capable of binding” means that a binding domain of the present invention does not bind a protein or antigen other than the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM or CD3 as effector, *i.e.*, does not show reactivity of more than 30%, preferably not more than 20%, more preferably not more than 10%, particularly preferably not more than 9%, 8%, 7%, 6% or 5% with proteins or antigens other than CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM or CD3 as effector, whereby binding to the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CDH3, MSLN, or EpCAM or CD3 as effector, respectively, is set to be 100%.

[150] Specific binding is believed to be effected by specific motifs in the amino acid sequence of the binding domain and the antigen. Thus, binding is achieved as a result of their primary, secondary and/or tertiary structure as well as the result of secondary modifications of said structures. The specific interaction of the antigen-interaction-side with its specific antigen may result in a simple binding of said side to the antigen. Moreover, the specific interaction of the antigen-interaction-side with its specific antigen may alternatively or additionally result in the initiation of a signal, *e.g.* due to the induction of a change of the conformation of the antigen, an oligomerization of the antigen, etc.

[151] The term “variable” refers to the portions of the antibody or immunoglobulin domains that exhibit variability in their sequence and that are involved in determining the specificity and binding affinity of a particular antibody (i.e., the “variable domain(s)”). The pairing of a variable heavy chain (VH) and a variable light chain (VL) together forms a single antigen-binding site.

5 [152] Variability is not evenly distributed throughout the variable domains of antibodies; it is concentrated in sub-domains of each of the heavy and light chain variable regions. These sub-domains are called “hypervariable regions” or “complementarity determining regions” (CDRs). The more conserved (i.e., non-hypervariable) portions of the variable domains are called the “framework” regions (FRM or FR) and provide a scaffold for the six CDRs in three dimensional
10 space to form an antigen-binding surface. The variable domains of naturally occurring heavy and light chains each comprise four FRM regions (FR1, FR2, FR3, and FR4), largely adopting a β -sheet configuration, connected by three hypervariable regions, which form loops connecting, and in some cases forming part of, the β -sheet structure. The hypervariable regions in each chain are held together in close proximity by the FRM and, with the hypervariable regions from the other chain, contribute
15 to the formation of the antigen-binding side (see Kabat *et al.*, *loc. cit.*).

[153] The terms “CDR”, and its plural “CDRs”, refer to the complementarity determining region of which three make up the binding character of a light chain variable region (CDR-L1, CDR-L2 and CDR-L3) and three make up the binding character of a heavy chain variable region (CDR-H1, CDR-H2 and CDR-H3). CDRs contain most of the residues responsible for specific interactions of the
20 antibody with the antigen and hence contribute to the functional activity of an antibody molecule: they are the main determinants of antigen specificity.

[154] The exact definitional CDR boundaries and lengths are subject to different classification and numbering systems. CDRs may therefore be referred to by Kabat, Chothia, contact or any other boundary definitions, including the numbering system described herein. Despite differing
25 boundaries, each of these systems has some degree of overlap in what constitutes the so called “hypervariable regions” within the variable sequences. CDR definitions according to these systems may therefore differ in length and boundary areas with respect to the adjacent framework region. See for example Kabat (an approach based on cross-species sequence variability), Chothia (an approach based on crystallographic studies of antigen-antibody complexes), and/or MacCallum (Kabat *et al.*,
30 *loc. cit.*; Chothia *et al.*, *J. Mol. Biol.*, 1987, 196: 901-917; and MacCallum *et al.*, *J. Mol. Biol.*, 1996, 262: 732). Still another standard for characterizing the antigen binding side is the AbM definition used by Oxford Molecular's AbM antibody modeling software. See, *e.g.*, Protein Sequence and Structure Analysis of Antibody Variable Domains. In: Antibody Engineering Lab Manual (Ed.: Duebel, S. and Kontermann, R., Springer-Verlag, Heidelberg). To the extent that two residue
35 identification techniques define regions of overlapping, but not identical regions, they can be

combined to define a hybrid CDR. However, the numbering in accordance with the so-called Kabat system is preferred.

[155] Typically, CDRs form a loop structure that can be classified as a canonical structure. The term “canonical structure” refers to the main chain conformation that is adopted by the antigen binding (CDR) loops. From comparative structural studies, it has been found that five of the six antigen binding loops have only a limited repertoire of available conformations. Each canonical structure can be characterized by the torsion angles of the polypeptide backbone. Correspondent loops between antibodies may, therefore, have very similar three dimensional structures, despite high amino acid sequence variability in most parts of the loops (Chothia and Lesk, *J. Mol. Biol.*, 1987, 196: 901; Chothia *et al.*, *Nature*, 1989, 342: 877; Martin and Thornton, *J. Mol. Biol.*, 1996, 263: 800). Furthermore, there is a relationship between the adopted loop structure and the amino acid sequences surrounding it. The conformation of a particular canonical class is determined by the length of the loop and the amino acid residues residing at key positions within the loop, as well as within the conserved framework (*i.e.*, outside of the loop). Assignment to a particular canonical class can therefore be made based on the presence of these key amino acid residues.

[156] The term “canonical structure” may also include considerations as to the linear sequence of the antibody, for example, as catalogued by Kabat (Kabat *et al.*, *loc. cit.*). The Kabat numbering scheme (system) is a widely adopted standard for numbering the amino acid residues of an antibody variable domain in a consistent manner and is the preferred scheme applied in the present invention as also mentioned elsewhere herein. Additional structural considerations can also be used to determine the canonical structure of an antibody. For example, those differences not fully reflected by Kabat numbering can be described by the numbering system of Chothia *et al.* and/or revealed by other techniques, for example, crystallography and two- or three-dimensional computational modeling. Accordingly, a given antibody sequence may be placed into a canonical class which allows for, among other things, identifying appropriate chassis sequences (*e.g.*, based on a desire to include a variety of canonical structures in a library). Kabat numbering of antibody amino acid sequences and structural considerations as described by Chothia *et al.*, *loc. cit.* and their implications for construing canonical aspects of antibody structure, are described in the literature. The subunit structures and three-dimensional configurations of different classes of immunoglobulins are well known in the art. For a review of the antibody structure, see *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, eds. Harlow *et al.*, 1988.

[157] The CDR3 of the light chain and, particularly, the CDR3 of the heavy chain may constitute the most important determinants in antigen binding within the light and heavy chain variable regions. In some antigen-binding molecules, the heavy chain CDR3 appears to constitute the major area of contact between the antigen and the antibody. *In vitro* selection schemes in which CDR3 alone is varied can be used to vary the binding properties of an antibody or determine which residues

contribute to the binding of an antigen. Hence, CDR3 is typically the greatest source of molecular diversity within the antibody-binding side. H3, for example, can be as short as two amino acid residues or greater than 26 amino acids.

5 [158] In a classical full-length antibody or immunoglobulin, each light (L) chain is linked to a heavy (H) chain by one covalent disulfide bond, while the two H chains are linked to each other by one or more disulfide bonds depending on the H chain isotype. The CH domain most proximal to VH is usually designated as CH1. The constant (“C”) domains are not directly involved in antigen binding, but exhibit various effector functions, such as antibody-dependent, cell-mediated cytotoxicity and complement activation. The Fc region of an antibody is comprised within the heavy chain constant
10 domains and is for example able to interact with cell surface located Fc receptors.

[159] The sequence of antibody genes after assembly and somatic mutation is highly varied, and these varied genes are estimated to encode 10^{10} different antibody molecules (Immunoglobulin Genes, 2nd ed., eds. Jonio et al., Academic Press, San Diego, CA, 1995). Accordingly, the immune system provides a repertoire of immunoglobulins. The term “repertoire” refers to at least one
15 nucleotide sequence derived wholly or partially from at least one sequence encoding at least one immunoglobulin. The sequence(s) may be generated by rearrangement *in vivo* of the V, D, and J segments of heavy chains, and the V and J segments of light chains. Alternatively, the sequence(s) can be generated from a cell in response to which rearrangement occurs, e.g., *in vitro* stimulation. Alternatively, part or all of the sequence(s) may be obtained by DNA splicing, nucleotide synthesis,
20 mutagenesis, and other methods, see, e.g., U.S. Patent 5,565,332. A repertoire may include only one sequence or may include a plurality of sequences, including ones in a genetically diverse collection.

[160] The term "Fc portion" or "Fc monomer" means in connection with this invention a polypeptide comprising at least one domain having the function of a CH2 domain and at least one domain having the function of a CH3 domain of an immunoglobulin molecule. As apparent from the term “Fc
25 monomer”, the polypeptide comprising those CH domains is a “polypeptide monomer”. An Fc monomer can be a polypeptide comprising at least a fragment of the constant region of an immunoglobulin excluding the first constant region immunoglobulin domain of the heavy chain (CH1), but maintaining at least a functional part of one CH2 domain and a functional part of one CH3 domain, wherein the CH2 domain is amino terminal to the CH3 domain. In a preferred aspect
30 of this definition, an Fc monomer can be a polypeptide constant region comprising a portion of the Ig-Fc hinge region, a CH2 region and a CH3 region, wherein the hinge region is amino terminal to the CH2 domain. It is envisaged that the hinge region of the present invention promotes dimerization. Such Fc polypeptide molecules can be obtained by papain digestion of an immunoglobulin region (of course resulting in a dimer of two Fc polypeptide), for example and not
35 limitation. In another aspect of this definition, an Fc monomer can be a polypeptide region comprising a portion of a CH2 region and a CH3 region. Such Fc polypeptide molecules can be

obtained by pepsin digestion of an immunoglobulin molecule, for example and not limitation. In one embodiment, the polypeptide sequence of an Fc monomer is substantially similar to an Fc polypeptide sequence of: an IgG₁ Fc region, an IgG₂ Fc region, an IgG₃ Fc region, an IgG₄ Fc region, an IgM Fc region, an IgA Fc region, an IgD Fc region and an IgE Fc region. (See, e.g., Padlan, Molecular Immunology, 31(3), 169-217 (1993)). Because there is some variation between immunoglobulins, and solely for clarity, Fc monomer refers to the last two heavy chain constant region immunoglobulin domains of IgA, IgD, and IgG, and the last three heavy chain constant region immunoglobulin domains of IgE and IgM. As mentioned, the Fc monomer can also include the flexible hinge N-terminal to these domains. For IgA and IgM, the Fc monomer may include the J chain. For IgG, the Fc portion comprises immunoglobulin domains CH2 and CH3 and the hinge between the first two domains and CH2. Although the boundaries of the Fc portion may vary an example for a human IgG heavy chain Fc portion comprising a functional hinge, CH2 and CH3 domain can be defined e.g. to comprise residues D231 (of the hinge domain– corresponding to D234 in Table 1 below) to P476, respectively L476 (for IgG₄) of the carboxyl-terminus of the CH3 domain, wherein the numbering is according to Kabat. The two Fc portion or Fc monomer, which are fused to each other via a peptide linker are a preferred example of the spacer between the two bispecific entities of the antigen-binding molecule of the invention, which may also be defined as scFc domain.

[161] In one embodiment of the invention it is envisaged that a scFc domain as disclosed herein, respectively the Fc monomers fused to each other are comprised only in the spacer of the antigen-binding molecule.

[162] In line with the present invention an IgG hinge region can be identified by analogy using the Kabat numbering as set forth in Table 1. In line with the above, it is envisaged that for a hinge domain/region of the present invention the minimal requirement comprises the amino acid residues corresponding to the IgG1 sequence stretch of D231 D234 to P243 according to the Kabat numbering. It is likewise envisaged that a hinge domain/region of the present invention comprises or consists of the IgG1 hinge sequence DKTHTCPPCP (SEQ ID NO: 330) (corresponding to the stretch D234 to P243 as shown in Table 1 below – variations of said sequence are also envisaged provided that the hinge region still promotes dimerization). In a preferred embodiment of the invention the glycosylation site at Kabat position 314 of the CH2 domains in the spacer of the antigen-binding molecule is removed by a N314X substitution, wherein X is any amino acid excluding Q. Said substitution is preferably a N314G substitution. In a more preferred embodiment, said CH2 domain additionally comprises the following substitutions (position according to Kabat) V321C and R309C (these substitutions introduce the intra domain cysteine disulfide bridge at Kabat positions 309 and 321).

[163] It is also envisaged that the spacer of the antigen-binding molecule of the invention is a scFc domain which may comprise or consist of an amino to carboxyl order: DKTHTCPPCP (SEQ ID NO: 330) (i.e. hinge) -CH₂-CH₃-linker- DKTHTCPPCP (SEQ ID NO: 330) (i.e. hinge) -CH₂-CH₃. The peptide linker of the aforementioned antigen-binding molecule is in a preferred embodiment characterized by the amino acid sequence Gly-Gly-Gly-Gly-Ser, i.e. Gly₄Ser (SEQ ID NO: 7), or polymers thereof, i.e. (Gly₄Ser)_x, where x is an integer of 5 or greater (e.g. 5, 6, 7, 8 etc. or greater), 6 being preferred ((Gly₄Ser)₆). According to the present invention, the Ser may advantageously be replaced by Gln as disclosed herein. Said construct may further comprise the aforementioned substitutions: N314X, preferably N314G, and/or the further substitutions V321C and R309C. In a preferred embodiment of the antigen-binding molecules of the invention as defined herein before, it is envisaged that the second domain binds to an extracellular epitope of the human and/or the *Macaca* CD3 ϵ chain. Table 1: Kabat numbering of the amino acid residues of the hinge region

IMGT numbering for the hinge	IgG ₁ amino acid translation	Kabat numbering
1	(E)	226
2	P	227
3	K	228
4	S	232
5	C	233
6	D	234
7	K	235
8	T	236
9	H	237
10	T	238
11	C	239
12	P	240
13	P	241
14	C	242
15	P	243

[164] In further embodiments of the present invention, the hinge domain/region comprises or consists of the IgG₂ subtype hinge sequence ERKCCVECPCPCP (SEQ ID NO: 331), the IgG₃ subtype hinge sequence ELKTPLDTHTCPRCP (SEQ ID NO: 332) or ELKTPLGDTTHTCPRCP (SEQ ID NO: 333), and/or the IgG₄ subtype hinge sequence ESKYGPPCPCPCP (SEQ ID NO: 444). The IgG₁ subtype hinge sequence may be the following one EPKSCDKTHTCPCPCP (as shown in Table 1 and SEQ ID NO: 445). These core hinge regions are thus also envisaged in the context of the present invention.

[165] The location and sequence of the IgG CH2 and IgG CD3 domain can be identified by analogy using the Kabat numbering as set forth in Table 2:

Table 2: Kabat numbering of the amino acid residues of the IgG CH2 and CH3 region

IgG subtype	CH2 aa translation	CH2 Kabat numbering	CH3 aa translation	CH3 Kabat numbering
IgG₁	APE KAK	244... ...360	GQP..... PGK	361... ...478
IgG₂	APP GTK	244... ...360	GQP..... PGK	361... ...478
IgG₃	APE GTK	244... ...360	GQP..... PGK	361... ...478
IgG₄	APE KAK	244... ...360	GQP..... LGK	361... ...478

[166] In one embodiment of the invention the emphasized bold amino acid residues in the CH3 domain of the first or both Fc monomers are deleted.

[167] The peptide linker, by whom the polypeptide monomers ("Fc portion" or "Fc monomer") of the spacer are fused to each other, preferably comprises at least 25 amino acid residues (25, 26, 27, 28, 29, 30 etc.). More preferably, this peptide linker comprises at least 30 amino acid residues (30, 31, 32, 33, 34, 35 etc.). It is also preferred that the linker comprises up to 40 amino acid residues, more preferably up to 35 amino acid residues, most preferably exactly 30 amino acid residues. A preferred embodiment of such peptide linker is characterized by the amino acid sequence Gly-Gly-Gly-Gly-Ser, i.e. Gly₄Ser (SEQ ID NO: 7), or polymers thereof, i.e. (Gly₄Ser)_x, where x is an integer of 5 or greater (e.g. 6, 7 or 8). Preferably the integer is 6 or 7, more preferably the integer is 6.

[168] In the event that a linker is used to fuse the first domain to the second domain, and/or the third to the fourth domain, and/or the second and the third domain to the spacer, this linker is preferably of a length and sequence sufficient to ensure that each of the first and second domains can, independently from one another, retain their differential binding specificities. For peptide linkers which connect the at least two binding domains (or two variable domains) in the antigen-binding molecule of the invention, those peptide linkers are preferred which comprise only a few number of amino acid residues, e.g. 12 amino acid residues or less. Thus, peptide linkers of 12, 11, 10, 9, 8, 7, 6 or 5 amino acid residues are preferred. An envisaged peptide linker with less than 5 amino acids comprises 4, 3, 2 or one amino acid(s), wherein Gly-rich linkers are preferred. A preferred embodiment of the peptide linker for a fusion the first and the second domain is depicted in SEQ ID NO:1. A preferred linker embodiment of the peptide linker for fusing the second and the third domain to the spacer is a (Gly)₄-linker, also called G₄-linker.

[169] A particularly preferred "single" amino acid in the context of one of the above described "peptide linker" is Gly. Accordingly, said peptide linker may consist of the single amino acid Gly. In a preferred embodiment of the invention a peptide linker is characterized by the amino acid sequence

Gly-Gly-Gly-Gly-Ser, i.e. Gly₄Ser (SEQ ID NO: 1), or polymers thereof, i.e. (Gly₄Ser)_x, where x is an integer of 1 or greater (e.g. 2 or 3). Preferred linkers are depicted in SEQ ID NOs: 1 to 12. The characteristics of said peptide linker, which comprise the absence of the promotion of secondary structures, are known in the art and are described e.g. in Dall'Acqua et al. (Biochem. (1998) 37, 9266-9273), Cheadle et al. (Mol Immunol (1992) 29, 21-30) and Raag and Whitlow (FASEB (1995) 9(1), 73-80). Peptide linkers which furthermore do not promote any secondary structures are preferred. The linkage of said domains to each other can be provided, e.g., by genetic engineering, as described in the examples. Methods for preparing fused and operatively linked bispecific single chain constructs and expressing them in mammalian cells or bacteria are well-known in the art (e.g. WO 99/54440 or Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York, 2001).

[170] In a preferred embodiment of the antigen-binding molecule of the present invention the first and second domain form an antigen-binding molecule in a format selected from the group consisting of (scFv)₂, scFv-single domain mAb, diabody and oligomers of any of these formats.

[171] According to a particularly preferred embodiment, and as documented in the appended examples, the first and the second domain of the antigen-binding molecule of the invention is a "bispecific single chain antigen-binding molecule", more preferably a bispecific "single chain Fv" (scFv). Although the two domains of the Fv fragment, VL and VH, are coded for by separate genes, they can be joined, using recombinant methods, by a synthetic linker – as described hereinbefore – that enables them to be made as a single protein chain in which the VL and VH regions pair to form a monovalent molecule; see e.g., Huston et al. (1988) Proc. Natl. Acad. Sci USA 85:5879-5883). These antibody fragments are obtained using conventional techniques known to those with skill in the art, and the fragments are evaluated for function in the same manner as are whole or full-length antibodies. A single-chain variable fragment (scFv) is hence a fusion protein of the variable region of the heavy chain (VH) and of the light chain (VL) of immunoglobulins, usually connected with a short linker peptide of about ten to about 25 amino acids, preferably about 15 to 20 amino acids. The linker is usually rich in glycine for flexibility, as well as serine or threonine for solubility, and can either connect the N-terminus of the VH with the C-terminus of the VL, or *vice versa*. This protein retains the specificity of the original immunoglobulin, despite removal of the constant regions and introduction of the linker.

[172] Bispecific single chain antigen-binding molecules are known in the art and are described in WO 99/54440, Mack, J. Immunol. (1997), 158, 3965-3970, Mack, PNAS, (1995), 92, 7021-7025, Kufer, Cancer Immunol. Immunother., (1997), 45, 193-197, Löffler, Blood, (2000), 95, 6, 2098-2103, Brühl, Immunol., (2001), 166, 2420-2426, Kipriyanov, J. Mol. Biol., (1999), 293, 41-56. Techniques described for the production of single chain antibodies (see, *inter alia*, US Patent

4,946,778, Kontermann and Dübel (2010), *loc. cit.* and Little (2009), *loc. cit.*) can be adapted to produce single chain antigen-binding molecules specifically recognizing (an) elected target(s).

5 [173] Bivalent (also called divalent) or bispecific single-chain variable fragments (bi-scFvs or di-scFvs having the format (scFv)₂ can be engineered by linking two scFv molecules (*e.g.* with linkers as described hereinbefore). If these two scFv molecules have the same binding specificity, the resulting (scFv)₂ molecule will preferably be called bivalent (*i.e.* it has two valences for the same target epitope). If the two scFv molecules have different binding specificities, the resulting (scFv)₂ molecule will preferably be called bispecific. The linking can be done by producing a single peptide chain with two VH regions and two VL regions, yielding tandem scFvs (see *e.g.* Kufer P. *et al.*, 10 (2004) Trends in Biotechnology 22(5):238-244). Another possibility is the creation of scFv molecules with linker peptides that are too short for the two variable regions to fold together (*e.g.* about five amino acids), forcing the scFvs to dimerize. This type is known as diabodies (see *e.g.* Hollinger, Philipp *et al.*, (July 1993) Proceedings of the National Academy of Sciences of the United States of America 90 (14): 6444-8).

15 [174] In line with this invention either the first, the second, the third and/or the fourth may comprise a single domain antibody, respectively the variable domain or at least the CDRs of a single domain antibody. Single domain antibodies comprise merely one (monomeric) antibody variable domain which is able to bind selectively to a specific antigen, independently of other V regions or domains. The first single domain antibodies were engineered from heavy chain antibodies found in camelids, 20 and these are called V_{HH} fragments. Cartilaginous fishes also have heavy chain antibodies (IgNAR) from which single domain antibodies called V_{NAR} fragments can be obtained. An alternative approach is to split the dimeric variable domains from common immunoglobulins *e.g.* from humans or rodents into monomers, hence obtaining VH or VL as a single domain Ab. Although most research into single domain antibodies is currently based on heavy chain variable domains, 25 nanobodies derived from light chains have also been shown to bind specifically to target epitopes. Examples of single domain antibodies are called sdAb, nanobodies or single variable domain antibodies.

30 [175] A (single domain mAb)₂ is hence a monoclonal antigen-binding molecule composed of (at least) two single domain monoclonal antibodies, which are individually selected from the group comprising V_H, V_L, V_{HH} and V_{NAR}. The linker is preferably in the form of a peptide linker. Similarly, an “scFv-single domain mAb” is a monoclonal antigen-binding molecule composed of at least one single domain antibody as described above and one scFv molecule as described above. Again, the linker is preferably in the form of a peptide linker.

35 [176] Whether or not an antigen-binding molecule competes for binding with another given antigen-binding molecule can be measured in a competition assay such as a competitive ELISA or a cell-

based competition assay. Avidin-coupled microparticles (beads) can also be used. Similar to an avidin-coated ELISA plate, when reacted with a biotinylated protein, each of these beads can be used as a substrate on which an assay can be performed. Antigen is coated onto a bead and then precoated with the first antibody. The second antibody is added and any additional binding is determined.
5 Possible means for the read-out includes flow cytometry.

[177] T cells or T lymphocytes are a type of lymphocyte (itself a type of white blood cell) that play a central role in cell-mediated immunity. There are several subsets of T cells, each with a distinct function. T cells can be distinguished from other lymphocytes, such as B cells and NK cells, by the presence of a T cell receptor (TCR) on the cell surface. The TCR is responsible for recognizing
10 antigens bound to major histocompatibility complex (MHC) molecules and is composed of two different protein chains. In 95% of the T cells, the TCR consists of an alpha (α) and beta (β) chain. When the TCR engages with antigenic peptide and MHC (peptide / MHC complex), the T lymphocyte is activated through a series of biochemical events mediated by associated enzymes, co-receptors, specialized adaptor molecules, and activated or released transcription factors.

[178] The CD3 receptor complex is a protein complex and is composed of four chains. In mammals, the complex contains a CD3 γ (gamma) chain, a CD3 δ (delta) chain, and two CD3 ϵ (epsilon) chains. These chains associate with the T cell receptor (TCR) and the so-called ζ (zeta) chain to form the T cell receptor CD3 complex and to generate an activation signal in T lymphocytes. The CD3 γ (gamma), CD3 δ (delta), and CD3 ϵ (epsilon) chains are highly related cell-surface proteins of the
20 immunoglobulin superfamily containing a single extracellular immunoglobulin domain. The intracellular tails of the CD3 molecules contain a single conserved motif known as an immunoreceptor tyrosine-based activation motif or ITAM for short, which is essential for the signaling capacity of the TCR. The CD3 epsilon molecule is a polypeptide which in humans is encoded by the *CD3E* gene which resides on chromosome 11. The most preferred epitope of
25 CD3 epsilon is comprised within amino acid residues 1-27 of the human CD3 epsilon extracellular domain. It is envisaged that antigen-binding molecules according to the present invention typically and advantageously show less unspecific T cell activation, which is not desired in specific immunotherapy. This translates to a reduced risk of side effects.

[179] The redirected lysis of target cells via the recruitment of T cells by a multichain multitargeting
30 least bispecific antigen-binding molecule involves cytolytic synapse formation and delivery of perforin and granzymes. The engaged T cells are capable of serial target cell lysis, and are not affected by immune escape mechanisms interfering with peptide antigen processing and presentation, or clonal T cell differentiation; see, for example, WO 2007/042261.

[180] Cytotoxicity mediated by antigen-binding molecules of the invention can be measured in
35 various ways. Effector cells can be e.g. stimulated enriched (human) CD8 positive T cells or

unstimulated (human) peripheral blood mononuclear cells (PBMC). If the target cells are of macaque origin or express or are transfected with macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM which is bound by the first domain, the effector cells should also be of macaque origin such as a macaque T cell line, e.g. 4119LnPx. The target cells should express (at least the extracellular domain of) CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM, e.g. human or macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM. Target cells can be a cell line (such as CHO) which is stably or transiently transfected with CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM, e.g. human or macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM. Usually EC_{50} values are expected to be lower with target cell lines expressing higher levels of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM on the cell surface. The effector to target cell (E:T) ratio is usually about 10:1, but can also vary. Cytotoxic activity of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM bispecific antigen-binding molecules can be measured in a ^{51}Cr -release assay (incubation time of about 18 hours) or in a FACS-based cytotoxicity assay (incubation time of about 48 hours). Modifications of the assay incubation time (cytotoxic reaction) are also possible. Other methods of measuring cytotoxicity are well-known to the skilled person and comprise MTT or MTS assays, ATP-based assays including bioluminescent assays, the sulforhodamine B (SRB) assay, WST assay, clonogenic assay and the ECIS technology.

[181] The cytotoxic activity mediated by CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecules of the present invention is preferably measured in a cell-based cytotoxicity assay. It may also be measured in a ^{51}Cr -release assay. It is represented by the EC_{50} value, which corresponds to the half maximal effective concentration (concentration of the antigen-binding molecule which induces a cytotoxic response halfway between the baseline and maximum). Preferably, the EC_{50} value of the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecules is ≤ 5000 pM or ≤ 4000 pM, more preferably ≤ 3000 pM or ≤ 2000 pM, even more preferably ≤ 1000 pM or ≤ 500 pM, even more preferably ≤ 400 pM or ≤ 300 pM, even more preferably ≤ 200 pM, even more preferably ≤ 100 pM, even more preferably ≤ 50 pM, even more preferably ≤ 20 pM or ≤ 10 pM, and most preferably ≤ 5 pM.

[182] The above given EC_{50} values can be measured in different assays. The skilled person is aware that an EC_{50} value can be expected to be lower when stimulated / enriched CD8^+ T cells are used as effector cells, compared with unstimulated PBMC. It can furthermore be expected that the EC_{50} values are lower when the target cells express a high number of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM compared with a low target expression rat. For example, when stimulated / enriched human CD8^+ T cells are used as effector cells (and either CS1, BCMA,

CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM transfected cells such as CHO cells or CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM positive human cell lines are used as target cells), the EC₅₀ value of the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecule is preferably
5 ≤1000 pM, more preferably ≤500 pM, even more preferably ≤250 pM, even more preferably ≤100 pM, even more preferably ≤50 pM, even more preferably ≤10 pM, and most preferably ≤5 pM. When human PBMCs are used as effector cells, the EC₅₀ value of the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecule is preferably ≤5000 pM or ≤4000 pM (in particular when the target cells are CS1, BCMA, CD20,
10 CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM positive human cell lines), more preferably ≤2000 pM (in particular when the target cells are CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM transfected cells such as CHO cells), more preferably ≤1000 pM or ≤500 pM, even more preferably ≤200 pM, even more preferably ≤150 pM, even more preferably ≤100 pM, and most preferably ≤50 pM, or lower. When a macaque T cell line such as LnPx4119 is
15 used as effector cells, and a macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM transfected cell line such as CHO cells is used as target cell line, the EC₅₀ value of the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecule is preferably ≤2000 pM or ≤1500 pM, more preferably ≤1000 pM or ≤500 pM, even more preferably ≤300 pM or ≤250 pM, even more preferably ≤100 pM, and most
20 preferably ≤50 pM.

[183] Preferably, the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecules of the present invention do not induce / mediate lysis or do not essentially induce / mediate lysis of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM negative cells such as CHO cells. The term “do not induce lysis”,
25 “do not essentially induce lysis”, “do not mediate lysis” or “do not essentially mediate lysis” means that an antigen-binding molecule of the present invention does not induce or mediate lysis of more than 30%, preferably not more than 20%, more preferably not more than 10%, particularly preferably not more than 9%, 8%, 7%, 6% or 5% of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM negative cells, whereby lysis of a CS1, BCMA, CD20, CD22, FLT3, CD123,
30 CLL1, CHD3, MSLN, or EpCAM positive human cell line is set to be 100%. This usually applies for concentrations of the antigen-binding molecule of up to 500 nM. The skilled person knows how to measure cell lysis without further ado. Moreover, the present specification teaches specific instructions how to measure cell lysis.

[184] The difference in cytotoxic activity between the monomeric and the dimeric isoform of individual CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3
35 bispecific antigen-binding molecules is referred to as “potency gap”. This potency gap can e.g. be

calculated as ratio between EC_{50} values of the molecule's monomeric and dimeric form. Potency gaps of the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM \times CD3 bispecific antigen-binding molecules of the present invention are preferably ≤ 5 , more preferably ≤ 4 , even more preferably ≤ 3 , even more preferably ≤ 2 and most preferably ≤ 1 .

5 [185] The first, second, third and/or the fourth binding domain of the antigen-binding molecule of the invention is/are preferably cross-species specific for members of the mammalian order of primates. Cross-species specific CD3 binding domains are, for example, those described herein and in WO 2008/119567. According to one embodiment, the first and third binding domain, in addition to binding to human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM
10 and human CD3, respectively, will also bind to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM / CD3 of primates including (but not limited to) new world primates (such as *Callithrix jacchus*, *Saguinus Oedipus* or *Saimiri sciureus*), old world primates (such as baboons and macaques), gibbons, and non-human *homininae*.

[186] In one embodiment of the antigen-binding molecule of the invention the first domain binds to
15 human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM and further binds to macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM, such as CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM of *Macaca fascicularis*, and more preferably, to macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM expressed on the surface of cells, e.g. such as CHO or 293 cells. The
20 affinity of the first domain for CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM, preferably for human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM, is preferably ≤ 100 nM or ≤ 50 nM, more preferably ≤ 25 nM or ≤ 20 nM, more preferably ≤ 15 nM or ≤ 10 nM, even more preferably ≤ 5 nM, even more preferably ≤ 2.5 nM or ≤ 2 nM, even more preferably ≤ 1 nM, even more preferably ≤ 0.6 nM, even more preferably ≤ 0.5 nM, and most
25 preferably ≤ 0.4 nM. The affinity can be measured for example in a BIAcore assay or in a Scatchard assay. Other methods of determining the affinity are also well-known to the skilled person. The affinity of the first domain for macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM is preferably ≤ 15 nM, more preferably ≤ 10 nM, even more preferably ≤ 5 nM, even more preferably ≤ 1 nM, even more preferably ≤ 0.5 nM, even more preferably ≤ 0.1 nM, and
30 most preferably ≤ 0.05 nM or even ≤ 0.01 nM.

[187] Preferably the affinity gap of the antigen-binding molecules according to the invention for binding macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM versus human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM [ma
35 CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM: hu CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM (as determined e.g. by surface plasmon resonance analysis such as BiaCoreTM or by Scatchard analysis) is < 100 , preferably < 20 ,

more preferably <15, further preferably <10, even more preferably <8, more preferably <6 and most preferably <2. Preferred ranges for the affinity gap of the antigen-binding molecules according to the invention for binding macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM versus human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM are between 0.1 and 20, more preferably between 0.2 and 10, even more preferably between 0.3 and 6, even more preferably between 0.5 and 3 or between 0.5 and 2.5, and most preferably between 0.5 and 2 or between 0.6 and 2.

[188] The second and the fourth binding domain of the antigen-binding molecule of the invention typically binds to human CD3 epsilon and/or to *Macaca* CD3 epsilon. In a preferred embodiment, where a selectivity gap is achieved, the second and the fourth binding domain, or alternatively, the first and the third binding domain, further binds to *Callithrix jacchus*, *Saguinus Oedipus* or *Saimiri sciureus* CD3 epsilon. *Callithrix jacchus* and *Saguinus oedipus* are both new world primate belonging to the family of *Callitrichidae*, while *Saimiri sciureus* is a new world primate belonging to the family of *Cebidae*. Said binding domains may preferably selected form sequences identified herein as “I2L” (or synonymously “I2L0”), “I2M” and “I2M2”, more preferably as “I2L” or “I2L0”.

[189] It is preferred for the antigen-binding molecule of the present invention that the preferably second and fourth binding domain which binds to an extracellular epitope of the human and/or the *Macaca* CD3 epsilon chain comprises a VL region comprising CDR-L1, CDR-L2 and CDR-L3 selected from:

- (a) VL region comprising CDR-L1, CDR-L2 and CDR-L3 selected from SEQ ID NOs 40 to 42, 48 to 50, 56 to 58, 64 to 66, 72 to 74 439 to 441, preferably 64 to 66
- (b) CDR-L1 as depicted in SEQ ID NO: 27 of WO 2008/119567, CDR-L2 as depicted in SEQ ID NO: 28 of WO 2008/119567 and CDR-L3 as depicted in SEQ ID NO: 29 of WO 2008/119567;
- (c) CDR-L1 as depicted in SEQ ID NO: 117 of WO 2008/119567, CDR-L2 as depicted in SEQ ID NO: 118 of WO 2008/119567 and CDR-L3 as depicted in SEQ ID NO: 119 of WO 2008/119567;
- (d) CDR-L1 as depicted in SEQ ID NO: 153 of WO 2008/119567, CDR-L2 as depicted in SEQ ID NO: 154 of WO 2008/119567 and CDR-L3 as depicted in SEQ ID NO: 155 of WO 2008/119567; and
- (e) VL region comprising CDR-L1, CDR-L2 and CDR-L3 of SEQ ID NOs 420 to 422.

[190] In a furthermore preferred embodiment of the antigen-binding molecule of the present invention, the preferably second and fourth binding domain which binds to an extracellular epitope of the human and/or the *Macaca* CD3 epsilon chain comprises a VH region comprising CDR-H1, CDR-H2 and CDR-H3 selected from:

- (a) VH region comprising CDR-H1, CDR-H2 and CDR-H3 selected from SEQ ID NOs 37 to 39, 45 to 47, 53 to 55, 61 to 63, 69 to 71 and 436 to 438, preferably 61 to 63;

- (b) CDR-H1 as depicted in SEQ ID NO: 12 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 13 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 14 of WO 2008/119567;
- (c) CDR-H1 as depicted in SEQ ID NO: 30 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 31 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 32 of WO 2008/119567;
- 5 (d) CDR-H1 as depicted in SEQ ID NO: 48 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 49 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 50 of WO 2008/119567;
- (e) CDR-H1 as depicted in SEQ ID NO: 66 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 67 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 68 of WO 2008/119567;
- (f) CDR-H1 as depicted in SEQ ID NO: 84 of WO 2008/119567, CDR-H2 as depicted in SEQ ID
10 NO: 85 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 86 of WO 2008/119567;
- (g) CDR-H1 as depicted in SEQ ID NO: 102 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 103 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 104 of WO 2008/119567;
- (h) CDR-H1 as depicted in SEQ ID NO: 120 of WO 2008/119567, CDR-H2 as depicted in
15 SEQ ID NO: 121 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 122 of WO 2008/119567;
- (i) CDR-H1 as depicted in SEQ ID NO: 138 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 139 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 140 of WO 2008/119567;
- 20 (j) CDR-H1 as depicted in SEQ ID NO: 156 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 157 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 158 of WO 2008/119567;
- (k) CDR-H1 as depicted in SEQ ID NO: 174 of WO 2008/119567, CDR-H2 as depicted in SEQ ID NO: 175 of WO 2008/119567 and CDR-H3 as depicted in SEQ ID NO: 176 of
25 WO 2008/119567; and
- (l) VH region comprising CDR-H 1, CDR-H2 and CDR-H3 of SEQ ID NOs 423 to 425.

[191] In a preferred embodiment of the antigen-binding molecule of the invention the above described three groups of VL CDRs are combined with the above described ten groups of VH CDRs within the third binding domain to form (30) groups, each comprising CDR-L 1-3 and CDR-H 1-3.

30 [192] It is preferred for the antigen-binding molecule of the present invention that the third domain which binds to CD3 comprises a VL region selected from the group consisting of those depicted in SEQ ID NOs: 17, 21, 35, 39, 53, 57, 71, 75, 89, 93, 107, 111, 125, 129, 143, 147, 161, 165, 179 or 183 of WO 2008/119567 or, preferably, as depicted in SEQ ID NO: 44, 52, 60, 68 and 76, preferably 68 according to the present invention.

35 [193] It is also preferred that the third domain which binds to CD3 comprises a VH region selected from the group consisting of those depicted in SEQ ID NO: 15, 19, 33, 37, 51, 55, 69, 73, 87, 91, 105,

109, 123, 127, 141, 145, 159, 163, 177 or 181 of WO 2008/119567 or, preferably, as depicted in SEQ ID NO: SEQ ID NOs 43, 51, 59, 67 and 75, preferably 67 according to the present invention.

[194] More preferably, the antigen-binding molecule of the present invention is characterized by a preferably second and fourth domain which binds to CD3 comprising a VL region and a VH region
5 selected from the group consisting of:

- (a) a VL region selected from SEQ ID NOs 44, 52, 60, 68, 76 and 443, and a VH region selected from SEQ ID NOs 43, 51, 59, 67, 75 and 442;
- (b) a VL region as depicted in SEQ ID NO: 17 or 21 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 15 or 19 of WO 2008/119567;
- 10 (c) a VL region as depicted in SEQ ID NO: 35 or 39 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 33 or 37 of WO 2008/119567;
- (d) a VL region as depicted in SEQ ID NO: 53 or 57 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 51 or 55 of WO 2008/119567;
- (e) a VL region as depicted in SEQ ID NO: 71 or 75 of WO 2008/119567 and a VH region as
15 depicted in SEQ ID NO: 69 or 73 of WO 2008/119567;
- (f) a VL region as depicted in SEQ ID NO: 89 or 93 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 87 or 91 of WO 2008/119567;
- (g) a VL region as depicted in SEQ ID NO: 107 or 111 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 105 or 109 of WO 2008/119567;
- 20 (h) a VL region as depicted in SEQ ID NO: 125 or 129 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 123 or 127 of WO 2008/119567;
- (i) a VL region as depicted in SEQ ID NO: 143 or 147 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 141 or 145 of WO 2008/119567;
- (j) a VL region as depicted in SEQ ID NO: 161 or 165 of WO 2008/119567 and a VH region as
25 depicted in SEQ ID NO: 159 or 163 of WO 2008/119567; and
- (k) a VL region as depicted in SEQ ID NO: 179 or 183 of WO 2008/119567 and a VH region as depicted in SEQ ID NO: 177 or 181 of WO 2008/119567.

[195] Also preferred in connection with the antigen-binding molecule of the present invention is a second and fourth domain which binds to CD3 comprising a VL region as depicted in SEQ ID NO: 68
30 and a VH region as depicted in SEQ ID NO: 67.

[196] According to a preferred embodiment of the antigen-binding molecule of the present invention, the first and/or the third domain have the following format: The pairs of VH regions and VL regions are in the format of a single chain antibody (scFv). The VH and VL regions are arranged in the order VH-VL or VL-VH. It is preferred that the VH-region is positioned N-terminally of a
35 linker sequence, and the VL-region is positioned C-terminally of the linker sequence.

[197] The invention further provides an antigen-binding molecule comprising or having an amino acid sequence (full bispecific antigen-binding molecule) selected from the group consisting of any of 673, 676, 679, 682, 685, 688, 691, 694, 697, 700, 703, 706, 709, 712, 715, 718, 721, 724, 727, 730, 733, 736, 739, 742, 745, 748, 751, 754, 757, 760, 763, 766, 769, 772, 775, 778, 781, 784, 787, 790, 5 793, 796, 799, 802, 805, 808, 811, 814, 817, 820, 823, 826, 829, 832, 835, 838, 841, 844, 847, 850, 853, 856, 859, 862, 865, 868, 871, 1437, 1440, 1443, 1446, 1449, 1452, 1455, 1458, 1461, 1464, 1467, 1470, 1473, 1476, 1479, 1482, 1485, 1488, 1499, 1667, 1670, 1673, 1676, 1679, 1682, 1685, 1688, 1691, 1694, 1697, 1700, 1703, 1706, 1709, 1712, 1715, 1718, 1721, 1724, 1727, 1730, 1733, 1736, 1739, 1742, 1745, 1748, 1751, 1754, 1757, 1760, 1763, 1766, 1769, 1772, 1775, 1778, 1781, 10 1784, 1787, 1790, 1793, 1796, 1799, 1802, 1805, 1808, 1811, 1814, 1817, 1820, 1823, 1826, and 1829, preferably 1437, or having an amino acid sequence having at least 90, 91, 92, 93, 94 95, 96, 97, 98 or 99% identity to said sequences.

[198] Covalent modifications of the antigen-binding molecules are also included within the scope of this invention, and are generally, but not always, done post-translationally. For example, several types 15 of covalent modifications of the antigen-binding molecule are introduced into the molecule by reacting specific amino acid residues of the antigen-binding molecule with an organic derivatizing agent that is capable of reacting with selected side chains or the N- or C-terminal residues.

[199] Cysteinyl residues most commonly are reacted with α -haloacetates (and corresponding amines), such as chloroacetic acid or chloroacetamide, to give carboxymethyl or carboxyamidomethyl 20 derivatives. Cysteinyl residues also are derivatized by reaction with bromotrifluoroacetone, α -bromo- β -(5-imidazolyl)propionic acid, chloroacetyl phosphate, N-alkylmaleimides, 3-nitro-2-pyridyl disulfide, methyl 2-pyridyl disulfide, p-chloromercuribenzoate, 2-chloromercuri-4-nitrophenol, or chloro-7-nitrobenzo-2-oxa-1,3-diazole.

[200] Histidyl residues are derivatized by reaction with diethylpyrocarbonate at pH 5.5-7.0 because 25 this agent is relatively specific for the histidyl side chain. Para-bromophenacyl bromide also is useful; the reaction is preferably performed in 0.1 M sodium cacodylate at pH 6.0. Lysinyl and amino terminal residues are reacted with succinic or other carboxylic acid anhydrides. Derivatization with these agents has the effect of reversing the charge of the lysinyl residues. Other suitable reagents for derivatizing alpha-amino-containing residues include imidoesters such as methyl picolinimidate; 30 pyridoxal phosphate; pyridoxal; chloroborohydride; trinitrobenzenesulfonic acid; O-methylisourea; 2,4-pentanedione; and transaminase-catalyzed reaction with glyoxylate.

[201] Arginyl residues are modified by reaction with one or several conventional reagents, among them phenylglyoxal, 2,3-butanedione, 1,2-cyclohexanedione, and ninhydrin. Derivatization of arginine residues requires that the reaction be performed in alkaline conditions because of the high pKa of the

guanidine functional group. Furthermore, these reagents may react with the groups of lysine as well as the arginine epsilon-amino group.

[202] The specific modification of tyrosyl residues may be made, with particular interest in introducing spectral labels into tyrosyl residues by reaction with aromatic diazonium compounds or tetranitromethane. Most commonly, N-acetylimidazole and tetranitromethane are used to form O-acetyl tyrosyl species and 3-nitro derivatives, respectively. Tyrosyl residues are iodinated using ¹²⁵I or ¹³¹I to prepare labeled proteins for use in radioimmunoassay, the chloramine T method described above being suitable.

[203] Carboxyl side groups (aspartyl or glutamyl) are selectively modified by reaction with carbodiimides (R'-N=C=N-R'), where R and R' are optionally different alkyl groups, such as 1-cyclohexyl-3-(2-morpholinyl-4-ethyl) carbodiimide or 1-ethyl-3-(4-azonia-4,4-dimethylpentyl) carbodiimide. Furthermore, aspartyl and glutamyl residues are converted to asparaginyl and glutaminyl residues by reaction with ammonium ions.

[204] Derivatization with bifunctional agents is useful for crosslinking the antigen-binding molecules of the present invention to a water-insoluble support matrix or surface for use in a variety of methods. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), and bifunctional maleimides such as bis-N-maleimido-1,8-octane. Derivatizing agents such as methyl-3-[(p-azidophenyl)dithio]propioimidate yield photoactivatable intermediates that are capable of forming crosslinks in the presence of light. Alternatively, reactive water-insoluble matrices such as cyanogen bromide-activated carbohydrates and the reactive substrates as described in U.S. Pat. Nos. 3,969,287; 3,691,016; 4,195,128; 4,247,642; 4,229,537; and 4,330,440 are employed for protein immobilization.

[205] Glutaminyl and asparaginyl residues are frequently deamidated to the corresponding glutamyl and aspartyl residues, respectively. Alternatively, these residues are deamidated under mildly acidic conditions. Either form of these residues falls within the scope of this invention.

[206] Other modifications include hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the α -amino groups of lysine, arginine, and histidine side chains (T. E. Creighton, *Proteins: Structure and Molecular Properties*, W. H. Freeman & Co., San Francisco, 1983, pp. 79-86), acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

[207] Another type of covalent modification of the antigen-binding molecules included within the scope of this invention comprises altering the glycosylation pattern of the protein. As is known in the

art, glycosylation patterns can depend on both the sequence of the protein (e.g., the presence or absence of particular glycosylation amino acid residues, discussed below), or the host cell or organism in which the protein is produced. Particular expression systems are discussed below.

5 [208] Glycosylation of polypeptides is typically either N-linked or O-linked. N-linked refers to the attachment of the carbohydrate moiety to the side chain of an asparagine residue. The tri-peptide sequences asparagine-X-serine and asparagine-X-threonine, where X is any amino acid except proline, are the recognition sequences for enzymatic attachment of the carbohydrate moiety to the asparagine side chain. Thus, the presence of either of these tri-peptide sequences in a polypeptide creates a potential glycosylation site. O-linked glycosylation refers to the attachment of one of the sugars N-
10 acetylgalactosamine, galactose, or xylose, to a hydroxyamino acid, most commonly serine or threonine, although 5-hydroxyproline or 5-hydroxylysine may also be used.

[209] Addition of glycosylation sites to the antigen-binding molecule is conveniently accomplished by altering the amino acid sequence such that it contains one or more of the above-described tri-peptide sequences (for N-linked glycosylation sites). The alteration may also be made by the addition
15 of, or substitution by, one or more serine or threonine residues to the starting sequence (for O-linked glycosylation sites). For ease, the amino acid sequence of an antigen-binding molecule is preferably altered through changes at the DNA level, particularly by mutating the DNA encoding the polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

[210] Another means of increasing the number of carbohydrate moieties on the antigen-binding
20 molecule is by chemical or enzymatic coupling of glycosides to the protein. These procedures are advantageous in that they do not require production of the protein in a host cell that has glycosylation capabilities for N- and O-linked glycosylation. Depending on the coupling mode used, the sugar(s) may be attached to (a) arginine and histidine, (b) free carboxyl groups, (c) free sulfhydryl groups such as those of cysteine, (d) free hydroxyl groups such as those of serine, threonine, or hydroxyproline, (e)
25 aromatic residues such as those of phenylalanine, tyrosine, or tryptophan, or (f) the amide group of glutamine. These methods are described in WO 87/05330, and in Aplin and Wriston, 1981, *CRC Crit. Rev. Biochem.*, pp. 259-306.

[211] Removal of carbohydrate moieties present on the starting antigen-binding molecule may be accomplished chemically or enzymatically. Chemical deglycosylation requires exposure of the protein
30 to the compound trifluoromethanesulfonic acid, or an equivalent compound. This treatment results in the cleavage of most or all sugars except the linking sugar (N-acetylglucosamine or N-acetylgalactosamine), while leaving the polypeptide intact. Chemical deglycosylation is described by Hakimuddin *et al.*, 1987, *Arch. Biochem. Biophys.* 259:52 and by Edge *et al.*, 1981, *Anal. Biochem.* 118:131. Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of
35 a variety of endo- and exo-glycosidases as described by Thotakura *et al.*, 1987, *Meth. Enzymol.*

138:350. Glycosylation at potential glycosylation sites may be prevented by the use of the compound tunicamycin as described by Duskin *et al.*, 1982, J. Biol. Chem. 257:3105. Tunicamycin blocks the formation of protein-N-glycoside linkages.

5 [212] Other modifications of the antigen-binding molecule are also contemplated herein. For example, another type of covalent modification of the antigen-binding molecule comprises linking the antigen-binding molecule to various non-proteinaceous polymers, including, but not limited to, various polyols such as polyethylene glycol, polypropylene glycol, polyoxyalkylenes, or copolymers of polyethylene glycol and polypropylene glycol, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337. In addition, as is known in the art, amino
10 acid substitutions may be made in various positions within the antigen-binding molecule, e.g. in order to facilitate the addition of polymers such as PEG.

[213] In some embodiments, the covalent modification of the antigen-binding molecules of the invention comprises the addition of one or more labels. The labelling group may be coupled to the antigen-binding molecule *via* spacer arms of various lengths to reduce potential steric hindrance.
15 Various methods for labelling proteins are known in the art and can be used in performing the present invention. The term “label” or “labelling group” refers to any detectable label. In general, labels fall into a variety of classes, depending on the assay in which they are to be detected – the following examples include, but are not limited to:

- a) isotopic labels, which may be radioactive or heavy isotopes, such as radioisotopes or radionuclides
20 (e.g., ³H, ¹⁴C, ¹⁵N, ³⁵S, ⁸⁹Zr, ⁹⁰Y, ⁹⁹Tc, ¹¹¹In, ¹²⁵I, ¹³¹I)
- b) magnetic labels (e.g., magnetic particles)
- c) redox active moieties
- d) optical dyes (including, but not limited to, chromophores, phosphors and fluorophores) such as fluorescent groups (e.g., FITC, rhodamine, lanthanide phosphors), chemiluminescent groups, and
25 fluorophores which can be either “small molecule” fluors or proteinaceous fluors
- e) enzymatic groups (e.g. horseradish peroxidase, β -galactosidase, luciferase, alkaline phosphatase)
- f) biotinylated groups
- g) predetermined polypeptide epitopes recognized by a secondary reporter (e.g., leucine zipper pair sequences, binding sites for secondary antibodies, metal binding domains, epitope tags, etc.)

30 [214] By “fluorescent label” is meant any molecule that may be detected *via* its inherent fluorescent properties. Suitable fluorescent labels include, but are not limited to, fluorescein, rhodamine, tetramethylrhodamine, eosin, erythrosin, coumarin, methyl-coumarins, pyrene, Malacite green, stilbene, Lucifer Yellow, Cascade BlueJ, Texas Red, IAEDANS, EDANS, BODIPY FL, LC Red 640, Cy 5, Cy 5.5, LC Red 705, Oregon green, the Alexa-Fluor dyes (Alexa Fluor 350, Alexa Fluor 430,
35 Alexa Fluor 488, Alexa Fluor 546, Alexa Fluor 568, Alexa Fluor 594, Alexa Fluor 633, Alexa Fluor 660, Alexa Fluor 680), Cascade Blue, Cascade Yellow and R-phycoerythrin (PE) (Molecular Probes,

Eugene, OR), FITC, Rhodamine, and Texas Red (Pierce, Rockford, IL), Cy5, Cy5.5, Cy7 (Amersham Life Science, Pittsburgh, PA). Suitable optical dyes, including fluorophores, are described in Molecular Probes Handbook by Richard P. Haugland.

[215] Suitable proteinaceous fluorescent labels also include, but are not limited to, green fluorescent protein, including a Renilla, Ptilosarcus, or Aequorea species of GFP (Chalfie *et al.*, 1994, *Science* 263:802-805), EGFP (Clontech Laboratories, Inc., Genbank Accession Number U55762), blue fluorescent protein (BFP, Quantum Biotechnologies, Inc. 1801 de Maisonneuve Blvd. West, 8th Floor, Montreal, Quebec, Canada H3H 1J9; Stauber, 1998, *Biotechniques* 24:462-471; Heim *et al.*, 1996, *Curr. Biol.* 6:178-182), enhanced yellow fluorescent protein (EYFP, Clontech Laboratories, Inc.), luciferase (Ichiki *et al.*, 1993, *J. Immunol.* 150:5408-5417), β galactosidase (Nolan *et al.*, 1988, *Proc. Natl. Acad. Sci. U.S.A.* 85:2603-2607) and Renilla (WO92/15673, WO95/07463, WO98/14605, WO98/26277, WO99/49019, U.S. Patent Nos. 5,292,658; 5,418,155; 5,683,888; 5,741,668; 5,777,079; 5,804,387; 5,874,304; 5,876,995; 5,925,558).

[216] The antigen-binding molecule of the invention may also comprise additional domains, which are *e.g.* helpful in the isolation of the molecule or relate to an adapted pharmacokinetic profile of the molecule. Domains helpful for the isolation of an antigen-binding molecule may be selected from peptide motives or secondarily introduced moieties, which can be captured in an isolation method, *e.g.* an isolation column. Non-limiting embodiments of such additional domains comprise peptide motives known as Myc-tag, HAT-tag, HA-tag, TAP-tag, GST-tag, chitin binding domain (CBD-tag), maltose binding protein (MBP-tag), Flag-tag, Strep-tag and variants thereof (*e.g.* StrepII-tag) and His-tag. All herein disclosed antigen-binding molecules may comprise a His-tag domain, which is generally known as a repeat of consecutive His residues in the amino acid sequence of a molecule, preferably of five, and more preferably of six His residues (hexa-histidine). The His-tag may be located *e.g.* at the N- or C-terminus of the antigen-binding molecule, preferably it is located at the C-terminus. Most preferably, a hexa-histidine tag (HHHHHH) (SEQ ID NO:16) is linked via peptide bond to the C-terminus of the antigen-binding molecule according to the invention. Additionally, a conjugate system of PLGA-PEG-PLGA may be combined with a poly-histidine tag for sustained release application and improved pharmacokinetic profile.

[217] Amino acid sequence modifications of the antigen-binding molecules described herein are also contemplated. For example, it may be desirable to improve the binding affinity and/or other biological properties of the antigen-binding molecule. Amino acid sequence variants of the antigen-binding molecules are prepared by introducing appropriate nucleotide changes into the antigen-binding molecules nucleic acid, or by peptide synthesis. All of the below described amino acid sequence modifications should result in an antigen-binding molecule which still retains the desired biological activity (binding to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM and to CD3) of the unmodified parental molecule.

[218] The term “amino acid” or “amino acid residue” typically refers to an amino acid having its art recognized definition such as an amino acid selected from the group consisting of: alanine (Ala or A); arginine (Arg or R); asparagine (Asn or N); aspartic acid (Asp or D); cysteine (Cys or C); glutamine (Gln or Q); glutamic acid (Glu or E); glycine (Gly or G); histidine (His or H); isoleucine (Ile or I); leucine (Leu or L); lysine (Lys or K); methionine (Met or M); phenylalanine (Phe or F); proline (Pro or P); serine (Ser or S); threonine (Thr or T); tryptophan (Trp or W); tyrosine (Tyr or Y); and valine (Val or V), although modified, synthetic, or rare amino acids may be used as desired. Generally, amino acids can be grouped as having a nonpolar side chain (e.g., Ala, Cys, Ile, Leu, Met, Phe, Pro, Val); a negatively charged side chain (e.g., Asp, Glu); a positively charged side chain (e.g., Arg, His, Lys); or an uncharged polar side chain (e.g., Asn, Cys, Gln, Gly, His, Met, Phe, Ser, Thr, Trp, and Tyr).

[219] Amino acid modifications include, for example, deletions from, and/or insertions into, and/or substitutions of, residues within the amino acid sequences of the antigen-binding molecules. Any combination of deletion, insertion, and substitution is made to arrive at the final construct, provided that the final construct possesses the desired characteristics. The amino acid changes also may alter post-translational processes of the antigen-binding molecules, such as changing the number or position of glycosylation sites.

[220] For example, 1, 2, 3, 4, 5, or 6 amino acids may be inserted, substituted or deleted in each of the CDRs (of course, dependent on their length), while 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, or 25 amino acids may be inserted, substituted or deleted in each of the FRs. Preferably, amino acid sequence insertions into the antigen-binding molecule include amino- and/or carboxyl-terminal fusions ranging in length from 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 residues to polypeptides containing a hundred or more residues, as well as intra-sequence insertions of single or multiple amino acid residues. An insertional variant of the antigen-binding molecule of the invention includes the fusion to the N-terminus or to the C-terminus of the antigen-binding molecule of an enzyme or the fusion to a polypeptide.

[221] The sites of greatest interest for substitutional mutagenesis include (but are not limited to) the CDRs of the heavy and/or light chain, in particular the hypervariable regions, but FR alterations in the heavy and/or light chain are also contemplated. The substitutions are preferably conservative substitutions as described herein. Preferably, 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 amino acids may be substituted in a CDR, while 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, or 25 amino acids may be substituted in the framework regions (FRs), depending on the length of the CDR or FR. For example, if a CDR sequence encompasses 6 amino acids, it is envisaged that one, two or three of these amino acids are substituted. Similarly, if a CDR sequence encompasses 15 amino acids it is envisaged that one, two, three, four, five or six of these amino acids are substituted.

[222] A useful method for identification of certain residues or regions of the antigen-binding molecules that are preferred locations for mutagenesis is called "alanine scanning mutagenesis" as described by Cunningham and Wells in *Science*, 244: 1081-1085 (1989). Here, a residue or group of target residues within the antigen-binding molecule is/are identified (e.g. charged residues such as arg, asp, his, lys, and glu) and replaced by a neutral or negatively charged amino acid (most preferably alanine or polyalanine) to affect the interaction of the amino acids with the epitope.

[223] Those amino acid locations demonstrating functional sensitivity to the substitutions are then refined by introducing further or other variants at, or for, the sites of substitution. Thus, while the site or region for introducing an amino acid sequence variation is predetermined, the nature of the mutation *per se* needs not to be predetermined. For example, to analyze or optimize the performance of a mutation at a given site, alanine scanning or random mutagenesis may be conducted at a target codon or region, and the expressed antigen-binding molecule variants are screened for the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in the DNA having a known sequence are well known, for example, M13 primer mutagenesis and PCR mutagenesis. Screening of the mutants is done using assays of antigen binding activities, such as CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM or CD3 binding.

[224] Generally, if amino acids are substituted in one or more or all of the CDRs of the heavy and/or light chain, it is preferred that the then-obtained "substituted" sequence is at least 60% or 65%, more preferably 70% or 75%, even more preferably 80% or 85%, and particularly preferably 90% or 95% identical to the "original" CDR sequence. This means that it is dependent of the length of the CDR to which degree it is identical to the "substituted" sequence. For example, a CDR having 5 amino acids is preferably 80% identical to its substituted sequence in order to have at least one amino acid substituted. Accordingly, the CDRs of the antigen-binding molecule may have different degrees of identity to their substituted sequences, e.g., CDRL1 may have 80%, while CDRL3 may have 90%.

[225] Preferred substitutions (or replacements) are conservative substitutions. However, any substitution (including non-conservative substitution or one or more from the "exemplary substitutions" listed in Table 3, below) is envisaged as long as the antigen-binding molecule retains its capability to bind to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM via the first domain and to CD3 epsilon via the second domain and/or its CDRs have an identity to the then substituted sequence (at least 60% or 65%, more preferably 70% or 75%, even more preferably 80% or 85%, and particularly preferably 90% or 95% identical to the "original" CDR sequence).

[226] Conservative substitutions are shown in Table 3 under the heading of "preferred substitutions". If such substitutions result in a change in biological activity, then more substantial changes, denominated "exemplary substitutions" in Table 3, or as further described below in reference to amino acid classes, may be introduced and the products screened for a desired characteristic.

Table 3: Amino acid substitutions

Original	Exemplary Substitutions	Preferred Substitutions
Ala (A)	val, leu, ile	Val
Arg (R)	lys, gln, asn	Lys
Asn (N)	gln, his, asp, lys, arg	Gln
Asp (D)	glu, asn	Glu
Cys (C)	ser, ala	ser
Gln (Q)	asn, glu	asn
Glu (E)	asp, gln	asp
Gly (G)	Ala	ala
His (H)	asn, gln, lys, arg	arg
Ile (I)	leu, val, met, ala, phe	leu
Leu (L)	norleucine, ile, val, met, ala	ile
Lys (K)	arg, gln, asn	arg
Met (M)	leu, phe, ile	leu
Phe (F)	leu, val, ile, ala, tyr	tyr
Pro (P)	Ala	ala
Ser (S)	Thr	thr
Thr (T)	Ser	ser
Trp (W)	tyr, phe	tyr
Tyr (Y)	trp, phe, thr, ser	phe
Val (V)	ile, leu, met, phe, ala	leu

[227] Substantial modifications in the biological properties of the antigen-binding molecule of the present invention are accomplished by selecting substitutions that differ significantly in their effect on maintaining (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain. Naturally occurring residues are divided into groups based on common side-chain properties: (1) hydrophobic: norleucine, met, ala, val, leu, ile; (2) neutral hydrophilic: cys, ser, thr; asn, gln (3) acidic: asp, glu; (4) basic: his, lys, arg; (5) residues that influence chain orientation: gly, pro; and (6) aromatic : trp, tyr, phe.

[228] Non-conservative substitutions will entail exchanging a member of one of these classes for another class. Any cysteine residue not involved in maintaining the proper conformation of the antigen-binding molecule may be substituted, generally with serine, to improve the oxidative stability of the molecule and prevent aberrant crosslinking. Conversely, cysteine bond(s) may be added to the

antibody to improve its stability (particularly where the antibody is an antibody fragment such as an Fv fragment).

[229] For amino acid sequences, sequence identity and/or similarity is determined by using standard techniques known in the art, including, but not limited to, the local sequence identity algorithm of Smith and Waterman, 1981, *Adv. Appl. Math.* 2:482, the sequence identity alignment algorithm of Needleman and Wunsch, 1970, *J. Mol. Biol.* 48:443, the search for similarity method of Pearson and Lipman, 1988, *Proc. Nat. Acad. Sci. U.S.A.* 85:2444, computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Drive, Madison, Wis.), the Best Fit sequence program described by Devereux *et al.*, 1984, *Nucl. Acid Res.* 12:387-395, preferably using the default settings, or by inspection. Preferably, percent identity is calculated by FastDB based upon the following parameters: mismatch penalty of 1; gap penalty of 1; gap size penalty of 0.33; and joining penalty of 30, "Current Methods in Sequence Comparison and Analysis," *Macromolecule Sequencing and Synthesis, Selected Methods and Applications*, pp 127-149 (1988), Alan R. Liss, Inc.

[230] An example of a useful algorithm is PILEUP. PILEUP creates a multiple sequence alignment from a group of related sequences using progressive, pairwise alignments. It can also plot a tree showing the clustering relationships used to create the alignment. PILEUP uses a simplification of the progressive alignment method of Feng & Doolittle, 1987, *J. Mol. Evol.* 35:351-360; the method is similar to that described by Higgins and Sharp, 1989, *CABIOS* 5:151-153. Useful PILEUP parameters including a default gap weight of 3.00, a default gap length weight of 0.10, and weighted end gaps.

[231] Another example of a useful algorithm is the BLAST algorithm, described in: Altschul *et al.*, 1990, *J. Mol. Biol.* 215:403-410; Altschul *et al.*, 1997, *Nucleic Acids Res.* 25:3389-3402; and Karin *et al.*, 1993, *Proc. Natl. Acad. Sci. U.S.A.* 90:5873-5787. A particularly useful BLAST program is the WU-BLAST-2 program which was obtained from Altschul *et al.*, 1996, *Methods in Enzymology* 266:460-480. WU-BLAST-2 uses several search parameters, most of which are set to the default values. The adjustable parameters are set with the following values: overlap span=1, overlap fraction=0.125, word threshold (T)=II. The HSP S and HSP S2 parameters are dynamic values and are established by the program itself depending upon the composition of the particular sequence and composition of the particular database against which the sequence of interest is being searched; however, the values may be adjusted to increase sensitivity.

[232] An additional useful algorithm is gapped BLAST as reported by Altschul *et al.*, 1993, *Nucl. Acids Res.* 25:3389-3402. Gapped BLAST uses BLOSUM-62 substitution scores; threshold T parameter set to 9; the two-hit method to trigger ungapped extensions, charges gap lengths of k a cost of 10+k; Xu set to 16, and Xg set to 40 for database search stage and to 67 for the output stage of the algorithms. Gapped alignments are triggered by a score corresponding to about 22 bits.

[233] Generally, the amino acid homology, similarity, or identity between individual variant CDRs or VH / VL sequences are at least 60% to the sequences depicted herein, and more typically with preferably increasing homologies or identities of at least 65% or 70%, more preferably at least 75% or 80%, even more preferably at least 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, and almost 100%. In a similar manner, “percent (%) nucleic acid sequence identity” with respect to the nucleic acid sequence of the binding proteins identified herein is defined as the percentage of nucleotide residues in a candidate sequence that are identical with the nucleotide residues in the coding sequence of the antigen-binding molecule. A specific method utilizes the BLASTN module of WU-BLAST-2 set to the default parameters, with overlap span and overlap fraction set to 1 and 0.125, respectively.

[234] Generally, the nucleic acid sequence homology, similarity, or identity between the nucleotide sequences encoding individual variant CDRs or VH / VL sequences and the nucleotide sequences depicted herein are at least 60%, and more typically with preferably increasing homologies or identities of at least 65%, 70%, 75%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99%, and almost 100%. Thus, a “variant CDR” or a “variant VH / VL region” is one with the specified homology, similarity, or identity to the parent CDR / VH / VL of the invention, and shares biological function, including, but not limited to, at least 60%, 65%, 70%, 75%, 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% of the specificity and/or activity of the parent CDR or VH / VL.

[235] In one embodiment, the percentage of identity to human germline of the antigen-binding molecules according to the invention is $\geq 70\%$ or $\geq 75\%$, more preferably $\geq 80\%$ or $\geq 85\%$, even more preferably $\geq 90\%$, and most preferably $\geq 91\%$, $\geq 92\%$, $\geq 93\%$, $\geq 94\%$, $\geq 95\%$ or even $\geq 96\%$. Identity to human antibody germline gene products is thought to be an important feature to reduce the risk of therapeutic proteins to elicit an immune response against the drug in the patient during treatment. Hwang & Foote (“Immunogenicity of engineered antibodies”; Methods 36 (2005) 3-10) demonstrate that the reduction of non-human portions of drug antigen-binding molecules leads to a decrease of risk to induce anti-drug antibodies in the patients during treatment. By comparing an exhaustive number of clinically evaluated antibody drugs and the respective immunogenicity data, the trend is shown that humanization of the V-regions of antibodies makes the protein less immunogenic (average 5.1 % of patients) than antibodies carrying unaltered non-human V regions (average 23.59 % of patients). A higher degree of identity to human sequences is hence desirable for V-region based protein therapeutics in the form of antigen-binding molecules. For this purpose of determining the germline identity, the V-regions of VL can be aligned with the amino acid sequences of human germline V segments and J segments (<http://vbase.mrc-cpe.cam.ac.uk/>) using Vector NTI software and the amino acid sequence calculated by dividing the identical amino acid residues by the total number of amino

acid residues of the VL in percent. The same can be for the VH segments (<http://vbase.mrc-cpe.cam.ac.uk/>) with the exception that the VH CDR3 may be excluded due to its high diversity and a lack of existing human germline VH CDR3 alignment partners. Recombinant techniques can then be used to increase sequence identity to human antibody germline genes.

5 [236] In a further embodiment, the bispecific antigen-binding molecules of the present invention exhibit high monomer yields under standard research scale conditions, *e.g.*, in a standard two-step purification process. Preferably the monomer yield of the antigen-binding molecules according to the invention is ≥ 0.25 mg/L supernatant, more preferably ≥ 0.5 mg/L, even more preferably ≥ 1 mg/L, and most preferably ≥ 3 mg/L supernatant.

10 [237] Likewise, the yield of the dimeric antigen-binding molecule isoforms and hence the monomer percentage (*i.e.*, monomer:(monomer+dimer)) of the antigen-binding molecules can be determined. The productivity of monomeric and dimeric antigen-binding molecules and the calculated monomer percentage can *e.g.* be obtained in the SEC purification step of culture supernatant from standardized research-scale production in roller bottles. In one embodiment, the monomer percentage of the
15 antigen-binding molecules is $\geq 80\%$, more preferably $\geq 85\%$, even more preferably $\geq 90\%$, and most preferably $\geq 95\%$.

[238] In one embodiment, the antigen-binding molecules have a preferred plasma stability (ratio of EC50 with plasma to EC50 w/o plasma) of ≤ 5 or ≤ 4 , more preferably ≤ 3.5 or ≤ 3 , even more preferably ≤ 2.5 or ≤ 2 , and most preferably ≤ 1.5 or ≤ 1 . The plasma stability of an antigen-binding
20 molecule can be tested by incubation of the construct in human plasma at 37°C for 24 hours followed by EC50 determination in a ⁵¹chromium release cytotoxicity assay. The effector cells in the cytotoxicity assay can be stimulated enriched human CD8 positive T cells. Target cells can *e.g.* be CHO cells transfected with human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM. The effector to target cell (E:T) ratio can be chosen as 10:1 or 5:1. The human plasma
25 pool used for this purpose is derived from the blood of healthy donors collected by EDTA coated syringes. Cellular components are removed by centrifugation and the upper plasma phase is collected and subsequently pooled. As control, antigen-binding molecules are diluted immediately prior to the cytotoxicity assay in RPMI-1640 medium. The plasma stability is calculated as ratio of EC50 (after plasma incubation) to EC50 (control).

30 [239] It is furthermore preferred that the monomer to dimer conversion of antigen-binding molecules of the invention is low. The conversion can be measured under different conditions and analyzed by high performance size exclusion chromatography. For example, incubation of the monomeric isoforms of the antigen-binding molecules can be carried out for 7 days at 37°C and concentrations of *e.g.* 100 µg/ml or 250 µg/ml in an incubator. Under these conditions, it is preferred that the antigen-
35 binding molecules of the invention show a dimer percentage that is $\leq 5\%$, more preferably $\leq 4\%$, even

more preferably $\leq 3\%$, even more preferably $\leq 2.5\%$, even more preferably $\leq 2\%$, even more preferably $\leq 1.5\%$, and most preferably $\leq 1\%$ or $\leq 0.5\%$ or even 0% .

[240] It is also preferred that the bispecific antigen-binding molecules of the present invention present with very low dimer conversion after a number of freeze/thaw cycles. For example, the antigen-binding molecule monomer is adjusted to a concentration of $250 \mu\text{g/ml}$ *e.g.* in generic formulation buffer and subjected to three freeze/thaw cycles (freezing at -80°C for 30 min followed by thawing for 30 min at room temperature), followed by high performance SEC to determine the percentage of initially monomeric antigen-binding molecule, which had been converted into dimeric antigen-binding molecule. Preferably the dimer percentages of the bispecific antigen-binding molecules are $\leq 5\%$, more preferably $\leq 4\%$, even more preferably $\leq 3\%$, even more preferably $\leq 2.5\%$, even more preferably $\leq 2\%$, even more preferably $\leq 1.5\%$, and most preferably $\leq 1\%$ or even $\leq 0.5\%$, for example after three freeze/thaw cycles.

[241] The bispecific antigen-binding molecules of the present invention preferably show a favorable thermostability with aggregation temperatures $\geq 45^\circ\text{C}$ or $\geq 50^\circ\text{C}$, more preferably $\geq 52^\circ\text{C}$ or $\geq 54^\circ\text{C}$, even more preferably $\geq 56^\circ\text{C}$ or $\geq 57^\circ\text{C}$, and most preferably $\geq 58^\circ\text{C}$ or $\geq 59^\circ\text{C}$. The thermostability parameter can be determined in terms of antibody aggregation temperature as follows: Antibody solution at a concentration $250 \mu\text{g/ml}$ is transferred into a single use cuvette and placed in a Dynamic Light Scattering (DLS) device. The sample is heated from 40°C to 70°C at a heating rate of $0.5^\circ\text{C}/\text{min}$ with constant acquisition of the measured radius. Increase of radius indicating melting of the protein and aggregation is used to calculate the aggregation temperature of the antibody.

[242] Alternatively, temperature melting curves can be determined by Differential Scanning Calorimetry (DSC) to determine intrinsic biophysical protein stabilities of the antigen-binding molecules. These experiments are performed using a MicroCal LLC (Northampton, MA, U.S.A) VP-DSC device. The energy uptake of a sample containing an antigen-binding molecule is recorded from 20°C to 90°C compared to a sample containing only the formulation buffer. The antigen-binding molecules are adjusted to a final concentration of $250 \mu\text{g/ml}$ *e.g.* in SEC running buffer. For recording of the respective melting curve, the overall sample temperature is increased stepwise. At each temperature T energy uptake of the sample and the formulation buffer reference is recorded. The difference in energy uptake C_p (kcal/mole/ $^\circ\text{C}$) of the sample minus the reference is plotted against the respective temperature. The melting temperature is defined as the temperature at the first maximum of energy uptake.

[243] The CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific antigen-binding molecules of the invention are also envisaged to have a turbidity (as measured by OD340 after concentration of purified monomeric antigen-binding molecule to

2.5 mg/ml and overnight incubation) of ≤ 0.2 , preferably of ≤ 0.15 , more preferably of ≤ 0.12 , even more preferably of ≤ 0.1 , and most preferably of ≤ 0.08 .

[244] In a further embodiment the antigen-binding molecule according to the invention is stable at physiologic or slightly lower pH, i.e. about pH 7.4 to 6.0. The more tolerant the antigen-binding molecule behaves at unphysiologic pH such as about pH 6.0, the higher is the recovery of the antigen-binding molecule eluted from an ion exchange column relative to the total amount of loaded protein. Recovery of the antigen-binding molecule from an ion (e.g., cation) exchange column at about pH 6.0 is preferably $\geq 30\%$, more preferably $\geq 40\%$, more preferably $\geq 50\%$, even more preferably $\geq 60\%$, even more preferably $\geq 70\%$, even more preferably $\geq 80\%$, even more preferably $\geq 90\%$, even more preferably $\geq 95\%$, and most preferably $\geq 99\%$.

[245] It is furthermore envisaged that the bispecific antigen-binding molecules of the present invention exhibit therapeutic efficacy or anti-tumor activity. This can e.g. be assessed in a study as disclosed in the following generalized example of an advanced stage human tumor xenograft model:

[246] On day 1 of the study, 5×10^6 cells of a human target cell antigen (here: CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM) positive cancer cell line are subcutaneously injected in the right dorsal flank of female NOD/SCID mice. When the mean tumor volume reaches about 100 mm^3 , in vitro expanded human CD3 positive T cells are transplanted into the mice by injection of about 2×10^7 cells into the peritoneal cavity of the animals. Mice of vehicle control group 1 do not receive effector cells and are used as an untransplanted control for comparison with vehicle control group 2 (receiving effector cells) to monitor the impact of T cells alone on tumor growth. The treatment with a bispecific antigen-binding molecule starts when the mean tumor volume reaches about 200 mm^3 . The mean tumor size of each treatment group on the day of treatment start should not be statistically different from any other group (analysis of variance). Mice are treated with 0.5 mg/kg/day of a CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM and CD3 bispecific antigen-binding molecule by intravenous bolus injection for about 15 to 20 days. Tumors are measured by caliper during the study and progress evaluated by intergroup comparison of tumor volumes (TV). The tumor growth inhibition T/C [%] is determined by calculating TV as $T/C\% = 100 \times (\text{median TV of analyzed group}) / (\text{median TV of control group 2})$.

[247] The skilled person knows how to modify or adapt certain parameters of this study, such as the number of injected tumor cells, the site of injection, the number of transplanted human T cells, the amount of bispecific antigen-binding molecules to be administered, and the timelines, while still arriving at a meaningful and reproducible result. Preferably, the tumor growth inhibition T/C [%] is ≤ 70 or ≤ 60 , more preferably ≤ 50 or ≤ 40 , even more preferably ≤ 30 or ≤ 20 and most preferably ≤ 10 or ≤ 5 or even ≤ 2.5 . Tumor growth inhibition is preferably close to 100%.

[248] In a preferred embodiment of the antigen-binding molecule of the invention the antigen-binding molecule is a single chain antigen-binding molecule.

[249] Also in a preferred embodiment of the antigen-binding molecule of the invention said spacer comprises in an amino to carboxyl order:

5 hinge-CH2-CH3-linker-hinge-CH2-CH3.

[250] In one embodiment of the invention each of said polypeptide monomers of the spacer has an amino acid sequence that is at least 90% identical to a sequence selected from the group consisting of: SEQ ID NO: 17-24. In a preferred embodiment of the invention each of said polypeptide monomers has an amino acid sequence selected from SEQ ID NO: 17-24.

10 [251] Also in one embodiment of the invention the CH2 domain of one or preferably each (both) polypeptide monomers of the spacer comprises an intra domain cysteine disulfide bridge. As known in the art the term "cysteine disulfide bridge" refers to a functional group with the general structure $R-S-S-R$. The linkage is also called an SS-bond or a disulfide bridge and is derived by the coupling of two thiol groups of cysteine residues. It is particularly preferred for the antigen-binding molecule of the invention that the cysteines forming the cysteine disulfide bridge in the mature antigen-binding molecule are introduced into the amino acid sequence of the CH2 domain corresponding to 309 and 321 (Kabat numbering).

15 [252] In one embodiment of the invention a glycosylation site in Kabat position 314 of the CH2 domain is removed. It is preferred that this removal of the glycosylation site is achieved by a N314X substitution, wherein X is any amino acid excluding Q. Said substitution is preferably a N314G. In a more preferred embodiment, said CH2 domain additionally comprises the following substitutions (position according to Kabat) V321C and R309C (these substitutions introduce the intra domain cysteine disulfide bridge at Kabat positions 309 and 321).

20 [253] It is assumed that the preferred features of the antigen-binding molecule of the invention compared e.g. to the bispecific heteroFc antigen-binding molecule known in the art may be inter alia related to the introduction of the above described modifications in the CH2 domain. Thus, it is preferred for the construct of the invention that the CH2 domains in the spacer of the antigen-binding molecule of the invention comprise the intra domain cysteine disulfide bridge at Kabat positions 309 and 321 and/or the glycosylation site at Kabat position 314 is removed, preferably by a N314G substitution.

25 [254] In a further preferred embodiment of the invention the CH2 domains in the spacer of the antigen-binding molecule of the invention comprise the intra domain cysteine disulfide bridge at Kabat positions 309 and 321 and the glycosylation site at Kabat position 314 is removed by a N314G substitution. Most preferably, the polypeptide monomer of the spacer of the antigen-binding molecule

of the invention has an amino acid sequence selected from the group consisting of SEQ ID NO: 17 and 18.

[255] In one embodiment the invention provides an antigen-binding molecule, wherein:

- 5 (i) the first domain comprises two antibody variable domains and the second domain comprises two antibody variable domains;
- (ii) the first domain comprises one antibody variable domain and the second domain comprises two antibody variable domains;
- (iii) the first domain comprises two antibody variable domains and the second domain comprises one antibody variable domain; or
- 10 (iv) the first domain comprises one antibody variable domain and the second domain comprises one antibody variable domain.

[256] Accordingly, the first and the second domain may be binding domains comprising each two antibody variable domains such as a VH and a VL domain. Examples for such binding domains comprising two antibody variable domains where described herein above and comprise e.g. Fv
15 fragments, scFv fragments or Fab fragments described herein above. Alternatively, either one or both of those binding domains may comprise only a single variable domain. Examples for such single domain binding domains where described herein above and comprise e.g. nanobodies or single variable domain antibodies comprising merely one variable domain, which may be VHH, VH or VL, that specifically bind an antigen or epitope independently of other V regions or domains.

20 **[257]** In a preferred embodiment of the antigen-binding molecule of the invention second and third binding domain are fused to the spacer via a peptide linker. Preferred peptide linker have been described herein above and are characterized by the amino acid sequence Gly-Gly-Gly-Gly-Ser, i.e. Gly₄Ser (SEQ ID NO: 7), or polymers thereof, i.e. (Gly₄Ser)_x, where x is an integer of 1 or greater (e.g. 2 or 3). A particularly preferred linker for the fusion of the first and second domain to the spacer
25 is depicted in SEQ ID NO: 7.

[258] The antigen-binding molecule of the present invention comprises a first domain which binds to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM, preferably to the extracellular domain(s) (ECD) of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM. It is understood that the term “binding to the extracellular domain of CS1, BCMA, CD20,
30 CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM”, in the context of the present invention, implies that the binding domain binds to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM expressed on the surface of a target cell. The first domain according to the invention hence preferably binds to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM when it is expressed by naturally expressing cells or cell lines, and/or by cells or cell lines
35 transformed or (stably / transiently) transfected with CS1, BCMA, CD20, CD22, FLT3, CD123,

CLL1, CHD3, MSLN, or EpCAM. In a preferred embodiment the first binding domain also binds to CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM when CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM is used as a "target" or "ligand" molecule in an *in vitro* binding assay such as BIAcore or Scatchard. The "target cell" can be any prokaryotic or eukaryotic cell expressing CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM on its surface; preferably the target cell is a cell that is part of the human or animal body, such as a specific CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM expressing cancer or tumor cell.

[259] Preferably, the first binding domain binds to human CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM / CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM ECD. In a further preferred embodiment, it binds to macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM / CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM ECD. According to the most preferred embodiment, it binds to both the human and the macaque CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM / CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM ECD. The "CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM extracellular domain" or "CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM ECD" refers to the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM region or sequence which is essentially free of transmembrane and cytoplasmic domains of CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM. It will be understood by the skilled artisan that the transmembrane domain identified for the CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM polypeptide of the present invention is identified pursuant to criteria routinely employed in the art for identifying that type of hydrophobic domain. The exact boundaries of a transmembrane domain may vary but most likely by no more than about 5 amino acids at either end of the domain specifically mentioned herein.

[260] Preferred binding domains which bind to CD3 are disclosed in WO 2010/037836, and WO 2011/121110. Any binding domain for CD3 described in these applications may be used in the context of the present invention.

[261] The invention further provides a polynucleotide / nucleic acid molecule encoding an antigen-binding molecule of the invention. A polynucleotide is a biopolymer composed of 13 or more nucleotide monomers covalently bonded in a chain. DNA (such as cDNA) and RNA (such as mRNA) are examples of polynucleotides with distinct biological function. Nucleotides are organic molecules that serve as the monomers or subunits of nucleic acid molecules like DNA or RNA. The nucleic acid molecule or polynucleotide can be double stranded and single stranded, linear and circular. It is preferably comprised in a vector which is preferably comprised in a host cell. Said host cell is, e.g. after transformation or transfection with the vector or the polynucleotide of the invention, capable of

expressing the antigen-binding molecule. For that purpose the polynucleotide or nucleic acid molecule is operatively linked with control sequences.

[262] The genetic code is the set of rules by which information encoded within genetic material (nucleic acids) is translated into proteins. Biological decoding in living cells is accomplished by the ribosome which links amino acids in an order specified by mRNA, using tRNA molecules to carry amino acids and to read the mRNA three nucleotides at a time. The code defines how sequences of these nucleotide triplets, called codons, specify which amino acid will be added next during protein synthesis. With some exceptions, a three-nucleotide codon in a nucleic acid sequence specifies a single amino acid. Because the vast majority of genes are encoded with exactly the same code, this particular code is often referred to as the canonical or standard genetic code. While the genetic code determines the protein sequence for a given coding region, other genomic regions can influence when and where these proteins are produced.

[263] Furthermore, the invention provides a vector comprising a polynucleotide / nucleic acid molecule of the invention. A vector is a nucleic acid molecule used as a vehicle to transfer (foreign) genetic material into a cell. The term “vector” encompasses – but is not restricted to – plasmids, viruses, cosmids and artificial chromosomes. In general, engineered vectors comprise an origin of replication, a multicloning site and a selectable marker. The vector itself is generally a nucleotide sequence, commonly a DNA sequence that comprises an insert (transgene) and a larger sequence that serves as the “backbone” of the vector. Modern vectors may encompass additional features besides the transgene insert and a backbone: promoter, genetic marker, antibiotic resistance, reporter gene, targeting sequence, protein purification tag. Vectors called expression vectors (expression constructs) specifically are for the expression of the transgene in the target cell, and generally have control sequences.

[264] The term “control sequences” refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

[265] A nucleic acid is “operably linked” when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, “operably linked” means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation

at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice.

[266] “Transfection” is the process of deliberately introducing nucleic acid molecules or polynucleotides (including vectors) into target cells. The term is mostly used for non-viral methods in eukaryotic cells. Transduction is often used to describe virus-mediated transfer of nucleic acid molecules or polynucleotides. Transfection of animal cells typically involves opening transient pores or “holes” in the cell membrane, to allow the uptake of material. Transfection can be carried out using calcium phosphate, by electroporation, by cell squeezing or by mixing a cationic lipid with the material to produce liposomes, which fuse with the cell membrane and deposit their cargo inside.

[267] The term “transformation” is used to describe non-viral transfer of nucleic acid molecules or polynucleotides (including vectors) into bacteria, and also into non-animal eukaryotic cells, including plant cells. Transformation is hence the genetic alteration of a bacterial or non-animal eukaryotic cell resulting from the direct uptake through the cell membrane(s) from its surroundings and subsequent incorporation of exogenous genetic material (nucleic acid molecules). Transformation can be effected by artificial means. For transformation to happen, cells or bacteria must be in a state of competence, which may occur as a time-limited response to environmental conditions such as starvation and cell density.

[268] Moreover, the invention provides a host cell transformed or transfected with the polynucleotide / nucleic acid molecule or with the vector of the invention. As used herein, the terms “host cell” or “recipient cell” are intended to include any individual cell or cell culture that can be or has/have been recipients of vectors, exogenous nucleic acid molecules, and polynucleotides encoding the antigen-binding molecule of the present invention; and/or recipients of the antigen-binding molecule itself. The introduction of the respective material into the cell is carried out by way of transformation, transfection and the like. The term “host cell” is also intended to include progeny or potential progeny of a single cell. Because certain modifications may occur in succeeding generations due to either natural, accidental, or deliberate mutation or due to environmental influences, such progeny may not, in fact, be completely identical (in morphology or in genomic or total DNA complement) to the parent cell, but is still included within the scope of the term as used herein. Suitable host cells include prokaryotic or eukaryotic cells, and also include but are not limited to bacteria, yeast cells, fungi cells, plant cells, and animal cells such as insect cells and mammalian cells, e.g., murine, rat, macaque or human.

[269] The antigen-binding molecule of the invention can be produced in bacteria. After expression, the antigen-binding molecule of the invention is isolated from the *E. coli* cell paste in a soluble fraction and can be purified through, e.g., affinity chromatography and/or size exclusion. Final

purification can be carried out similar to the process for purifying antibody expressed *e.g.*, in CHO cells.

[270] In addition to prokaryotes, eukaryotic microbes such as filamentous fungi or yeast are suitable cloning or expression hosts for the antigen-binding molecule of the invention. *Saccharomyces cerevisiae*, or common baker's yeast, is the most commonly used among lower eukaryotic host microorganisms. However, a number of other genera, species, and strains are commonly available and useful herein, such as *Schizosaccharomyces pombe*, Kluyveromyces hosts such as *K. lactis*, *K. fragilis* (ATCC 12424), *K. bulgaricus* (ATCC 16045), *K. wickerhamii* (ATCC 24178), *K. waltii* (ATCC 56500), *K. drosophilae* (ATCC 36906), *K. thermotolerans*, and *K. marxianus*; yarrowia (EP 402 226); *Pichia pastoris* (EP 183 070); *Candida*; *Trichoderma reesia* (EP 244 234); *Neurospora crassa*; Schwanniomyces such as *Schwanniomyces occidentalis*; and filamentous fungi such as *Neurospora*, *Penicillium*, *Tolypocladium*, and *Aspergillus* hosts such as *A. nidulans* and *A. niger*.

[271] Suitable host cells for the expression of glycosylated antigen-binding molecule of the invention are derived from multicellular organisms. Examples of invertebrate cells include plant and insect cells. Numerous baculoviral strains and variants and corresponding permissive insect host cells from hosts such as *Spodoptera frugiperda* (caterpillar), *Aedes aegypti* (mosquito), *Aedes albopictus* (mosquito), *Drosophila melanogaster* (fruit fly), and *Bombyx mori* have been identified. A variety of viral strains for transfection are publicly available, *e.g.*, the L-1 variant of *Autographa californica* NPV and the Bm-5 strain of *Bombyx mori* NPV, and such viruses may be used as the virus herein according to the present invention, particularly for transfection of *Spodoptera frugiperda* cells.

[272] Plant cell cultures of cotton, corn, potato, soybean, petunia, tomato, Arabidopsis and tobacco can also be used as hosts. Cloning and expression vectors useful in the production of proteins in plant cell culture are known to those of skill in the art. See *e.g.* Hiatt *et al.*, *Nature* (1989) 342: 76-78, Owen *et al.* (1992) *Bio/Technology* 10: 790-794, Artsaenko *et al.* (1995) *The Plant J* 8: 745-750, and Fecker *et al.* (1996) *Plant Mol Biol* 32: 979-986.

[273] However, interest has been greatest in vertebrate cells, and propagation of vertebrate cells in culture (tissue culture) has become a routine procedure. Examples of useful mammalian host cell lines are monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 or 293 cells subcloned for growth in suspension culture, Graham *et al.*, *J. Gen Virol.* 36 : 59 (1977)); baby hamster kidney cells (BHK, ATCC CCL 10); Chinese hamster ovary cells/-DHFR (CHO, Urlaub *et al.*, *Proc. Natl. Acad. Sci. USA* 77: 4216 (1980)); mouse sertoli cells (TM4, Mather, *Biol. Reprod.* 23: 243-251 (1980)); monkey kidney cells (CV1 ATCC CCL 70); African green monkey kidney cells (VERO-76, ATCC CRL1587); human cervical carcinoma cells (HELA, ATCC CCL 2); canine kidney cells (MDCK, ATCC CCL 34); buffalo rat liver cells (BRL 3A, ATCC CRL 1442); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2,1413 8065); mouse

mammary tumor (MMT 060562, ATCC CCL5 1); TRI cells (Mather *et al.*, Annals N. Y Acad. Sci. (1982) 383: 44-68); MRC 5 cells; FS4 cells; and a human hepatoma line (Hep G2).

[274] In a further embodiment the invention provides a process for the production of an antigen-binding molecule of the invention, said process comprising culturing a host cell of the invention under conditions allowing the expression of the antigen-binding molecule of the invention and recovering the produced antigen-binding molecule from the culture.

[275] As used herein, the term “culturing” refers to the *in vitro* maintenance, differentiation, growth, proliferation and/or propagation of cells under suitable conditions in a medium. The term “expression” includes any step involved in the production of an antigen-binding molecule of the invention including, but not limited to, transcription, post-transcriptional modification, translation, post-translational modification, and secretion.

[276] When using recombinant techniques, the antigen-binding molecule can be produced intracellularly, in the periplasmic space, or directly secreted into the medium. If the antigen-binding molecule is produced intracellularly, as a first step, the particulate debris, either host cells or lysed fragments, are removed, for example, by centrifugation or ultrafiltration. Carter *et al.*, Bio/Technology 10: 163-167 (1992) describe a procedure for isolating antibodies which are secreted to the periplasmic space of *E. coli*. Briefly, cell paste is thawed in the presence of sodium acetate (pH 3.5), EDTA, and phenylmethylsulfonylfluoride (PMSF) over about 30 min. Cell debris can be removed by centrifugation. Where the antibody is secreted into the medium, supernatants from such expression systems are generally first concentrated using a commercially available protein concentration filter, for example, an Amicon or Millipore Pellicon ultrafiltration unit. A protease inhibitor such as PMSF may be included in any of the foregoing steps to inhibit proteolysis and antibiotics may be included to prevent the growth of adventitious contaminants.

[277] The antigen-binding molecule of the invention prepared from the host cells can be recovered or purified using, for example, hydroxylapatite chromatography, gel electrophoresis, dialysis, and affinity chromatography. Other techniques for protein purification such as fractionation on an ion-exchange column, ethanol precipitation, Reverse Phase HPLC, chromatography on silica, chromatography on heparin SEPHAROSE™, chromatography on an anion or cation exchange resin (such as a polyaspartic acid column), chromato-focusing, SDS-PAGE, and ammonium sulfate precipitation are also available depending on the antibody to be recovered. Where the antigen-binding molecule of the invention comprises a CH3 domain, the Bakerbond ABX resin (J.T. Baker, Phillipsburg, NJ) is useful for purification.

[278] Affinity chromatography is a preferred purification technique. The matrix to which the affinity ligand is attached is most often agarose, but other matrices are available. Mechanically stable matrices

such as controlled pore glass or poly (styrene-divinyl) benzene allow for faster flow rates and shorter processing times than can be achieved with agarose.

[279] Moreover, the invention provides a pharmaceutical composition comprising an antigen-binding molecule of the invention or an antigen-binding molecule produced according to the process of the invention. It is preferred for the pharmaceutical composition of the invention that the homogeneity of the antigen-binding molecule is $\geq 80\%$, more preferably $\geq 81\%$, $\geq 82\%$, $\geq 83\%$, $\geq 84\%$, or $\geq 85\%$, further preferably $\geq 86\%$, $\geq 87\%$, $\geq 88\%$, $\geq 89\%$, or $\geq 90\%$, still further preferably, $\geq 91\%$, $\geq 92\%$, $\geq 93\%$, $\geq 94\%$, or $\geq 95\%$ and most preferably $\geq 96\%$, $\geq 97\%$, $\geq 98\%$ or $\geq 99\%$.

[280] As used herein, the term "pharmaceutical composition" relates to a composition which is suitable for administration to a patient, preferably a human patient. The particularly preferred pharmaceutical composition of this invention comprises one or a plurality of the antigen-binding molecule(s) of the invention, preferably in a therapeutically effective amount. Preferably, the pharmaceutical composition further comprises suitable formulations of one or more (pharmaceutically effective) carriers, stabilizers, excipients, diluents, solubilizers, surfactants, emulsifiers, preservatives and/or adjuvants. Acceptable constituents of the composition are preferably nontoxic to recipients at the dosages and concentrations employed. Pharmaceutical compositions of the invention include, but are not limited to, liquid, frozen, and lyophilized compositions.

[281] The inventive compositions may comprise a pharmaceutically acceptable carrier. In general, as used herein, "pharmaceutically acceptable carrier" means any and all aqueous and non-aqueous solutions, sterile solutions, solvents, buffers, e.g. phosphate buffered saline (PBS) solutions, water, suspensions, emulsions, such as oil/water emulsions, various types of wetting agents, liposomes, dispersion media and coatings, which are compatible with pharmaceutical administration, in particular with parenteral administration. The use of such media and agents in pharmaceutical compositions is well known in the art, and the compositions comprising such carriers can be formulated by well-known conventional methods.

[282] Certain embodiments provide pharmaceutical compositions comprising the antigen-binding molecule of the invention and further one or more excipients such as those illustratively described in this section and elsewhere herein. Excipients can be used in the invention in this regard for a wide variety of purposes, such as adjusting physical, chemical, or biological properties of formulations, such as adjustment of viscosity, and or processes of the invention to improve effectiveness and or to stabilize such formulations and processes against degradation and spoilage due to, for instance, stresses that occur during manufacturing, shipping, storage, pre-use preparation, administration, and thereafter.

[283] In certain embodiments, the pharmaceutical composition may contain formulation materials for the purpose of modifying, maintaining or preserving, e.g., the pH, osmolarity, viscosity, clarity, color, isotonicity, odor, sterility, stability, rate of dissolution or release, adsorption or penetration of the composition (see, REMINGTON'S PHARMACEUTICAL SCIENCES, 18th Edition, (A.R. Genrmo, ed.), 1990, Mack Publishing Company). In such embodiments, suitable formulation materials may include, but are not limited to:

- amino acids such as glycine, alanine, glutamine, asparagine, threonine, proline, 2-phenylalanine, including charged amino acids, preferably lysine, lysine acetate, arginine, glutamate and/or histidine
- 10 • antimicrobials such as antibacterial and antifungal agents
- antioxidants such as ascorbic acid, methionine, sodium sulfite or sodium hydrogen-sulfite;
- buffers, buffer systems and buffering agents which are used to maintain the composition at physiological pH or at a slightly lower pH, preferably a lower pH of 4.0 to 6.5; examples of buffers are borate, bicarbonate, Tris-HCl, citrates, phosphates or other organic acids, succinate, phosphate, and histidine; for example Tris buffer of about pH 7.0-8.5;
- 15 • non-aqueous solvents such as propylene glycol, polyethylene glycol, vegetable oils such as olive oil, and injectable organic esters such as ethyl oleate;
- aqueous carriers including water, alcoholic/aqueous solutions, emulsions or suspensions, including saline and buffered media;
- 20 • biodegradable polymers such as polyesters;
- bulking agents such as mannitol or glycine;
- chelating agents such as ethylenediamine tetraacetic acid (EDTA);
- isotonic and absorption delaying agents;
- complexing agents such as caffeine, polyvinylpyrrolidone, beta-cyclodextrin or hydroxypropyl-beta-cyclodextrin)
- 25 • fillers;
- monosaccharides; disaccharides; and other carbohydrates (such as glucose, mannose or dextrans); carbohydrates may be non-reducing sugars, preferably trehalose, sucrose, octasulfate, sorbitol or xylitol;
- 30 • (low molecular weight) proteins, polypeptides or proteinaceous carriers such as human or bovine serum albumin, gelatin or immunoglobulins, preferably of human origin;
- coloring and flavouring agents;
- sulfur containing reducing agents, such as glutathione, thiocetic acid, sodium thioglycolate, thioglycerol, [alpha]-monothioglycerol, and sodium thio sulfate
- 35 • diluting agents;
- emulsifying agents;
- hydrophilic polymers such as polyvinylpyrrolidone)

- salt-forming counter-ions such as sodium;
- preservatives such as antimicrobials, anti-oxidants, chelating agents, inert gases and the like; examples are: benzalkonium chloride, benzoic acid, salicylic acid, thimerosal, phenethyl alcohol, methylparaben, propylparaben, chlorhexidine, sorbic acid or hydrogen peroxide);
- 5 • metal complexes such as Zn-protein complexes;
- solvents and co-solvents (such as glycerin, propylene glycol or polyethylene glycol);
- sugars and sugar alcohols, such as trehalose, sucrose, octasulfate, mannitol, sorbitol or xylitol stachyose, mannose, sorbose, xylose, ribose, myoinisitol, galactose, lactitol, ribitol, myoinisitol, galactitol, glycerol, cyclitols (e.g., inositol), polyethylene glycol; and polyhydric sugar alcohols;
- 10 • suspending agents;
- surfactants or wetting agents such as pluronics, PEG, sorbitan esters, polysorbates such as polysorbate 20, polysorbate, triton, tromethamine, lecithin, cholesterol, tyloxapal; surfactants may be detergents, preferably with a molecular weight of >1.2 KD and/or a polyether, preferably with a molecular weight of >3 KD; non-limiting examples for preferred detergents are Tween 20,
- 15 Tween 40, Tween 60, Tween 80 and Tween 85; non-limiting examples for preferred polyethers are PEG 3000, PEG 3350, PEG 4000 and PEG 5000;
- stability enhancing agents such as sucrose or sorbitol;
- tonicity enhancing agents such as alkali metal halides, preferably sodium or potassium chloride, mannitol sorbitol;
- 20 • parenteral delivery vehicles including sodium chloride solution, Ringer's dextrose, dextrose and sodium chloride, lactated Ringer's, or fixed oils;
- intravenous delivery vehicles including fluid and nutrient replenishers, electrolyte replenishers (such as those based on Ringer's dextrose).

[284] In the context of the present invention, a pharmaceutical composition, which is preferably a liquid composition or may be a solid composition obtained by lyophilisation or may be a reconstituted liquid composition comprises

- (a) an antigen-binding molecule comprising at least four binding domains, wherein:
- a first and a third domain binds to a target cell surface antigen and has an isoelectric point (pI) in the range of 4 to 9,5;
 - 30 • a second and a fourth domain binds to CD3; and has a pI in the range of 8 to 10, preferably 8.5 to 9.0; and
 - a spacer comprising preferably two polypeptide monomers, each comprising a hinge, a CH2 domain and a CH3 domain, wherein said two polypeptide monomers are fused to each other via a peptide linker;
- 35 (b) at least one buffer agent;
- (c) at least one saccharide; and

(d) at least one surfactant;

and wherein the pH of the pharmaceutical composition is in the range of 3.5 to 6.

5 [285] It is further envisaged in the context of the present invention that the at least one buffer agent is present at a concentration range of 5 to 200 mM, more preferably at a concentration range of 10 to 50 mM. It is envisaged in the context of the present invention that the at least one saccharide is selected from the group consisting of monosaccharide, disaccharide, cyclic polysaccharide, sugar alcohol, linear branched dextran or linear non-branched dextran. It is also envisaged in the context of the present invention that the disaccharide is selected from the group consisting of sucrose, trehalose
10 and mannitol, sorbitol, and combinations thereof. It is further envisaged in the context of the present invention that the sugar alcohol is sorbitol. It is envisaged in the context of the present invention that the at least one saccharide is present at a concentration in the range of 1 to 15% (m/V), preferably in a concentration range of 9 to 12% (m/V).

15 [286] It is also envisaged in the context of the present invention that the at least one surfactant is selected from the group consisting of polysorbate 20, polysorbate 40, polysorbate 60, polysorbate 80, poloxamer 188, pluronic F68, triton X-100, polyoxyethylen, PEG 3350, PEG 4000 and combinations thereof. It is further envisaged in the context of the present invention that the at least one surfactant is present at a concentration in the range of 0.004 to 0.5 % (m/V), preferably in the range of 0.001 to 0.01% (m/V). It is envisaged in the context of the present invention that the pH of the composition is
20 in the range of 4.0 to 5.0, preferably 4.2. It is also envisaged in the context of the present invention that the pharmaceutical composition has an osmolarity in the range of 150 to 500 mOsm. It is further envisaged in the context of the present invention that the pharmaceutical composition further comprises an excipient selected from the group consisting of, one or more polyol and one or more amino acid. It is envisaged in the context of the present invention that said one or more excipient is
25 present in the concentration range of 0.1 to 15 % (w/V).

[287] It is also envisaged in the context of the present invention that the pharmaceutical composition comprises

- 30 (a) the antigen-binding molecule as discussed above,
(b) 10 mM glutamate or acetate,
(c) 9% (m/V) sucrose or 6% (m/V) sucrose and 6% (m/V) hydroxypropyl- β -cyclodextrin,
(d) 0.01% (m/V) polysorbate 80

and wherein the pH of the liquid pharmaceutical composition is 4.2.

35 [288] It is further envisaged in the context of the present invention that the antigen-binding molecule is present in a concentration range of 0.1 to 8 mg/ml, preferably of 0.2-2.5 mg/ml, more preferably of 0.25-1.0 mg/ml.

[289] It is evident to those skilled in the art that the different constituents of the pharmaceutical composition (e.g., those listed above) can have different effects, for example, and amino acid can act as a buffer, a stabilizer and/or an antioxidant; mannitol can act as a bulking agent and/or a tonicity enhancing agent; sodium chloride can act as delivery vehicle and/or tonicity enhancing agent; etc.

5 [290] It is envisaged that the composition of the invention may comprise, in addition to the polypeptide of the invention defined herein, further biologically active agents, depending on the intended use of the composition. Such agents may be drugs acting on the gastro-intestinal system, drugs acting as cytostatica, drugs preventing hyperurikemia, drugs inhibiting immunoreactions (e.g. corticosteroids), drugs modulating the inflammatory response, drugs acting on the circulatory system
10 and/or agents such as cytokines known in the art. It is also envisaged that the antigen-binding molecule of the present invention is applied in a co-therapy, i.e., in combination with another anti-cancer medicament.

[291] In certain embodiments, optimal pharmaceutical compositions may influence the physical state, stability, rate of in vivo release and rate of in vivo clearance of the antigen-binding molecule of
15 the invention. In certain embodiments, the primary vehicle or carrier in a pharmaceutical composition may be either aqueous or non-aqueous in nature. For example, a suitable vehicle or carrier may be water for injection, physiological saline solution or artificial cerebrospinal fluid, possibly supplemented with other materials common in compositions for parenteral administration. Neutral buffered saline or saline mixed with serum albumin are further exemplary vehicles. In certain
20 embodiments, the antigen-binding molecule of the invention compositions may be prepared for storage by mixing the selected composition having the desired degree of purity with optional formulation agents (REMINGTON'S PHARMACEUTICAL SCIENCES, supra) in the form of a lyophilized cake or an aqueous solution. Further, in certain embodiments, the antigen-binding molecule of the invention may be formulated as a lyophilizate using appropriate excipients such as
25 sucrose.

[292] When parenteral administration is contemplated, the therapeutic compositions for use in this invention may be provided in the form of a pyrogen-free, parenterally acceptable aqueous solution comprising the desired antigen-binding molecule of the invention in a pharmaceutically acceptable vehicle. A particularly suitable vehicle for parenteral injection is sterile distilled water in which the
30 antigen-binding molecule of the invention is formulated as a sterile, isotonic solution, properly preserved. In certain embodiments, the preparation can involve the formulation of the desired molecule with an agent, such as injectable microspheres, bio-erodible particles, polymeric compounds (such as polylactic acid or polyglycolic acid), beads or liposomes, that may provide controlled or sustained release of the product which can be delivered via depot injection. In certain embodiments,
35 hyaluronic acid may also be used, having the effect of promoting sustained duration in the circulation.

In certain embodiments, implantable drug delivery devices may be used to introduce the desired antigen-binding molecule.

[293] Additional pharmaceutical compositions will be evident to those skilled in the art, including formulations involving the antigen-binding molecule of the invention in sustained- or controlled-
5 delivery / release formulations. Techniques for formulating a variety of other sustained- or controlled- delivery means, such as liposome carriers, bio-erodible microparticles or porous beads and depot injections, are also known to those skilled in the art. See, for example, International Patent Application No. PCT/US93/00829, which describes controlled release of porous polymeric microparticles for delivery of pharmaceutical compositions. Sustained-release preparations may include semipermeable
10 polymer matrices in the form of shaped articles, e.g., films, or microcapsules. Sustained release matrices may include polyesters, hydrogels, polylactides (as disclosed in U.S. Pat. No. 3,773,919 and European Patent Application Publication No. EP 058481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman et al., 1983, *Biopolymers* 2:547-556), poly (2-hydroxyethyl-methacrylate) (Langer et al., 1981, *J. Biomed. Mater. Res.* 15:167-277 and Langer, 1982, *Chem. Tech.* 12:98-105),
15 ethylene vinyl acetate (Langer et al., 1981, *supra*) or poly-D(-)-3-hydroxybutyric acid (European Patent Application Publication No. EP 133,988). Sustained release compositions may also include liposomes that can be prepared by any of several methods known in the art. See, e.g., Eppstein et al., 1985, *Proc. Natl. Acad. Sci. U.S.A.* 82:3688-3692; European Patent Application Publication Nos. EP 036,676; EP 088,046 and EP 143,949.

[294] The antigen-binding molecule may also be entrapped in microcapsules prepared, for example,
20 by coacervation techniques or by interfacial polymerization (for example, hydroxymethylcellulose or gelatine-microcapsules and poly (methylmethacrylate) microcapsules, respectively), in colloidal drug delivery systems (for example, liposomes, albumin microspheres, microemulsions, nanoparticles and nanocapsules), or in macroemulsions. Such techniques are disclosed in Remington's *Pharmaceutical*
25 *Sciences*, 16th edition, Oslo, A., Ed., (1980).

[295] Pharmaceutical compositions used for in vivo administration are typically provided as sterile preparations. Sterilization can be accomplished by filtration through sterile filtration membranes. When the composition is lyophilized, sterilization using this method may be conducted either prior to
30 or following lyophilization and reconstitution. Compositions for parenteral administration can be stored in lyophilized form or in a solution. Parenteral compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

[296] Another aspect of the invention includes self-buffering antigen-binding molecule of the invention formulations, which can be used as pharmaceutical compositions, as described in
35 international patent application WO 06138181A2 (PCT/US2006/022599). A variety of expositions are

available on protein stabilization and formulation materials and methods useful in this regard, such as Arakawa et al., "Solvent interactions in pharmaceutical formulations," *Pharm Res.* 8(3): 285-91 (1991); Kendrick et al., "Physical stabilization of proteins in aqueous solution" in: *RATIONAL DESIGN OF STABLE PROTEIN FORMULATIONS: THEORY AND PRACTICE*, Carpenter and Manning, eds. *Pharmaceutical Biotechnology*. 13: 61-84 (2002), and Randolph et al., "Surfactant-protein interactions", *Pharm Biotechnol.* 13: 159-75 (2002), see particularly the parts pertinent to excipients and processes of the same for self-buffering protein formulations in accordance with the current invention, especially as to protein pharmaceutical products and processes for veterinary and/or human medical uses.

10 [297] Salts may be used in accordance with certain embodiments of the invention to, for example, adjust the ionic strength and/or the isotonicity of a formulation and/or to improve the solubility and/or physical stability of a protein or other ingredient of a composition in accordance with the invention. As is well known, ions can stabilize the native state of proteins by binding to charged residues on the protein's surface and by shielding charged and polar groups in the protein and reducing the strength of
15 their electrostatic interactions, attractive, and repulsive interactions. Ions also can stabilize the denatured state of a protein by binding to, in particular, the denatured peptide linkages (--CONH) of the protein. Furthermore, ionic interaction with charged and polar groups in a protein also can reduce intermolecular electrostatic interactions and, thereby, prevent or reduce protein aggregation and insolubility.

20 [298] Ionic species differ significantly in their effects on proteins. A number of categorical rankings of ions and their effects on proteins have been developed that can be used in formulating pharmaceutical compositions in accordance with the invention. One example is the Hofmeister series, which ranks ionic and polar non-ionic solutes by their effect on the conformational stability of proteins in solution. Stabilizing solutes are referred to as "kosmotropic". Destabilizing solutes are referred to as
25 "chaotropic". Kosmotropes commonly are used at high concentrations (e.g., >1 molar ammonium sulfate) to precipitate proteins from solution ("salting-out"). Chaotropes commonly are used to denature and/or to solubilize proteins ("salting-in"). The relative effectiveness of ions to "salt-in" and "salt-out" defines their position in the Hofmeister series.

30 [299] Free amino acids can be used in the antigen-binding molecule of the invention formulations in accordance with various embodiments of the invention as bulking agents, stabilizers, and antioxidants, as well as other standard uses. Lysine, proline, serine, and alanine can be used for stabilizing proteins in a formulation. Glycine is useful in lyophilization to ensure correct cake structure and properties. Arginine may be useful to inhibit protein aggregation, in both liquid and lyophilized formulations. Methionine is useful as an antioxidant.

[300] Polyols include sugars, e.g., mannitol, sucrose, and sorbitol and polyhydric alcohols such as, for instance, glycerol and propylene glycol, and, for purposes of discussion herein, polyethylene glycol (PEG) and related substances. Polyols are kosmotropic. They are useful stabilizing agents in both liquid and lyophilized formulations to protect proteins from physical and chemical degradation processes. Polyols also are useful for adjusting the tonicity of formulations. Among polyols useful in select embodiments of the invention is mannitol, commonly used to ensure structural stability of the cake in lyophilized formulations. It ensures structural stability to the cake. It is generally used with a lyoprotectant, e.g., sucrose. Sorbitol and sucrose are among preferred agents for adjusting tonicity and as stabilizers to protect against freeze-thaw stresses during transport or the preparation of bulks during the manufacturing process. Reducing sugars (which contain free aldehyde or ketone groups), such as glucose and lactose, can glycate surface lysine and arginine residues. Therefore, they generally are not among preferred polyols for use in accordance with the invention. In addition, sugars that form such reactive species, such as sucrose, which is hydrolyzed to fructose and glucose under acidic conditions, and consequently engenders glycation, also is not among preferred polyols of the invention in this regard. PEG is useful to stabilize proteins and as a cryoprotectant and can be used in the invention in this regard.

[301] Embodiments of the antigen-binding molecule of the invention formulations further comprise surfactants. Protein molecules may be susceptible to adsorption on surfaces and to denaturation and consequent aggregation at air-liquid, solid-liquid, and liquid-liquid interfaces. These effects generally scale inversely with protein concentration. These deleterious interactions generally scale inversely with protein concentration and typically are exacerbated by physical agitation, such as that generated during the shipping and handling of a product. Surfactants routinely are used to prevent, minimize, or reduce surface adsorption. Useful surfactants in the invention in this regard include polysorbate 20, polysorbate 80, other fatty acid esters of sorbitan polyethoxylates, and poloxamer 188. Surfactants also are commonly used to control protein conformational stability. The use of surfactants in this regard is protein-specific since, any given surfactant typically will stabilize some proteins and destabilize others.

[302] Polysorbates are susceptible to oxidative degradation and often, as supplied, contain sufficient quantities of peroxides to cause oxidation of protein residue side-chains, especially methionine. Consequently, polysorbates should be used carefully, and when used, should be employed at their lowest effective concentration. In this regard, polysorbates exemplify the general rule that excipients should be used in their lowest effective concentrations.

[303] Embodiments of the antigen-binding molecule of the invention formulations further comprise one or more antioxidants. To some extent deleterious oxidation of proteins can be prevented in pharmaceutical formulations by maintaining proper levels of ambient oxygen and temperature and by avoiding exposure to light. Antioxidant excipients can be used as well to prevent oxidative

degradation of proteins. Among useful antioxidants in this regard are reducing agents, oxygen/free-radical scavengers, and chelating agents. Antioxidants for use in therapeutic protein formulations in accordance with the invention preferably are water-soluble and maintain their activity throughout the shelf life of a product. EDTA is a preferred antioxidant in accordance with the invention in this regard.

5 Antioxidants can damage proteins. For instance, reducing agents, such as glutathione in particular, can disrupt intramolecular disulfide linkages. Thus, antioxidants for use in the invention are selected to, among other things, eliminate or sufficiently reduce the possibility of themselves damaging proteins in the formulation.

[304] Formulations in accordance with the invention may include metal ions that are protein co-
10 factors and that are necessary to form protein coordination complexes, such as zinc necessary to form certain insulin suspensions. Metal ions also can inhibit some processes that degrade proteins. However, metal ions also catalyze physical and chemical processes that degrade proteins. Magnesium ions (10-120 mM) can be used to inhibit isomerization of aspartic acid to isoaspartic acid. Ca^{+2} ions (up to 100 mM) can increase the stability of human deoxyribonuclease. Mg^{+2} , Mn^{+2} , and Zn^{+2} ,
15 however, can destabilize rhDNase. Similarly, Ca^{+2} and Sr^{+2} can stabilize Factor VIII, it can be destabilized by Mg^{+2} , Mn^{+2} and Zn^{+2} , Cu^{+2} and Fe^{+2} , and its aggregation can be increased by Al^{+3} ions.

[305] Embodiments of the antigen-binding molecule of the invention formulations further comprise one or more preservatives. Preservatives are necessary when developing multi-dose parenteral formulations that involve more than one extraction from the same container. Their primary function is
20 to inhibit microbial growth and ensure product sterility throughout the shelf-life or term of use of the drug product. Commonly used preservatives include benzyl alcohol, phenol and m-cresol. Although preservatives have a long history of use with small-molecule parenterals, the development of protein formulations that includes preservatives can be challenging. Preservatives almost always have a destabilizing effect (aggregation) on proteins, and this has become a major factor in limiting their use
25 in multi-dose protein formulations. To date, most protein drugs have been formulated for single-use only. However, when multi-dose formulations are possible, they have the added advantage of enabling patient convenience, and increased marketability. A good example is that of human growth hormone (hGH) where the development of preserved formulations has led to commercialization of more convenient, multi-use injection pen presentations. At least four such pen devices containing preserved
30 formulations of hGH are currently available on the market. Norditropin (liquid, Novo Nordisk), Nutropin AQ (liquid, Genentech) & Genotropin (lyophilized--dual chamber cartridge, Pharmacia & Upjohn) contain phenol while Somatropo (Eli Lilly) is formulated with m-cresol. Several aspects need to be considered during the formulation and development of preserved dosage forms. The effective preservative concentration in the drug product must be optimized. This requires testing a given
35 preservative in the dosage form with concentration ranges that confer anti-microbial effectiveness without compromising protein stability.

[306] As may be expected, development of liquid formulations containing preservatives are more challenging than lyophilized formulations. Freeze-dried products can be lyophilized without the preservative and reconstituted with a preservative containing diluent at the time of use. This shortens the time for which a preservative is in contact with the protein, significantly minimizing the associated stability risks. With liquid formulations, preservative effectiveness and stability should be maintained over the entire product shelf-life (about 18 to 24 months). An important point to note is that preservative effectiveness should be demonstrated in the final formulation containing the active drug and all excipient components.

[307] The antigen-binding molecules disclosed herein may also be formulated as immunoliposomes. A "liposome" is a small vesicle composed of various types of lipids, phospholipids and/or surfactant which is useful for delivery of a drug to a mammal. The components of the liposome are commonly arranged in a bilayer formation, similar to the lipid arrangement of biological membranes. Liposomes containing the antigen-binding molecule are prepared by methods known in the art, such as described in Epstein et al., Proc. Natl. Acad. Sci. USA, 82: 3688 (1985); Hwang et al., Proc. Natl. Acad. Sci. USA, 77: 4030 (1980); US Pat. Nos. 4,485,045 and 4,544,545; and WO 97/38731. Liposomes with enhanced circulation time are disclosed in US Patent No. 5,013, 556. Particularly useful liposomes can be generated by the reverse phase evaporation method with a lipid composition comprising phosphatidylcholine, cholesterol and PEG-derivatized phosphatidylethanolamine (PEG-PE). Liposomes are extruded through filters of defined pore size to yield liposomes with the desired diameter. Fab' fragments of the antigen-binding molecule of the present invention can be conjugated to the liposomes as described in Martin et al. J. Biol. Chem. 257: 286-288 (1982) via a disulfide interchange reaction. A chemotherapeutic agent is optionally contained within the liposome. See Gabizon et al. J. National Cancer Inst. 81 (19) 1484 (1989).

[308] Once the pharmaceutical composition has been formulated, it may be stored in sterile vials as a solution, suspension, gel, emulsion, solid, crystal, or as a dehydrated or lyophilized powder. Such formulations may be stored either in a ready-to-use form or in a form (e.g., lyophilized) that is reconstituted prior to administration.

[309] The biological activity of the pharmaceutical composition defined herein can be determined for instance by cytotoxicity assays, as described in the following examples, in WO 99/54440 or by Schlereth et al. (Cancer Immunol. Immunother. 20 (2005), 1-12). "Efficacy" or "in vivo efficacy" as used herein refers to the response to therapy by the pharmaceutical composition of the invention, using e.g. standardized NCI response criteria. The success or in vivo efficacy of the therapy using a pharmaceutical composition of the invention refers to the effectiveness of the composition for its intended purpose, i.e. the ability of the composition to cause its desired effect, i.e. depletion of pathologic cells, e.g. tumor cells. The in vivo efficacy may be monitored by established standard methods for the respective disease entities including, but not limited to white blood cell counts,

differentials, Fluorescence Activated Cell Sorting, bone marrow aspiration. In addition, various disease specific clinical chemistry parameters and other established standard methods may be used. Furthermore, computer-aided tomography, X-ray, nuclear magnetic resonance tomography (e.g. for National Cancer Institute-criteria based response assessment [Cheson BD, Horning SJ, Coiffier B, Shipp MA, Fisher RI, Connors JM, Lister TA, Vose J, Grillo-Lopez A, Hagenbeek A, Cabanillas F, Klippensten D, Hiddemann W, Castellino R, Harris NL, Armitage JO, Carter W, Hoppe R, Canellos GP. Report of an international workshop to standardize response criteria for non-Hodgkin's lymphomas. NCI Sponsored International Working Group. J Clin Oncol. 1999 Apr;17(4):1244]), positron-emission tomography scanning, white blood cell counts, differentials, Fluorescence Activated Cell Sorting, bone marrow aspiration, lymph node biopsies/histologies, and various lymphoma specific clinical chemistry parameters (e.g. lactate dehydrogenase) and other established standard methods may be used.

[310] Another major challenge in the development of drugs such as the pharmaceutical composition of the invention is the predictable modulation of pharmacokinetic properties. To this end, a pharmacokinetic profile of the drug candidate, i.e. a profile of the pharmacokinetic parameters that affect the ability of a particular drug to treat a given condition, can be established. Pharmacokinetic parameters of the drug influencing the ability of a drug for treating a certain disease entity include, but are not limited to: half-life, volume of distribution, hepatic first-pass metabolism and the degree of blood serum binding. The efficacy of a given drug agent can be influenced by each of the parameters mentioned above. It is an envisaged characteristic of the antigen-binding molecules of the present invention provided with the specific FC modality that they comprise, for example, differences in pharmacokinetic behavior. A half-life extended targeting antigen-binding molecule according to the present invention preferably shows a surprisingly increased residence time in vivo in comparison to "canonical" non-HLE versions of said antigen-binding molecule.

[311] "Half-life" means the time where 50% of an administered drug are eliminated through biological processes, e.g. metabolism, excretion, etc. By "hepatic first-pass metabolism" is meant the propensity of a drug to be metabolized upon first contact with the liver, i.e. during its first pass through the liver. "Volume of distribution" means the degree of retention of a drug throughout the various compartments of the body, like e.g. intracellular and extracellular spaces, tissues and organs, etc. and the distribution of the drug within these compartments. "Degree of blood serum binding" means the propensity of a drug to interact with and bind to blood serum proteins, such as albumin, leading to a reduction or loss of biological activity of the drug.

[312] Pharmacokinetic parameters also include bioavailability, lag time (Tlag), Tmax, absorption rates, more onset and/or Cmax for a given amount of drug administered. "Bioavailability" means the amount of a drug in the blood compartment. "Lag time" means the time delay between the administration of the drug and its detection and measurability in blood or plasma. "Tmax" is the time

after which maximal blood concentration of the drug is reached, and “C_{max}” is the blood concentration maximally obtained with a given drug. The time to reach a blood or tissue concentration of the drug which is required for its biological effect is influenced by all parameters. Pharmacokinetic parameters of bispecific antigen-binding molecules exhibiting cross-species specificity, which may be determined in preclinical animal testing in non-chimpanzee primates as outlined above, are also set forth e.g. in the publication by Schlereth et al. (Cancer Immunol. Immunother. 20 (2005), 1-12).

[313] In a preferred aspect of the invention the pharmaceutical composition is stable for at least four weeks at about -20°C. As apparent from the appended examples the quality of an antigen-binding molecule of the invention vs. the quality of corresponding state of the art antigen-binding molecules may be tested using different systems. Those tests are understood to be in line with the “ICH Harmonised Tripartite Guideline: *Stability Testing of Biotechnological/Biological Products Q5C and Specifications: Test procedures and Acceptance Criteria for Biotech Biotechnological/Biological Products Q6B*” and, thus are elected to provide a stability-indicating profile that provides certainty that changes in the identity, purity and potency of the product are detected. It is well accepted that the term purity is a relative term. Due to the effect of glycosylation, deamidation, or other heterogeneities, the absolute purity of a biotechnological/biological product should be typically assessed by more than one method and the purity value derived is method-dependent. For the purpose of stability testing, tests for purity should focus on methods for determination of degradation products.

[314] For the assessment of the quality of a pharmaceutical composition comprising an antigen-binding molecule of the invention may be analyzed e.g. by analyzing the content of soluble aggregates in a solution (HMWS per size exclusion). It is preferred that stability for at least four weeks at about -20°C is characterized by a content of less than 1.5% HMWS, preferably by less than 1% HMWS.

[315] A preferred formulation for the antigen-binding molecule as a pharmaceutical composition may e.g. comprise the components of a formulation as described below:

25• Formulation:

potassium phosphate, L-arginine hydrochloride, trehalose dihydrate, polysorbate 80 at pH 6.0

[316] Other examples for the assessment of the stability of an antigen-binding molecule of the invention in form of a pharmaceutical composition are provided in the appended examples 4-12. In those examples embodiments of antigen-binding molecules of the invention are tested with respect to different stress conditions in different pharmaceutical formulations and the results compared with other half-life extending (HLE) formats of bispecific T cell engaging antigen-binding molecule known from the art. In general, it is envisaged that antigen-binding molecules provided with the specific FC modality according to the present invention are typically more stable over a broad range of stress conditions such as temperature and light stress, both compared to antigen-binding molecules provided

with different HLE formats and without any HLE format (e.g. “canonical” antigen-binding molecules). Said temperature stability may relate both to decreased (below room temperature including freezing) and increased (above room temperature including temperatures up to or above body temperature) temperature. As the person skilled in the art will acknowledge, such improved stability with regard to stress, which is hardly avoidable in clinical practice, makes the antigen-binding molecule safer because less degradation products will occur in clinical practice. In consequence, said increased stability means increased safety.

[317] One embodiment provides the antigen-binding molecule of the invention or the antigen-binding molecule produced according to the process of the invention for use in the prevention, treatment or amelioration of a cancer correlating with, CD20, CD22, FLT3, CLL1, CHD3, MSLN, or EpCAM expression or CD20, CD22, FLT3, , CLL1, CHD3, MSLN, or EpCAM overexpression, such as prostate cancer.

[318] The formulations described herein are useful as pharmaceutical compositions in the treatment, amelioration and/or prevention of the pathological medical condition as described herein in a patient in need thereof. The term "treatment" refers to both therapeutic treatment and prophylactic or preventative measures. Treatment includes the application or administration of the formulation to the body, an isolated tissue, or cell from a patient who has a disease/disorder, a symptom of a disease/disorder, or a predisposition toward a disease/disorder, with the purpose to cure, heal, alleviate, relieve, alter, remedy, ameliorate, improve, or affect the disease, the symptom of the disease, or the predisposition toward the disease.

[319] The term “amelioration” as used herein refers to any improvement of the disease state of a patient having a disease as specified herein below, by the administration of an antigen-binding molecule according to the invention to a subject in need thereof. Such an improvement may also be seen as a slowing or stopping of the progression of the patient’s disease. The term “prevention” as used herein means the avoidance of the occurrence or re-occurrence of a patient having a tumor or cancer or a metastatic cancer as specified herein below, by the administration of an antigen-binding molecule according to the invention to a subject in need thereof.

[320] The term “disease” refers to any condition that would benefit from treatment with the antigen-binding molecule or the pharmaceutical composition described herein. This includes chronic and acute disorders or diseases including those pathological conditions that predispose the mammal to the disease in question.

[321] A “neoplasm” is an abnormal growth of tissue, usually but not always forming a mass. When also forming a mass, it is commonly referred to as a “tumor”. Neoplasms or tumors or can be benign, potentially malignant (pre-cancerous), or malignant. Malignant neoplasms are commonly called

cancer. They usually invade and destroy the surrounding tissue and may form metastases, i.e., they spread to other parts, tissues or organs of the body. Hence, the term “metastatic cancer” encompasses metastases to other tissues or organs than the one of the original tumor. Lymphomas and leukemias are lymphoid neoplasms. For the purposes of the present invention, they are also encompassed by the terms “tumor” or “cancer”.

[322] The term “viral disease” describes diseases, which are the result of a viral infection of a subject.

[323] The term “immunological disorder” as used herein describes in line with the common definition of this term immunological disorders such as autoimmune diseases, hypersensitivities, immune deficiencies.

[324] In one embodiment the invention provides a method for the treatment or amelioration of a cancer correlating with CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM expression or CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM overexpression, comprising the step of administering to a subject in need thereof the antigen-binding molecule of the invention, or the antigen-binding molecule produced according to the process of the invention. The CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAMxCD3 bispecific single chain antibody is particularly advantageous for the therapy of cancer, preferably solid tumors, more preferably carcinomas and prostate cancer.

[325] The terms “subject in need” or those “in need of treatment” includes those already with the disorder, as well as those in which the disorder is to be prevented. The subject in need or “patient” includes human and other mammalian subjects that receive either prophylactic or therapeutic treatment.

[326] The antigen-binding molecule of the invention will generally be designed for specific routes and methods of administration, for specific dosages and frequencies of administration, for specific treatments of specific diseases, with ranges of bio-availability and persistence, among other things. The materials of the composition are preferably formulated in concentrations that are acceptable for the site of administration.

[327] Formulations and compositions thus may be designed in accordance with the invention for delivery by any suitable route of administration. In the context of the present invention, the routes of administration include, but are not limited to

- topical routes (such as epicutaneous, inhalational, nasal, ophthalmic, auricular / aural, vaginal, mucosal);
- enteral routes (such as oral, gastrointestinal, sublingual, sublabial, buccal, rectal); and

- parenteral routes (such as intravenous, intraarterial, intraosseous, intramuscular, intracerebral, intracerebroventricular, epidural, intrathecal, subcutaneous, intraperitoneal, extra-amniotic, intraarticular, intracardiac, intradermal, intralesional, intrauterine, intravesical, intravitreal, transdermal, intranasal, transmucosal, intrasynovial, intraluminal).

5 [328] The pharmaceutical compositions and the antigen-binding molecule of this invention are particularly useful for parenteral administration, e.g., subcutaneous or intravenous delivery, for example by injection such as bolus injection, or by infusion such as continuous infusion. Pharmaceutical compositions may be administered using a medical device. Examples of medical devices for administering pharmaceutical compositions are described in U.S. Patent Nos. 4,475,196;
10 4,439,196; 4,447,224; 4,447, 233; 4,486,194; 4,487,603; 4,596,556; 4,790,824; 4,941,880; 5,064,413; 5,312,335; 5,312,335; 5,383,851; and 5,399,163.

[329] In particular, the present invention provides for an uninterrupted administration of the suitable composition. As a non-limiting example, uninterrupted or substantially uninterrupted, i.e. continuous administration may be realized by a small pump system worn by the patient for metering the influx of
15 therapeutic agent into the body of the patient. The pharmaceutical composition comprising the antigen-binding molecule of the invention can be administered by using said pump systems. Such pump systems are generally known in the art, and commonly rely on periodic exchange of cartridges containing the therapeutic agent to be infused. When exchanging the cartridge in such a pump system, a temporary interruption of the otherwise uninterrupted flow of therapeutic agent into the body of the
20 patient may ensue. In such a case, the phase of administration prior to cartridge replacement and the phase of administration following cartridge replacement would still be considered within the meaning of the pharmaceutical means and methods of the invention together make up one “uninterrupted administration” of such therapeutic agent.

[330] The continuous or uninterrupted administration of the antigen-binding molecules of the
25 invention may be intravenous or subcutaneous by way of a fluid delivery device or small pump system including a fluid driving mechanism for driving fluid out of a reservoir and an actuating mechanism for actuating the driving mechanism. Pump systems for subcutaneous administration may include a needle or a cannula for penetrating the skin of a patient and delivering the suitable composition into the patient’s body. Said pump systems may be directly fixed or attached to the skin of the patient
30 independently of a vein, artery or blood vessel, thereby allowing a direct contact between the pump system and the skin of the patient. The pump system can be attached to the skin of the patient for 24 hours up to several days. The pump system may be of small size with a reservoir for small volumes. As a non-limiting example, the volume of the reservoir for the suitable pharmaceutical composition to be administered can be between 0.1 and 50 ml.

[331] The continuous administration may also be transdermal by way of a patch worn on the skin and replaced at intervals. One of skill in the art is aware of patch systems for drug delivery suitable for this purpose. It is of note that transdermal administration is especially amenable to uninterrupted administration, as exchange of a first exhausted patch can advantageously be accomplished
5 simultaneously with the placement of a new, second patch, for example on the surface of the skin immediately adjacent to the first exhausted patch and immediately prior to removal of the first exhausted patch. Issues of flow interruption or power cell failure do not arise.

[332] If the pharmaceutical composition has been lyophilized, the lyophilized material is first reconstituted in an appropriate liquid prior to administration. The lyophilized material may be
10 reconstituted in, e.g., bacteriostatic water for injection (BWFI), physiological saline, phosphate buffered saline (PBS), or the same formulation the protein had been in prior to lyophilization.

[333] The compositions of the present invention can be administered to the subject at a suitable dose which can be determined e.g. by dose escalating studies by administration of increasing doses of the antigen-binding molecule of the invention exhibiting cross-species specificity described herein to non-
15 chimpanzee primates, for instance macaques. As set forth above, the antigen-binding molecule of the invention exhibiting cross-species specificity described herein can be advantageously used in identical form in preclinical testing in non-chimpanzee primates and as drug in humans.

[334] The term "effective dose" or "effective dosage" is defined as an amount sufficient to achieve or at least partially achieve the desired effect. The term "therapeutically effective dose" is defined as
20 an amount sufficient to cure or at least partially arrest the disease and its complications in a patient already suffering from the disease. Amounts or doses effective for this use will depend on the condition to be treated (the indication), the delivered antigen-binding molecule, the therapeutic context and objectives, the severity of the disease, prior therapy, the patient's clinical history and response to the therapeutic agent, the route of administration, the size (body weight, body surface or organ size)
25 and/or condition (the age and general health) of the patient, and the general state of the patient's own immune system.

[335] A typical dosage may range from about 0.1 $\mu\text{g}/\text{kg}$ to up to about 30 mg/kg or more, depending on the factors mentioned above. In specific embodiments, the dosage may range from 1.0 $\mu\text{g}/\text{kg}$ up to about 20 mg/kg , optionally from 10 $\mu\text{g}/\text{kg}$ up to about 10 mg/kg or from 100 $\mu\text{g}/\text{kg}$ up to about
30 5 mg/kg .

[336] A therapeutic effective amount of an antigen-binding molecule of the invention preferably results in a decrease in severity of disease symptoms, an increase in frequency or duration of disease symptom-free periods or a prevention of impairment or disability due to the disease affliction. For treating diseases correlating with CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or

EpCAM expression as described herein above, a therapeutically effective amount of the antigen-binding molecule of the invention, here: an anti-CS1, BCMA, CD20, CD22, FLT3, CD123, CLL1, CHD3, MSLN, or EpCAM/anti-CD3 antigen-binding molecule, preferably inhibits cell growth or tumor growth by at least about 20%, at least about 40%, at least about 50%, at least about 60%, at least about 70%, at least about 80%, or at least about 90% relative to untreated patients. The ability of a compound to inhibit tumor growth may be evaluated in an animal model predictive of efficacy

[337] The pharmaceutical composition can be administered as a sole therapeutic or in combination with additional therapies such as anti-cancer therapies as needed, e.g. other proteinaceous and non-proteinaceous drugs. These drugs may be administered simultaneously with the composition comprising the antigen-binding molecule of the invention as defined herein or separately before or after administration of said antigen-binding molecule in timely defined intervals and doses.

[338] The term “effective and non-toxic dose” as used herein refers to a tolerable dose of an inventive antigen-binding molecule which is high enough to cause depletion of pathologic cells, tumor elimination, tumor shrinkage or stabilization of disease without or essentially without major toxic effects. Such effective and non-toxic doses may be determined e.g. by dose escalation studies described in the art and should be below the dose inducing severe adverse side events (dose limiting toxicity, DLT).

[339] The term “toxicity” as used herein refers to the toxic effects of a drug manifested in adverse events or severe adverse events. These side events may refer to a lack of tolerability of the drug in general and/or a lack of local tolerance after administration. Toxicity could also include teratogenic or carcinogenic effects caused by the drug.

[340] The term “safety”, “in vivo safety” or “tolerability” as used herein defines the administration of a drug without inducing severe adverse events directly after administration (local tolerance) and during a longer period of application of the drug. “Safety”, “in vivo safety” or “tolerability” can be evaluated e.g. at regular intervals during the treatment and follow-up period. Measurements include clinical evaluation, e.g. organ manifestations, and screening of laboratory abnormalities. Clinical evaluation may be carried out and deviations to normal findings recorded/coded according to NCI-CTC and/or MedDRA standards. Organ manifestations may include criteria such as allergy/immunology, blood/bone marrow, cardiac arrhythmia, coagulation and the like, as set forth e.g. in the Common Terminology Criteria for adverse events v3.0 (CTCAE). Laboratory parameters which may be tested include for instance hematology, clinical chemistry, coagulation profile and urine analysis and examination of other body fluids such as serum, plasma, lymphoid or spinal fluid, liquor and the like. Safety can thus be assessed e.g. by physical examination, imaging techniques (i.e. ultrasound, x-ray, CT scans, Magnetic Resonance Imaging (MRI), other measures with technical devices (i.e. electrocardiogram), vital signs, by measuring laboratory parameters and recording

adverse events. For example, adverse events in non-chimpanzee primates in the uses and methods according to the invention may be examined by histopathological and/or histochemical methods.

5 [341] The above terms are also referred to e.g. in the Preclinical safety evaluation of biotechnology-derived pharmaceuticals S6; ICH Harmonised Tripartite Guideline; ICH Steering Committee meeting on July 16, 1997.

[342] Finally, the invention provides a kit comprising an antigen-binding molecule of the invention or produced according to the process of the invention, a pharmaceutical composition of the invention, a polynucleotide of the invention, a vector of the invention and/or a host cell of the invention.

10 [343] In the context of the present invention, the term “kit” means two or more components – one of which corresponding to the antigen-binding molecule, the pharmaceutical composition, the vector or the host cell of the invention – packaged together in a container, recipient or otherwise. A kit can hence be described as a set of products and/or utensils that are sufficient to achieve a certain goal, which can be marketed as a single unit.

15 [344] The kit may comprise one or more recipients (such as vials, ampoules, containers, syringes, bottles, bags) of any appropriate shape, size and material (preferably waterproof, e.g. plastic or glass) containing the antigen-binding molecule or the pharmaceutical composition of the present invention in an appropriate dosage for administration (see above). The kit may additionally contain directions for use (e.g. in the form of a leaflet or instruction manual), means for administering the antigen-binding molecule of the present invention such as a syringe, pump, infuser or the like, means for reconstituting
20 the antigen-binding molecule of the invention and/or means for diluting the antigen-binding molecule of the invention.

[345] The invention also provides kits for a single-dose administration unit. The kit of the invention may also contain a first recipient comprising a dried / lyophilized antigen-binding molecule and a second recipient comprising an aqueous formulation. In certain embodiments of this invention, kits
25 containing single-chambered and multi-chambered pre-filled syringes (e.g., liquid syringes and lyosyringes) are provided.

30 [346] It is noted that as used herein, the singular forms “a”, “an”, and “the”, include plural references unless the context clearly indicates otherwise. Thus, for example, reference to “a reagent” includes one or more of such different reagents and reference to “the method” includes reference to equivalent steps and methods known to those of ordinary skill in the art that could be modified or substituted for the methods described herein.

[347] Unless otherwise indicated, the term "at least" preceding a series of elements is to be understood to refer to every element in the series. Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the present invention.

[348] The term "and/or" wherever used herein includes the meaning of "and", "or" and "all or any other combination of the elements connected by said term".

[349] The term "about" or "approximately" as used herein means within 20%, preferably within 10%, and more preferably within 5% of a given value or range. It includes, however, also the concrete number, e.g., about 20 includes 20.

[350] The term "less than" or "greater than" includes the concrete number. For example, less than 20 means less than or equal to. Similarly, more than or greater than means more than or equal to, or greater than or equal to, respectively.

[351] Throughout this specification and the claims which follow, unless the context requires otherwise, the word "comprise", and variations such as "comprises" and "comprising", will be understood to imply the inclusion of a stated integer or step or group of integers or steps but not the exclusion of any other integer or step or group of integer or step. When used herein the term "comprising" can be substituted with the term "containing" or "including" or sometimes when used herein with the term "having".

[352] When used herein "consisting of" excludes any element, step, or ingredient not specified in the claim element. When used herein, "consisting essentially of" does not exclude materials or steps that do not materially affect the basic and novel characteristics of the claim.

[353] In each instance herein any of the terms "comprising", "consisting essentially of" and "consisting of" may be replaced with either of the other two terms.

[354] It should be understood that this invention is not limited to the particular methodology, protocols, material, reagents, and substances, etc., described herein and as such can vary. The terminology used herein is for the purpose of describing particular embodiments only, and is not intended to limit the scope of the present invention, which is defined solely by the claims.

[355] All publications and patents cited throughout the text of this specification (including all patents, patent applications, scientific publications, manufacturer's specifications, instructions, etc.), whether supra or infra, are hereby incorporated by reference in their entirety. Nothing herein is to be construed as an admission that the invention is not entitled to antedate such disclosure by virtue of prior invention. To the extent the material incorporated by reference contradicts or is inconsistent with this specification, the specification will supersede any such material.

[356] A better understanding of the present invention and of its advantages will be obtained from the following examples, offered for illustrative purposes only. The examples are not intended to limit the scope of the present invention in any way.

EXAMPLES

5 [357] Example 1: T-cell dependent cellular cytotoxicity (TDCC) assay with unstimulated human Tcells on multichain multitargeting bispecific antigen-binding molecules to determine beneficial efficacy gap

Isolation of effector cells

Human peripheral blood mononuclear cells (PBMC) were prepared by Ficoll density gradient
10 centrifugation from enriched lymphocyte preparations (buffy coats). Buffy coats were supplied by a local blood bank and PBMC were prepared on the day after blood collection. After Ficoll density centrifugation and washes with Dulbecco's PBS (Gibco), erythrocytes were removed from PBMC via incubation with erythrocyte lysis buffer (155 mM NH₄Cl, 10 mM KHCO₃, 100 μM EDTA). Remaining lymphocytes mainly encompass B and T lymphocytes, NK cells and monocytes. PBMC
15 were kept in culture at 37°C/5% CO₂ in RPMI complete medium (RPMI1640 (Biochrom AG, #FG1215) supplemented with 10% Fetal Bovine Serum (FBS) (Bio West, #S1810), 1x non-essential amino acids (Biochrom AG, #K0293), 1 mM sodium pyruvate (Biochrom AG, #L0473) and 100 U/mL penicillin/streptomycin (Biochrom AG, #A2213)).

20 Isolation of human T-cells

For isolation of human T-cells, Pan T Cell Isolation Kit, human (Miltenyi Biotec, MACS, #130-096-535) was used to deplete non-target cells, i.e., monocytes, neutrophils, eosinophils, B cells, stem cells, dendritic cells, NK cells, granulocytes, or erythroid cells from the PBMC cell solution. Cell were isolated according to the manufacturer's protocol and stored in RPMI complete medium at 37°C/5%
25 CO₂ until needed.

Setup of luciferase-based T-cell-dependent cellular cytotoxicity (TDCC) assay and

analysis Luciferase(LUC)-positive target-cells and effector cells (i.e., Pan T-cells) were mixed at an effector to target-cell (E:T) ratio of 10:1 and incubated with serial dilutions of the corresponding multichain multitargeting bispecific T-cell engager molecules in 384-well plates. Plates were
30 incubated for 48 hours in a 5% CO₂ humidified incubator at 37°C.

Following target cell lines were used for the Luciferase-based cytotoxicity assay:

- **HCT116 WT**

Parental cell line, wildtype (WT), transfected with luciferase

- **HCT116 MSLN KO**

35 Parental cell line HCT 116 LUC, in which MSLN gene was knocked out (KO)

- **HCT116 CDH3 KO**

Parental cell line HCT 116 LUC, in which CDH3 gene was knocked out (KO)

- **GSU WT**

Parental cell line, wildtype (WT), transfected with luciferase

- **GSU MSLN KO**

5 Parental cell line GSU LUC, in which MSLN gene was knocked out (KO)

- **GSU CDH3 KO**

Parental cell line GSU LUC, in which CDH3 gene was knocked out (KO)

10 Upon measurement of cytotoxicity, luciferase substrate (Steady-Glo® Reagent, Promega) was added to the 384-well plates. Living cells are lysed, thereby internal luciferase is released into the supernatant creating a luminescence signal through interaction with the substrate. Samples were measured with a SPARK microplate reader (TECAN) and analyzed by Spark Control Magellan software (TECAN).

Percentage of cytotoxicity was calculated as follows:

$$\text{Cytotoxicity [\%]} = \left(1 - \frac{\text{RLU}_{\text{Sample}}}{\text{RLU}_{\text{Negative Control}}}\right) \times 100$$

15

RLU = relative light units

Negative-Control = cells without multitargeting bispecific T-cell engager polypeptides

Using GraphPad Prism 8.4.3 software (Graph Pad Software, San Diego), the percentage of cytotoxicity was plotted against the corresponding molecule concentration. Sigmoidal dose response curves were analyzed with the four parametric logistic regression models with variable slope and EC50 values were calculated.

25

Table 4: EC50 values and selectivity gaps of parental HCT116 WT cells versus target knockout HCT116 cells. b.c.t. : below calculation threshold

	EC50 HCT116 KO CDH3 [pM]	-fold selectivity gap	EC50 HCT116 WT [pM]	-fold selectivity gap	EC50 HCT116 KO MSLN [pM]
CDH3 T-cell engager molecule 1	b.c.t.	-	0.53	1	0.65
MSLN T-cell engager molecule 1	0.86	1	0.76	-	b.c.t.
MSLN-CDH3 T-cell engager molecule 1	1.76	600	0.003	367	1.08
MSLN-CDH3 T-cell engager molecule 2	3.48	266	0.013	135	1.77

MSLN-CDH3 T-cell engager molecule 3	3.19	623	0.005	74	0.38
MSLN-CDH3 T-cell engager molecule 4	3.50	130	0.027	23	0.62
MSLN-CDH3 T-cell engager molecule 5	8.07	207	0.039	58	2.25
MSLN-CDH3 T-cell engager molecule 6	0.58	279	0.002	132	0.27
MSLN-CDH3 T-cell engager molecule 7	25.5	635	0.040	37	1.50

Results: CDH3 T-cell engager molecule 1 and MSLN T-cell engager molecule 1, that only target either CDH3 or MSLN, demonstrate comparable activity on single positive knockout cells vs. double positive cells HCT116 WT cells. MSLN-CDH3 T-cell engager molecules 1, 2, 3, 4, 5, 6 and 7 all show highly increased activity on double positive HCT116 WT cells compared to target knockout HCT116 cells. The EC50 selectivity gaps between double positive WT cells and CDH3 knockout cells varies within the T-cell engager molecules between 130- and 635-fold, the EC50 selectivity gaps between double positive WT cells and MSLN knockout cells varies between 23- and 367-fold. Table 5: EC50 values and selectivity gaps of parental GSU WT cells versus target knockout GSU cells. b.c.t. : below calculation threshold

	EC50 GSU KO CDH3 [pM]	-fold selectivity gap	EC50 GSU WT [pM]	-fold selectivity gap	EC50 GSU KO MSLN [pM]
CDH3 T-cell engager molecule 1	b.c.t.	-	9.9	2	23.8
MSLN T-cell engager molecule 1	0.31	1	0.41	-	b.c.t.
MSLN-CDH3 T-cell engager molecule 1	0.47	37	0.013	8238	103.2
MSLN-CDH3 T-cell engager molecule 2	2.50	64	0.039	8913	349.4
MSLN-CDH3 T-cell engager molecule 3	7.50	103	0.073	761	55.3
MSLN-CDH3 T-cell engager molecule 4	2.89	29	0.098	374	36.8
MSLN-CDH3 T-cell engager molecule 5	10.5	82	0.128	1369	174.8
MSLN-CDH3 T-cell engager molecule 6	0.89	37	0.024	483	11.6
MSLN-CDH3 T-cell engager molecule 7	54.6	209	0.261	363	95.0

Results: CDH3 T-cell engager molecule 1 and MSLN T-cell engager molecule 1, that only target either CDH3 or MSLN, demonstrate comparable activity on single positive knockout cells vs. double positive cells GSU WT cells (1- to 2-fold selectivity gap). MSLN-CDH3 T-cell engager molecules 1, 2, 3, 4, 5, 6 and 7 all show increased activity on double positive GSU WT cells compared to target knockout GSU cells. The EC50 selectivity gaps between double positive and CDH3 knockout cells varies within the T-cell engager molecules between 29- and 209-fold, the EC50 selectivity gaps between double positive and MSLN knockout cells varies between 363- and 8238- fold.

MSLN-CDH3 T-cell engager molecule 1 is a single chain multitargeting bispecific antigen-binding molecule, more specifically a T-cell engager molecule), with one bispecific entity (target binding domain and CD3 binding domain) at the N-terminus of a spacer and one bispecific entity at the C-terminus of the polypeptide, separated by a single chain Fc-domain as spacer. In MSLN-CDH3 T-cell engager molecule 2, the bispecific entities are separated by a heterodimer domain (heteroFc) that connects two multitargeting bispecific T-cell engager polypeptides and spaces apart the first and the second bispecific entity. MSLN-CDH3 T-cell engager molecules 3-7 are multichain multitargeting bispecific T-cell engager polypeptides (MMBiTEP), with one target binding domain and one CD3 binding domain forming a bispecific entity at the N-terminus of the polypeptide chains and one target binding domain and one CD3 binding domain forming another bispecific entity at the C-terminus of the polypeptide chains, separated by a hetero Fc domain spacer. The target and CD3 binding domains and their arrangements vary between constructs 3-7, but they all share the separation of the bispecific entities between N- and C-terminus of the polypeptides of the heteroFc spacer.

Legend to tables/figures	Seq ID	Binder detailed description
CDH3 T-cell engager molecule 1 (control)	Seq ID 1284	CH3-G8A_6-B12 x I2Cx scFc
MSLN T-cell engager molecule 1 (control)	Seq ID 1285	MS 5-F11 x I2C x scFc
MSLN-CDH3 T-cell engager molecule 1 (scFc single chain)	Seq ID 1272	antiCDH3_01 scFv x I2L scFv x scFc x antiMSLN_01 scFv x I2L scFv
MSLN-CDH3 T-cell engager molecule 2 (heteroFc multichain)	Seq ID 1259 + 1251	antiCDH3_01 scFv x I2L scFv x heFc(A)*heFc(B) x antiMSLN_01 scFv x I2L
MSLN-CDH3 T-cell engager molecule 3 (example to Fig. 2A)	Seq ID 1247 + 1248	antiCDH3_01 scFv x heFc(A) x antiMSLN_01 scFv *I2L scFv x heFc(B) x I2L scFv
MSLN-CDH3 T-cell engager molecule 4 (example to Fig. 2B)	Seq ID 1249 + 1250	antiCDH3_01 scFv x heFc(A) x I2L scFv * I2L scFv x heFc(B) x antiMSLN_01 scFv
MSLN-CDH3 T-cell engager molecule 5 (example to Fig. 2D)	Seq ID 1254 + 1255 + 1253	antiCDH3_01 Fab x heFc(A) x I2L scFv* I2L scFv x heFc(B) x antiMSLN_01 scFv

MSLN-CDH3 T-cell engager molecule 6 (example to Fig. 2L)	Seq ID 1252 + 1257 + 1253 + 1256	antiCDH3_01 Fab x heFc(A) * I2L Fab x heFc(B) x antiMSLN_01 scFv x I2L scFv
MSLN-CDH3 T-cell engager molecule 7 (example to Fig. 2K)	Seq ID 1254 + 1258 + 1253 + 1256	antiCDH3_01 Fab x heFc(A) x I2L scFv * I2L Fab x heFc(B) x antiMSLN_01 scFv

Table 6: EC50 values and selectivity gaps of parental GSU WT cells versus target knockout GSU cells. b.c.t.: below calculation threshold involving a Fab comprising multichain multitargeting bispecific T-cell engager molecule

	EC50 GSU KO CDH3 [pM]	-fold selectivity gap	EC50 GSU WT [pM]	-fold selectivity gap	EC50 GSU KO MSLN [pM]
CDH3 T-cell engager molecule 1	b.c.t.	b.c.t.	125.52	0.82	102.58
MSLN T-cell engager molecule 1	2.83	1.4	2.01	b.c.t.	b.c.t.
MSLN-CDH3 T-cell engager molecule 24	735	5.6	132.0	276.6	36525

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Results are shown in Figure 6, i.e. Cytotoxicity curves of CDH3 T-cell engager molecule 1, MSLN T-cell engager molecule 1 and MSLN-CDH3 T-cell engager molecule 24 on parental double positive GSU WT cells versus target-knockout GSU cells. Effector cells were unstimulated Pan T-cells.

10 **Results:** CDH3 T-cell engager molecule 1 and MSLN T-cell engager molecule 1 that only target either CDH3 or MSLN, demonstrate comparable activity on single positive knockout cells vs. double positive GSU WT cells (0.8- to 1.4-fold selectivity gap). MSLN-CDH3 T-cell engager molecule 24 shows an increased activity on double positive GSU WT cells compared to target knockout GSU cells. The EC50 selectivity gap between double positive WT cells and CDH3 knockout cells was 5.6-fold, 15 the EC50 selectivity gap between double positive WT cells and MSLN knockout cells was 276.6-fold.

MSLN-CDH3 T-cell engager molecule 24 is a multichain multitargeting bispecific T-cell engager molecule, with one Fab target binding domain and one CD3 scFv binding domain forming a bispecific entity at the N-terminus of the polypeptide chain and one Fab target binding domain and one scFv

CD3 binding domain forming another bispecific entity at the C-terminus of the polypeptide chain, separated by a scFc domain.

Legend

CDH3 T-cell engager molecule 1 CH3-G8A_6-B12 x I2Cx scFc

(control)

MSLN T-cell engager molecule 1 MS 5-F11 x I2C x scFc

(control)

MSLN-CDH3 T-cell engager molecule 24 antiCDH3_01 Fab x I2L scFv x scFc

x antiMSLN_01 Fab x I2L scFv

(example to Fig. 3I)

5 [358] Example 2 Thermal stability of multitargeting bispecific antigen-binding molecules consisting of a singlechain or multichains with various domains or arrangements

Determination of aggregation and melting temperature T_{agg} and T_m

Multitargeting bispecific antigen-binding molecules were measured in a NanoTemper Prometheus

10 15 Pantia in triplicates and the aggregation temperature T_{agg} and melting temperature T_m was determined. The thermal unfolding assay was performed at 25°C – 95°C with a heating rate of 1°C/min and high sensitivity mode ON. The aggregation temperature T_{agg} (°C) is defined as the onset of the cumulant radius (nm), measured with dynamic light scattering (DLS). The melting temperature T_m (°C) is based on the changes in fluorescence to evaluate protein unfolding and / or aggregation and defines the point at which 50% of the molecule is unfolded. The T_m is defined as the first inflection point of the ratio 350nm/330nm of a thermal unfolding assay (first maxima of the first derivative of the ratio 350nm/330nm).

	T_{agg} [°C]	T_m [°C]
MSLN-CDH3 T-cell engager molecule 1	64.0	68.8
MSLN-CDH3 T-cell engager molecule 5	65.7	68.2
MSLN-CDH3 T-cell engager molecule 6	66.2	69.9
MSLN-CDH3 T-cell engager molecule 7	65.2	70.3

Table X: Aggregation temperature (T_{agg}) and melting temperature (T_m)

Results: MSLN-CDH3 T-cell engager molecules 1, 5, 6 and 7 show aggregation temperatures over 64°C, with MSLN-CDH3 T-cell engager molecule 6 showing the highest aggregation temperature of 66.2 °C. The melting temperature of all 4 molecules is higher than 68.2 °C, with MSLN-CDH3 T-cell engager molecule 7 exhibiting the highest melting temperature of 70.3 °C. The molecules were also tested regarding long-term storage stability and freeze-thaw stability and all molecules exhibited comparable characteristics.

10

MSLN-CDH3 T-cell engager molecule 1 is a single chain multitargeting bispecific antigen-binding molecule, with one bispecific entity (target binding domain and CD3 binding domain) at the N-terminus and one bispecific entity at the C-terminus of the molecule, separated by a single chain Fc-domain.

15

MSLN-CDH3 T-cell engager molecules 5,6 and 7 are multichain multitargeting bispecific antigen-binding molecules, with one target binding domain and one CD3 binding domain forming a bispecific entity at the N-terminus of the polypeptide chains and one target binding domain and one CD3 binding domain forming another bispecific entity at the C-terminus of the polypeptide chains, separated by a hetero Fc domain. The target and CD3 binding domains and their arrangements vary between constructs 5-7. The presented data demonstrates, that multichain multitargeting bispecific antigen-binding molecules are at least as resistant to high temperatures as single chain multitargeting bispecific antigen-binding molecules.

20

Legend	Seq ID	
MSLN-CDH3 T-cell engager molecule 1	Seq ID 1272	antiCDH3_01 scFv x I2L scFv x scFc x antiMSLN_01 scFv x I2L scFv
MSLN-CDH3 T-cell engager molecule 5	Seq ID 1254 + 1255 + 1253	antiCDH3_01 Fab x heFc(A) x I2L scFv* I2L scFv x heFc(B) x antiMSLN_01 scFv
MSLN-CDH3 T-cell engager molecule 6	Seq ID 1252 + 1257 + 1253 + 1256	antiCDH3_01 Fab x heFc(A) * I2L Fab x heFc(B) x antiMSLN_01 scFv x I2L scFv
MSLN-CDH3 T-cell engager molecule 7	Seq ID 1254 + 1258 + 1253 + 1256	antiCDH3_01 Fab x heFc(A) x I2L scFv * I2L Fab x heFc(B) x antiMSLN_01 scFv

Table 6: Sequence Table: Linkers, which may be indicated in the description as “G4”, “(G4S)n”, “(G4Q)n” or the like are not necessarily indicated in the table with such linked binding domain in order to maintain readability. The absence of such linker indication does not mean that the molecule in the table differs from the corresponding molecule in the description under a denomination which comprises the linker information. “CC” indicates disulfide bonds within a binding domain, “I2L”, “I2C”, “I2M” and “I2M2” indicate CD3 binding domains, respectively. Target binding domains may be abbreviated such as “CH3” for “CDH3”, “CL1” for “CLL1”, “FL” for “FLT3” and “MS” for “MSLN”. For most positions in the consensus sequences, “X” is the most restrictive ambiguity symbol. The amino acids “X” stands for are listed for CDRs of CDH3 binding domain in claim 35, for CDRs of MSLN binding domains in claim 36, for VH/VL of CDH3 binding domain in claim 37 and for VH/VL of MSLN binding domain in claim 38.

SEQ ID NO:	Designation	Source		Sequence
1.	(G4Q)3 - Linker	artificial	Aa	GGGGQGGGGQGGGGQ
2.	(G4S)10 - Linker	artificial	aa	GGGGSGGGGSGGGGSGGGGSGGGGSGGGGS GGGSGGGGSGGGGSGGGGS
3.	(G4S)3 - Linker	artificial	aa	GGGGSGGGGSGGGGS
4.	G(EAAAK)10 - Linker	artificial	aa	GEAAAKEAAAKEAAAKEAAAKEAAAKEAAA KEAAAKEAAAKEAAAKEAAAKEAAAKEAAK
5.	G4 - Linker	artificial	aa	GGGG
6.	G4Q - Linker	artificial	aa	GGGGQ
7.	G4S - Linker, spacer control	artificial	aa	GGGGS
8.	S(G4S)10 - Linker	artificial	aa	SGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS GGGGSGGGGSGGGGSGGGGS
9.	S(G4S)3 - Linker	artificial	aa	SGGGGSGGGGSGGGGS
10.	SG(EAAAK)10 - Linker	artificial	aa	SGEAAAKEAAAKEAAAKEAAAKEAAAKEAA AKEAAAKEAAAKEAAAKEAAAKEAAAKEAAK
11.	SG4Q - Linker	artificial	aa	SGGGGQ
12.	SG4S - Linker	artificial	aa	SGGGGS
13.	(EAAAK)10 - Spacer	artificial	aa	EAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEAAK EAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEAAK
14.	(G4S)10 - Spacer control	artificial	aa	GGGGSGGGGSGGGGSGGGGSGGGGSGGGGS GGGSGGGGSGGGGSGGGGS
15.	Human Serum Albumin (HSA) - Spacer	artificial	Aa	DAHKSEVAHRFKDLGEENFKALVLIQFAQYLQ QCPFEDHVKLVNEVTEFAKTCVADESAENCCK SLHTLFGDKLCTVATLRETYGEMADCCAKQEP ERNECFLQHKDDPNLPRLRPEVDVMCTAFH DNEETFLLKLYEYIARRHPYFYAPELFFAKRY KAAFTECCQAADKAAACLLPKLDELREDEGKASS AKQRLKCASLQKFGERAFAKAWAVARLSQRFP KAFAEAVSKLVTDLTKVHTECCHGDLLECADD RADLAKYICENQDSISSKLEKCEKPLLEKSHCI AEVENDEMPADLPSLAADFVESKDVCKNYAE AKDVFLLGMFLYEYARRHPDYSVLLLLRLAKT YETTLEKCCAAADPHECYAKVFDEFKPLVEEP QNLIKQNCLEFELGELYKFNALLVRYTKKVP QVSTPTLVEVSRNLGKVGSKCCKHPEAKRMP

				AEDYLSVVLNQLCVLHEKTPVSDRVTKCCTES LVNRRPCFSALEVDETYVPKEFNAETFTFHADI CTLSEKERQIKKQTALVELVKHKPKATKEQLK AVMDDFAAFVEKCKKADDKETCF AE EGKKL V AASQAALGL
16.	PD1 (ECD 25-167) - Spacer	artificial	Aa	LDSPDRPWNPTFSPALLVVTEGDNATFTCSFS NTSESFVLNWyRMSPSNQTDKLAAPEDRSQP GQDCRFRVTQLPNGRDFHMSVVRARRNDSGT YLCGAISLAPKAQIKESLRAELRV TERRAEVPT AHPSPSPRPAGQFQ
17.	Fc monomer-1 - c/+g	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
18.	Fc monomer-2 - c/+g/delGK	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
19.	Fc monomer-3 - c/+g	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYNSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
20.	Fc monomer-4 - c/+g/delGK	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYNSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
21.	Fc monomer-5 - c/+g	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYGSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
22.	Fc monomer-6 - c/+g/delGK	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYGSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
23.	Fc monomer-7 -	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS

	c/+g			RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYNSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
24.	Fc monomer-8- c/+g/delGK	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYNSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
25.	scFc - Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
26.	scFc-2 Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG GGGSGGGGSGGGGSGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
27.	scFc-3 Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYNSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYNSTYRVVSVLTVLHQDWLNG

				KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
28.	scFc-4 Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYNSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG GGGSGGGGSGGGGSGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPREEQYNSTYRVVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
29.	scFc-5 Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYGSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYGSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
30.	scFc-6 Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPREEQYGSTYRVVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG GGGSGGGGSGGGGSGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPREEQYGSTYRVVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSP
31.	scFc-7 Spacer	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYNSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE

				SNGQPENNYKTTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYNSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG K
32.	scFc-8 Spacer	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYNSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG GGGSGGGGSGGGGSGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYNSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSP
33.	scFc_mod_GQ_clippingvariant – Spacer	artificial	Aa	CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEEPEVKFNWYVDGVEVHNAKTK PCEEQYGSTYRCVSVLTVLHQDWLNGKEYKC KVSNAKALPAPIEKTISKAKGQPREPQVYTLPPS REEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ PENNYKTTTPPVLDSDGSFFLYSKLTVDKSRWQ QGNVFCSCVMHEALHNHYTQKSLSLSPGKGG GGQGGGGQGGGGQGGGGQGGGGQGGGGQCP PCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTC VVVDVSHEEPEVKFNWYVDGVEVHNAKTKPC EEQYGSTYRCVSVLTVLHQDWLNGKEYKCKV SNKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPPVLDSDGSFFLYSKLTVDKSRWQQGN VFCSCVMHEALHNHYTQKSLSLSPGK
34.	2x scFc – double size Spacer	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGYSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGYSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG

				KDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSGDSFFLYSKLTV DKSRWQQGNVFCFSVMHEALHNHYTQKSLSL SPGKGGGGSGGGGSGGGGSGGGGSGGGGSGG GGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSGDSFFLYSK LTVDKSRWQQGNVFCFSVMHEALHNHYTQKS LSLSPGK
35.	heteroFc (A) – Spacer	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLN KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYDTTPVLDSGDSFFLYSDLTVDKS RWQQGNVFCFSVMHEALHNHYTQDSLSLSPG K
36.	heteroFc (B) – Spacer	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLN KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTTPVLKSDGDSFFLYSKLTVDK SRWQQGNVFCFSVMHEALHNHYTQKSLSLSP GK
37.	I2C - HCDR1	artificial	Aa	KYAMN
38.	I2C - HCDR2	artificial	Aa	RIRSKYNNYATYYADSVKD
39.	I2C - HCDR3	artificial	Aa	HGNFGNSYISYWAY
40.	I2C - LCDR1	artificial	Aa	GSSTGAVTSGNYPN
41.	I2C - LCDR2	artificial	aa	GTKFLAP
42.	I2C - LCDR3	artificial	aa	VLWYSNRWV
43.	I2C – VH	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGLEWVARIRSKYNNYATYY ADSVKDRFTISRDDSKNTAYLQMNNLKTEDA VYYCVRHGNFGNSYISYWAYWGQGLTVVSS
44.	I2C – VL	artificial	aa	QTVVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWVQQKPGQAPRGLIGGKFLAPGTPARFS GSLGGAALTLGSGVQPEDEAEYYCVLWYSN RWVFGGGTKLTVL
45.	I2C_44/100cc - HCDR1	artificial	aa	KYAMN
46.	I2C_44/100cc - HCDR2	artificial	aa	RIRSKYNNYATYYADSVKD
47.	I2C_44/100cc - HCDR3	artificial	aa	HGNFGNSYISYWAY
48.	I2C_44/100cc - LCDR1	artificial	aa	GSSTGAVTSGNYPN
49.	I2C_44/100cc -	artificial	Aa	GTKFLAP

	LCDR2			
50.	I2C_44/100cc - LCDR3	artificial	Aa	VLWYSNRWV
51.	I2C_44/100cc - VH	artificial	Aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKCLEWVARIRSKYNNYATYY ADSVKDRFTISRDDSKNTAYLQMNNLKTEDTA VYYCVRHGNFGNSYISYWAYWGQGLTVTVSS
52.	I2C_44/100cc - VL	artificial	Aa	QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWWVQKPGQAPRGLIGGTKFLAPGTPARFS GSLGGAALTLSGVQPEDEAEYYCVLWYSN RWVFGCGTKLTVL
53.	I2E - HCDR1	artificial	Aa	KYAIN
54.	I2E - HCDR2	artificial	Aa	RIRSKYNNYATYYADAVKD
55.	I2E - HCDR3	artificial	Aa	AGNFGSSYISYWAY
56.	I2E - LCDR1	artificial	Aa	GSSTGAVTSGNYPN
57.	I2E - LCDR2	artificial	Aa	GTKFLAP
58.	I2E - LCDR3	artificial	aa	VLWYSNRWV
59.	I2E - VH	artificial	aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AINWVRQAPGKGLEWVARIRSKYNNYATYYA DAVKDRFTISRDDSKNTVYLQMNNLKTEDTA VYYCARAGNFGSSYISYWAYWGQGLTVTVSS
60.	I2E - VL	artificial	aa	QTVVTQEPSLTVSPGGTVTITCGSSTGAVTSGN YPNWWVQKPGQAPRGLIGGTKFLAPGTPARFS GSLSGGAALTLSGVQPEDEAEYYCVLWYSN RWVFGSGTKLTVL
61.	I2L - HCDR1	artificial	aa	KYAMN
62.	I2L - HCDR2	artificial	aa	RIRSKYNNYATYYADAVKD
63.	I2L - HCDR3	artificial	aa	AGNFGSSYISYFAY
64.	I2L - LCDR1	artificial	aa	GSSTGAVTSGNYPN
65.	I2L - LCDR2	artificial	aa	GTKFLAP
66.	I2L - LCDR3	artificial	Aa	VLYYSNRWV
67.	I2L - VH	artificial	Aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVTVSS
68.	I2L - VL	artificial	Aa	QTVVTQEPSLTVSPGGTVTITCGSSTGAVTSGN YPNWIQKPGQAPRGLIGGTKFLAPGTPARFSG SLEGGKAALTLSGVQPEDEAEYYCVLYYSNR WVFGSGTKLTVL
69.	I2M2 - HCDR1	artificial	Aa	KYAIN
70.	I2M2 - HCDR2	artificial	Aa	RIRSKYNNYATYYADAVKD
71.	I2M2 - HCDR3	artificial	Aa	NANFGTSYISYFAY
72.	I2M2 - LCDR1	artificial	Aa	GSSTGAVTSGNYPN
73.	I2M2 - LCDR2	artificial	Aa	GTKFLAP
74.	I2M2 - LCDR3	artificial	Aa	VLWYSNRWV
75.	I2M2 - VH	artificial	aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AINWVREAPGKGLEWVARIRSKYNNYATYYA DAVKDRFTISRDDSKNTAYLQMNNLKTEDTA VYYCVRNANFGTSYISYFAYWGQGLTVTVSS
76.	I2M2 - VL	artificial	aa	QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWWVQKPGQAPRGLIGGTKFLAPGTPARFS GSLGGAALTLSGVQPEDEAEYYCVLWYSN RWVFGSGTKLTVL
77.	MS 01-G11 CC - HCDR1	artificial	aa	DYYMT

78.	MS 01-G11 CC - HCDR2	artificial	aa	YISSSGSTIYYAEAVKG
79.	MS 01-G11 CC - HCDR3	artificial	aa	DRNSHFDY
80.	MS 01-G11 CC - LCDR1	artificial	aa	RASQGIRTWLA
81.	MS 01-G11 CC - LCDR2	artificial	aa	GASGLQS
82.	MS 01-G11 CC - LCDR3	artificial	aa	QQAESFPRT
83.	MS 01-G11 CC - VH	artificial	Aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDY YMTWIRQAPGKCLEWLSYISSSGSTIYYAEAV KGRFTISRDNKNSLFLQMNSLRAEDTAVYYC ARDRNSHFDYWGQGLVTVSS
84.	MS 01-G11 CC - VL	artificial	Aa	DIMTQSPSSVSASVGDRVTITCRASQGIRTWLA WYQQKPGKAPKLLIYGASGLQSGVPSRFSGSG SGTDFTLTISSLQPEDFATYYCQQAESFPRTFGC GTKVEIK
85.	MS 01-G11 CC EI - VL	artificial	Aa	EIMTQSPSSVSASVGDRVTITCRASQGIRTWLA WYQQKPGKAPKLLIYGASGLQSGVPSRFSGSG SGTDFTLTISSLQPEDFATYYCQQAESFPRTFGC GTKVEIK
86.	MS 15-B12 CC - HCDR1	artificial	Aa	SSSYFWG
87.	MS 15-B12 CC - HCDR2	artificial	Aa	NIYYSGSSNYNPSLKS
88.	MS 15-B12 CC - HCDR3	artificial	Aa	LPRGDRDAFDI
89.	MS 15-B12 CC - LCDR1	artificial	Aa	RASQGISNYLA
90.	MS 15-B12 CC - LCDR2	artificial	Aa	AASTLQS
91.	MS 15-B12 CC - LCDR3	artificial	Aa	QQSYSTPFT
92.	MS 15-B12 CC - VH	artificial	aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSS
93.	MS 15-B12 CC - VL	artificial	aa	DIVMTQSPSSLSASVGDRVTITCRASQGISNYL AWYQQKPGKVPKLLIYAASTLQSGVPSRFSGSG GSGTDFTLTISSLQPEDFATYYCQQSYSTPFTFG CGTKVEIK
94.	MS 15-B12 CC EI - VL	artificial	aa	EIVMTQSPSSLSASVGDRVTITCRASQGISNYLA WYQQKPGKVPKLLIYAASTLQSGVPSRFSGSG SGTDFTLTISSLQPEDFATYYCQQSYSTPFTFGC GTKVEIK
95.	MS 25-E3 CC - HCDR1	artificial	aa	SSSYFWV
96.	MS 25-E3 CC - HCDR2	artificial	aa	SIYYSGSTYYNPSLKS
97.	MS 25-E3 CC - HCDR3	artificial	aa	LPRGDRMTFDI
98.	MS 25-E3 CC - LCDR1	artificial	aa	RASQSVSSSYLA
99.	MS 25-E3 CC -	artificial	aa	GASSRAT

	LCDR2			
100.	MS 25-E3 CC - LCDR3	artificial	Aa	QQYGSSPFT
101.	MS 25-E3 CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWVWIRQPPGKCLEWIGSIYYSGSTYYNPSLKS RVTISVDTSKNQFSLKLNSTVTAADTAVYYCAR LPRGDRMTFDIWGQGTMVTVSS
102.	MS 25-E3 CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPFTFGC GTKLEIK
103.	MS 36-C5 CC - HCDR1	artificial	Aa	SYAMS
104.	MS 36-C5 CC - HCDR2	artificial	Aa	AISGSGEQWYYAPSVKG
105.	MS 36-C5 CC - HCDR3	artificial	Aa	VRNYYGSGSLDY
106.	MS 36-C5 CC - LCDR1	artificial	Aa	RASQSFSSAYLA
107.	MS 36-C5 CC - LCDR2	artificial	Aa	GASIRAT
108.	MS 36-C5 CC - LCDR3	artificial	Aa	QQYGSSLT
109.	MS 36-C5 CC - VH	artificial	aa	EVQLLESGGGVVQPGRSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGEQWYYAPSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKVRNYYGSGSLDYWGQGTLLVTVSS
110.	MS 36-C5 CC - VL	artificial	aa	EIVLTQSPGTLSPGERATLSCRASQSFSSAYL AWYQQKPGQAPRLLIYGASIRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSLTFGC GTKVEIK
111.	MS 36-G7 CC - HCDR1	artificial	aa	SYAMS
112.	MS 36-G7 CC - HCDR2	artificial	aa	AISGSGEGDYYANSVKG
113.	MS 36-G7 CC - HCDR3	artificial	aa	VRNYYGSGSLDY
114.	MS 36-G7 CC - LCDR1	artificial	aa	RASQSVSSTYLA
115.	MS 36-G7 CC - LCDR2	artificial	aa	GASIRAT
116.	MS 36-G7 CC - LCDR3	artificial	aa	QQYGSSLT
117.	MS 36-G7 CC - VH	artificial	Aa	EVQLLESGGGVVQPGRSLRLSCAASGFTFSSYA MSWVRQAPGMCLEWVSAISGSGEGDYYANSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKVRNYYGSGSLDYWGQGTLLVTVSS
118.	MS 36-G7 CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSTYL AWYQQKPGQAPRLLIYGASIRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSLTFGC GTKVEIK
119.	MS 37-E5 CC - HCDR1	artificial	Aa	SYAMS
120.	MS 37-E5 CC - HCDR2	artificial	Aa	AISGSGGSTYYAIDVKG

121.	MS 37-E5 CC - HCDR3	artificial	Aa	EGYYPGSGYPLYYYFGMDV
122.	MS 37-E5 CC - LCDR1	artificial	Aa	RASQSVSSSYLA
123.	MS 37-E5 CC - LCDR2	artificial	Aa	GASSRAT
124.	MS 37-E5 CC - LCDR3	artificial	Aa	QQYGSSPIFT
125.	MS 37-E5 CC - VH	artificial	Aa	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGGSTYYAIDV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKEGYYPGSGYPLYYYFGMDVWGQGTTVTVS S
126.	MS 37-E5 CC - VL	artificial	aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPIFTFG CGTKVEIK
127.	MS 46-A3 CC - HCDR1	artificial	aa	SYGMG
128.	MS 46-A3 CC - HCDR2	artificial	aa	VISYHGSKNYADAVKG
129.	MS 46-A3 CC - HCDR3	artificial	aa	EGAHFGSGSYPLYYYAMDV
130.	MS 46-A3 CC - LCDR1	artificial	aa	RASQSVSSSYLA
131.	MS 46-A3 CC - LCDR2	artificial	aa	GASIRAT
132.	MS 46-A3 CC - LCDR3	artificial	aa	QQTGSSPIFT
133.	MS 46-A3 CC - VH	artificial	aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMGWVRQAPGKCLEWVAVISYHGSKNYAD AVKGRFTISRDNKNTLYLQMNSLRAEDTAVY YCAREGAHFGSGSYPLYYYAMDVWGQGT TVTSS
134.	MS 46-A3 CC - VL	artificial	Aa	EIVTQSPGTLSPGERATLSCRASQSVSSSYLA WYQQKPGQAPRLLIYGASIRATGIPDRFSGSGS GTDFTLTISRLEPEDFAVYYCQQTGSSPIFTFGC GTKVEIK
135.	MS R195L CC - HCDR1	artificial	Aa	SYAMS
136.	MS R195L CC - HCDR2	artificial	Aa	AISGSGEFSYYAAAVKG
137.	MS R195L CC - HCDR3	artificial	Aa	VRNYYGSGSLDY
138.	MS R195L CC - LCDR1	artificial	Aa	RASQSVSSTYLA
139.	MS R195L CC - LCDR2	artificial	Aa	GASIRAT
140.	MS R195L CC - LCDR3	artificial	Aa	QQYQSSLT
141.	MS R195L CC - VH	artificial	Aa	EVQLLESGGGVVQPGRSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGEFSYYAAAV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKVRNYYGSGSLDYWGQGTTLTVSS
142.	MS R195L CC -	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSTYL

	VL			AWYQQKPGQAPRLLIYGASIRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYQSSLTFGC GTKVEIK
143.	MS R4L CC - HCDR1	artificial	aa	GYyIH
144.	MS R4L CC - HCDR2	artificial	aa	WINPNSGGTNYAQKFQG
145.	MS R4L CC - HCDR3	artificial	aa	VEAVAGREYYYFSGMDV
146.	MS R4L CC - LCDR1	artificial	aa	SGEKLGDKYVY
147.	MS R4L CC - LCDR2	artificial	aa	QSTKRPS
148.	MS R4L CC - LCDR3	artificial	aa	QAYHASTAV
149.	MS R4L CC - VH	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTG YYIHWVRQAPGQCLEWMGWINPNSGGTNYA QKFQGRVTMTRDTSISTAYMELSLRSDDTAV YYCARVEAVAGREYYYFSGMDVWGQGTVT VSS
150.	MS R4L CC - VL	artificial	aa	SYELTQPPSVSVSPGQTASITCSGEKLGDKYVY WYQQKPGQSPVLVIYQSTKRPSGVPERFSGSNS GNTATLTISGTQAMDEADYYCQAYHASTAVF GCGTKLTVL
151.	MS H2 - HCDR1	artificial	Aa	SYGMG
152.	MS H2 - HCDR2	artificial	Aa	VISYDGSNKYYADSVKG
153.	MS H2 - HCDR3	artificial	Aa	EGAHFGSGSYYPlyYYYAMDV
154.	MS H2 - LCDR1	artificial	Aa	RASQSVSSSYLA
155.	MS H2 - LCDR2	artificial	Aa	GASIRAT
156.	MS H2 - LCDR3	artificial	Aa	QQYGSSPIFT
157.	MS H2 - VH	artificial	Aa	EVQLLESgggVVQPGRSLRLSCAASGFTfSSyG MGWVRQAPGKGLEWVAVISYDGSNKYYADS VKGRFTISRDNskNTLYLQMNSLRAEDTAVYY CAREGAHFGSGSYYPlyYYYAMDVWGQGT TVSS
158.	MS H2 - VL	artificial	Aa	ELTLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASIRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYgSSPIFTFG PGTKVEIK
159.	CH3 005-D5 CC - HCDR1	artificial	Aa	SYPIN
160.	CH3 005-D5 CC - HCDR2	artificial	aa	VIWTGGGTNYASSVKG
161.	CH3 005-D5 CC - HCDR3	artificial	aa	SRGVYDFKGRGAMDY
162.	CH3 005-D5 CC - LCDR1	artificial	aa	KSSQSLlySSNQKNyFA
163.	CH3 005-D5 CC - LCDR2	artificial	aa	WASTRES
164.	CH3 005-D5 CC - LCDR3	artificial	aa	QQYYSYPYT
165.	CH3 005-D5 CC - VH	artificial	aa	EVQLLESggGLVQPGGSLRLSCAASGfSfSSyPI NWVRQAPGKCLEWVGVIWTGGGTNYASSVK GRFTISRDNskNTVYLQMNSLRAEDTAVYYCA KSRGVYDFKGRGAMDYWGQGTlTVSS

166.	CH3 005-D5 CC - VL	artificial	aa	DIVMTQSPDSLAVSLGERATINCKSSQSLLYSS NQKNYFAWYQQKPGQPPKLLIYWASTRESGV PDRFSGSGSGTDFTLTISLQAEDVAVYYCQQY YSYPYTFGCGTKLEIK
167.	CH3 005-D5 CC EI - VL	artificial	aa	EIVMTQSPDSLAVSLGERATINCKSSQSLLYSSN QKNYFAWYQQKPGQPPKLLIYWASTRESGVP DRFSGSGSGTDFTLTISLQAEDVAVYYCQQY YSYPYTFGCGTKLEIK
168.	CH3 03-C8 CC - HCDR1	artificial	Aa	SYWMH
169.	CH3 03-C8 CC - HCDR2	artificial	Aa	VISGSKSYTIYNQKVKG
170.	CH3 03-C8 CC - HCDR3	artificial	Aa	SGPGYFDV
171.	CH3 03-C8 CC - LCDR1	artificial	Aa	RASENIYSYLA
172.	CH3 03-C8 CC - LCDR2	artificial	Aa	NAKTLAE
173.	CH3 03-C8 CC - LCDR3	artificial	Aa	QHLNMTPYT
174.	CH3 03-C8 CC - VH	artificial	Aa	EVQLLESGGGLVQPGGSLRLSCAASGYTFSSY WMHWVRQAPGKCLEWMGVISGSKSYTIYNQ KVKGRFTISRDNKNTVYLQMNSLRAGDTAV YYCARSGPGYFDVWGQGTMTVSS
175.	CH3 03-C8 CC - VL	artificial	Aa	DIQLTQSPSFLSASVGDRVTITCRASENIYSYLA WYQQKPGKAPKLLIYNAKTLAEGVPSRFSGSG SGTEFTLTISLQPEDFGTYQCQHLNMTPYTFG CGTKLEIK
176.	CH3 03-C8 CC EI - VL	artificial	Aa	EIQLTQSPSFLSASVGDRVTITCRASENIYSYLA WYQQKPGKAPKLLIYNAKTLAEGVPSRFSGSG SGTEFTLTISLQPEDFGTYQCQHLNMTPYTFG CGTKLEIK
177.	CH3 08-A11 CC - HCDR1	artificial	aa	SYWMH
178.	CH3 08-A11 CC - HCDR2	artificial	aa	KIDPSDDYTNYNQKVKG
179.	CH3 08-A11 CC - HCDR3	artificial	aa	WDYNYFDV
180.	CH3 08-A11 CC - LCDR1	artificial	aa	RASSSVSYM
181.	CH3 08-A11 CC - LCDR2	artificial	aa	GTSNLVS
182.	CH3 08-A11 CC - LCDR3	artificial	aa	QQWSSYPLT
183.	CH3 08-A11 CC - VH	artificial	aa	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSY WMHWVRQTPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDKSKNTLYLQMNSLRAEDTAVY YCARWDYNYFDVWGQGTTVTSS
184.	CH3 08-A11 CC - VL	artificial	aa	EIVMTQSPATLSVSPGERATLTCRASSSVSYM WYQQKPGQAPRLLIYGTSNLVSGVPSRFSGSG SGTEFTLTISLQSEDFVYYCQQWSSYPLTFG CGTKVEIK
185.	CH3 14-D1 CC - HCDR1	artificial	Aa	SYWMH
186.	CH3 14-D1 CC -	artificial	Aa	VIYTSYTYIYNQKFQG

	HCDR2			
187.	CH3 14-D1 CC - HCDR3	artificial	Aa	SGPGYFDV
188.	CH3 14-D1 CC - LCDR1	artificial	Aa	RASGNIHNYLA
189.	CH3 14-D1 CC - LCDR2	artificial	Aa	NAKTLAE
190.	CH3 14-D1 CC - LCDR3	artificial	Aa	QHFAWTPYT
191.	CH3 14-D1 CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKV SCKASGYTFTSY WMHWVRQAPGQCLEWMGV IYTSGSYTIYNQ KFQGRVTMTRDTSTSTAY MELSSLRSEDVAVY YCARSGPGYFDVWGQGT MVTVSS
192.	CH3 14-D1 CC - VL	artificial	Aa	DIQLTQSPSFLSASVGDRVT ITCRASGNIHNYLA WYQQKPGKAPKLLIYNAK TLAEGVPSRFSGSG SGTEFTLKISLQPEDFATY YCQHFAWTPYTFG CGTKLEIK
193.	CH3 14-D1 CC EI - VL	artificial	Aa	EIQLTQSPSFLSASVGDRVT ITCRASGNIHNYLA WYQQKPGKAPKLLIYNAK TLAEGVPSRFSGSG SGTEFTLKISLQPEDFATY YCQHFAWTPYTFG CGTKLEIK
194.	CH3 15-E11 CC - HCDR1	artificial	aa	NYWMN
195.	CH3 15-E11 CC - HCDR2	artificial	aa	NIAYGVKGTNYNQKFQG
196.	CH3 15-E11 CC - HCDR3	artificial	aa	RYFYVMDY
197.	CH3 15-E11 CC - LCDR1	artificial	aa	RASQDISNYLN
198.	CH3 15-E11 CC - LCDR2	artificial	aa	YTSRLHS
199.	CH3 15-E11 CC - LCDR3	artificial	aa	VQYAQFPLT
200.	CH3 15-E11 CC - VH	artificial	aa	QVQLVQSGAEVKKPGASVK VSCASGYTFTN YWMNWVRQAPGQCLEWM GNIAYGVKGTNY NQKFQGRVTMTVDTSSST AYMELSRRLRSDDTA VYYCATRYFYVMDYWGQ GTLVTVSS
201.	CH3 15-E11 CC - VL	artificial	aa	DIQMTQSPSSLSASVGDRVT ITCRASQDISNYL NWXQQKPGKVPKLLIYYT SRLHSGVPSRFSGSG GSGTDFLTISSLQPEDVAT YYCVQYAQFPLTF GCGTKVEIK
202.	CH3 15-E11 CC EI - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVT ITCRASQDISNYLN WYQQKPGKVPKLLIYYT SRLHSGVPSRFSGSGS GTDFTLTISSLQPEDVAT YYCVQYAQFPLTFGC GTKVEIK
203.	CH3 22-A12 CC - HCDR1	artificial	Aa	SSWMN
204.	CH3 22-A12 CC - HCDR2	artificial	Aa	RIYGTGETKYSKGFQG
205.	CH3 22-A12 CC - HCDR3	artificial	Aa	QRDYGALYAMDY
206.	CH3 22-A12 CC - LCDR1	artificial	Aa	RASDDIYSYLA
207.	CH3 22-A12 CC - LCDR2	artificial	Aa	NAKTLAE

208.	CH3 22-A12 CC - LCDR3	artificial	Aa	QNHDRTPFT
209.	CH3 22-A12 CC - VH	artificial	Aa	QVQLVQSGAEVVKPGASVKV SCKASGYTFTSS WMNWVRQAPGQCLEWMGRIY TGTGETKYSG KFQGRVTITRDTSASTAYMEL SSLTSEDVAVYY CARQRDYGALYAMDYWGQGL VTVSS
210.	CH3 22-A12 CC - VL	artificial	Aa	DIQLTQSPSFLSASVGDRVTIT CRASDDIYSYLA WYQQKPGKAPKLLVYNAKTLA EGVPSRFSGS GSGTEFTLTISLQPEDFATYYC QNHDRTPFTFG CGTKVDIK
211.	CH3 22-A12 CC EI - VL	artificial	aa	EIQLTQSPSFLSASVGDRVTIT CRASDDIYSYLA WYQQKPGKAPKLLVYNAKTLA EGVPSRFSGS GSGTEFTLTISLQPEDFATYYC QNHDRTPFTFG CGTKVDIK
212.	CH3 24-D7 CC - HCDR1	artificial	aa	NYWMN
213.	CH3 24-D7 CC - HCDR2	artificial	aa	NIHSKAHGTNYNQKFQG
214.	CH3 24-D7 CC - HCDR3	artificial	aa	RYFYVMDY
215.	CH3 24-D7 CC - LCDR1	artificial	aa	RASQDISNYLN
216.	CH3 24-D7 CC - LCDR2	artificial	aa	YTSRLHS
217.	CH3 24-D7 CC - LCDR3	artificial	aa	VQYAQFPLT
218.	CH3 24-D7 CC - VH	artificial	aa	QVQLVQSGAEVKKPGASVKV SCKASGYTFTN YWMNWVRQAPGQCLEWMGNI HSKAHGTNY NQKFQGRVTMTVDTSSSTAY MELSRRLRSDDTA VYYCATRYFYVMDYWGQGL VTVSS
219.	CH3 24-D7 CC - VL	artificial	Aa	DIQMTQSPSSLSASVGDRVTIT CRASQDISNYL NWXQQKPGKVPKLLIYYTSR LHSGVPSRFSGS GSGTDFTLTISLQPEDVATYYC VQYAQFPLTF GCGTKVEIK
220.	CH3 24-D7 CC EI - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVTIT CRASQDISNYLN WYQQKPGKVPKLLIYYTSR LHSGVPSRFSGS GSGTDFTLTISLQPEDVATYYC VQYAQFPLTFGC GTKVEIK
221.	CH3 26-E5 CC - HCDR1	artificial	Aa	SYWMH
222.	CH3 26-E5 CC - HCDR2	artificial	Aa	VIRTSTSYTIYNQKFKG
223.	CH3 26-E5 CC - HCDR3	artificial	Aa	SGPGYFDV
224.	CH3 26-E5 CC - LCDR1	artificial	Aa	RASENIYSYLA
225.	CH3 26-E5 CC - LCDR2	artificial	Aa	NAKTLAE
226.	CH3 26-E5 CC - LCDR3	artificial	Aa	QHNYGTPYT
227.	CH3 26-E5 CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKV SCKASGYTFTSY WMHWVRQAPGQCLEWMGVIR TSTSYTIYNQK FKGRVTMTRDTSTSTVYME LSSLRSEDVAVYY CARSGPGYFDVWGQGMVTVSS
228.	CH3 26-E5 CC -	artificial	aa	DIQLTQSPSFLSASVGDRVTIT CRASENIYSYLA

	VL			WYQQKPGKAPKLLIYNAKTLAEGVPSRFSGSG SGTEFTLTISSLQPEDFATYYCQHNYGTPYTFG CGTKLEIK
229.	CH3 26-E5 CC EI - VL	artificial	aa	EIQLTQSPSFLSASVGDRTITCRASENIYSYLA WYQQKPGKAPKLLIYNAKTLAEGVPSRFSGSG SGTEFTLTISSLQPEDFATYYCQHNYGTPYTFG CGTKLEIK
230.	CH3 R164L CC - HCDR1	artificial	aa	SYWMY
231.	CH3 R164L CC - HCDR2	artificial	aa	KIDPSDDYTNYNQKVKG
232.	CH3 R164L CC - HCDR3	artificial	aa	WDYTHFDV
233.	CH3 R164L CC - LCDR1	artificial	aa	RASSSVSYMH
234.	CH3 R164L CC - LCDR2	artificial	aa	GTSNLAS
235.	CH3 R164L CC - LCDR3	artificial	aa	QQWSSYPLT
236.	CH3 R164L CC - VH	artificial	Aa	EVQLLESGLVQPGGSLRSLCAASGFTFSSY WMYWVRQAPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDNSKNTLYLQMNSLRAEDSAVY YCARWDYTHFDVWGQGTITVTVSS
237.	CH3 R164L CC - VL	artificial	Aa	EIVMTQSPATLSVSPGERATLSCRASSSVSYMH WYQQKPGQAPRLLIYGTSNLSGVPVRFSGSG SGTEFTLTISSLQSEDAVYYCQQWSSYPLTFG CGTKVEIK
238.	CH3 R170R CC - HCDR1	artificial	Aa	SYWMH
239.	CH3 R170R CC - HCDR2	artificial	Aa	KIDPSDDYTNYNQKVKG
240.	CH3 R170R CC - HCDR3	artificial	Aa	WDYSHFDV
241.	CH3 R170R CC - LCDR1	artificial	Aa	RASSSVSYMH
242.	CH3 R170R CC - LCDR2	artificial	Aa	GTSNLVS
243.	CH3 R170R CC - LCDR3	artificial	Aa	QQWSSYPLT
244.	CH3 R170R CC - VH	artificial	Aa	EVQLLESGLVQPGGSLRSLCAASGFTFSSY WMHWVRQTPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDKSKNTLYLQMNSLRAEDTAVY YCARWDYSHFDVWGQGTITVTVSS
245.	CH3 R170R CC - VL	artificial	aa	EIVMTQSPATLSVSPGERATLTCRASSSVSYMH WYQQKPGQAPRLLIYGTSNLVSGVPARFSGSG SGTEFTLTISSLQSEDAVYYCQQWSSYPLTFG CGTKVEIK
246.	CH3 005-D5 CCx I2Ccc(44/100)x (G4)x scFc x (G4) x MS 01-G11 CCx I2Ccc(44/100) - Full Sequence	artificial	aa	EVQLLESGLVQPGGSLRSLCAASGFSFSSYPI NWRQAPGKCLEWVGVIWTGGGTNYASSVK GRFTISRDNKNTVYLQMNSLRAEDTAVYYCA KSRGVYDFKGRGAMDYWGQGLTVTVSSGGG GSGGGGGGGSDIVMTQSPDSLAVSLGERATI NCKSSQSLLYSSNQKNYFAWYQQKPGPPKLL IYWASTRESGVPDRFSGSGSGTDFTLTISSLQAE DVAVYYCQYYSPYTFGCGTKLEIKSGGGGS

				<p>EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKCLEWVARIRSKYNNYATYY ADSVKDRFTISRDDSKNTAYLQMNNLKTEDTA VYYCVRHGNFGNSYISYWAYWGQGLTVTVSS GGGGSGGGGSGGGGSQTVVTQEPSLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQQKPGQAPRG LIGGTKFLAPGTPARFSGSLLGGKAALTLSGVQ PEDEAEYYCVLWYSNRWVFGCGTKLTVLGGG GDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPPEVTCVVVDVSHEDPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSGDGSFFLYSKLTV DKSRWQQGNVFSCSVMHEALHNHYTQKSLSL SPGKGGGGSGGGGSGGGGSGGGGSGGGGSGG GGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNQKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSGDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGQVQLVESGGGLVKPGGSLRLS CAASGFTFSDYYMTWIRQAPGKCLEWLSYISS GSTIYYAEAVKGRFTISRDNKNSLFLQMNSLR AEDTAVYYCARDNRNSHFYDWGQGLTVTVSSG GGGSGGGGSGGGGSDIMTQSPSSVSASVGDRV TITCRASQGIRTWLAWYQQKPGKAPKLLIYGA SGLQSGVPSRFSGSGSGTDFTLTISSLQPEDFAT YYCQQAESFPRTFGCGTKVEIKSGGGGSEVQL VESGGGLVQPGGSLKLSAASGFTFNKYAMN WVRQAPGKCLEWVARIRSKYNNYATYYADSV KDRFTISRDDSKNTAYLQMNNLKTEDTAVYYC VRHGNFGNSYISYWAYWGQGLTVTVSSGGGG SGGGGSGGGGSGGGGSQTVVTQEPSLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIGG TKFLAPGTPARFSGSLLGGKAALTLSGVQPEDEA EYYCVLWYSNRWVFGCGTKLTVL</p>
247.	<p>CH3 08-A11 CC x I2Ccc(44/100)x (G4S)3x scFcx (G4S)3x MS R4L CCx I2Ccc(44/100) - Full Sequence</p>	artificial	aa	<p>EVQLLESGGGLVQPGGSLRLSCAASGFTFSSY WMHWVRQTPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDKSKNTLYLQMNSLRAEDTAVY YCARWDYNYFDVWGQGTTVTVSSGGGGSGG GGSGGGGSEIVMTQSPATLSVSPGERATLTCRA SSVSVMHWYQQKPGQAPRLLIYGTSNLVSGV PARFSGSGGTEFTLTISSLQSEDFAVYYCQQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGLTVTVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGT VTLTCGSSTGA VTSNYPNWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLGGGGSGGGGSGGG GSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTL</p>

				<p>RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGSGGGGSQVQLVQSGAEVKKPG ASVKVSCKASGYTFTGYYIHWVRQAPGQCLE WMGWINPNSGGTNYAQKFQGRVTMTRDTSIS TAYMELSRLSDDTAVYYCARVEAVAGREYY YFSGMDVWGQGTITVTVSSGGGGSGGGGSGGG GSSYELTQPPSVSVSPGQTASITCSGEKLGDKY VYWYQQKPGQSPVLVIYQSTKRPSPVPERFSG SNSGNTATLTISGTQAMDEADYYCQAYHASTA VFGCGTKLTVLSSGGGGSEVQLVESGGGLVQPG GSLKLSCAASGFTFNKYAMNWVRQAPGKGM WVARIRSKYNNYATYYADAVKDRFTISRDDSK NTLYLQMNNLKTEDTAVYYCVRAGNFGSSYIS YFAYWGQGTITVTVSSGGGGSGGGGSGGGGSQ TVVTQEPSLTVSPGGTITITCGSSTGAVTSGNY PNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGS LEGGKAALTLSGVQPEDEAEYYCVLYYSNRW VFGSGTKLTVL</p>
<p>249.</p>	<p>CH3 08-A11 CCx I2Ccc(44/100)x (G4)x scFc x (G4) x MS R4L CCx I2Ccc(44/100) - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>EVQLLESGLVQPGGSLRLSCAASGFTFSSY WMHWVRQTPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDKSKNTLYLQMNSLRAEDTAVY YCARWDYNYFDVWGQGTITVTVSSGGGGSGG GGSGGGGSEIVMTQSPATLSVSPGERATLTCRA SSVSVMHWYQQKPGQAPRLLIYGTSLVSGV PARFSGSGSGTEFTLTISSLSQSEDFAVYYCQQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGTITVTVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGTITITCGSSTGA VTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLGGGGDKTHTCPPCP APPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VDVSHEDPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPEN YKTTTPVLDSGDSFFLYSKLTVDKSRWQQGN FSCSVMHEALHNHYTQKSLSLSPGKGGGGG GGSGGGGSGGGSGGGGSGGGGSDKHTHTCPP CPPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VVDVSHEDPEVKFNWYVDGVEVHNAKTKPCE EQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSGDSFFLYSKLTVDKSRWQQGN VFCFSVMHEALHNHYTQKSLSLSPGKGGGGG VQLVQSGAEVKKPGASVKVSCKASGYTFTGY YIHWVRQAPGQCLEWMGWINPNSGGTNYAQ</p>

				<p>KFQGRVTMTRDTSISTAYMELSRLRSDDTAVY YCARVEAVAGREYYFSGMDVWGQTTVTV SSGGGSGGGGSGGGGSSYELTQPPSVSVSPG QTASITCSGEKLGDKYVYWYQQKPGQSPVLVI YQSTKRPSGVPERFSGSNSGNTATLTISGTQAM DEADYYCQAYHASTAVFGCGTKLTVLSSGGGG SEVQLVESGGGLVQPGGSLKLSCAASGFTFNK YAMNWVRQAPGKCLEWVARIRSKYNNYATY YADSVKDRFTISRDDSKNTAYLQMNNLKTEDT AVYYCVRHGNFGNSYISYWAYWGGQTLTVTS SGGGGSGGGGSGGGGSGTQVVTQEPSLTVSPGG TVTLTCGSSTGAVTSGNYPNWVQKPGQAPR GLIGGTKFLAPGTPARFSGSLLGGKAALTLVSGV QPEDEAEYYCVLWYSNRWVFGCGTKLTVL</p>
<p>250.</p>	<p>CH3 15-E11 CC x I2L x (G4Q)3x scFcmo d x (G4Q)3 x MS 15- B12 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLVQSGAEVKKPGASVKVSKCASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQTLTVVSSGGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYC VQY AQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSCAASGFTFNKYAMNWVR QAPGKGM EWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQTLTVVSSGGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGTKFL APGTPARFSGSLEGGKAALTLVSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGQGGGGQ GGGGQCPPCPAPELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEEPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSDGSFFLYSKLTV DKSRWQQGNV FSCSVMHEALHNHYTQKSLSL SPGKGGGGQGGGGQGGGGQGGGGQGGGGQGG GGGQCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEEPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNV FSCSVMHEALHNHYTQKSLSLSPG KGGGGQGGGGQGGGGQGVQLQESGPGLVKPS ETLSLTCTVSGGSISSSSYFWGWIRQPPGKCLE WIGNIYSGSSNYPNPSLKSRTISVDTSKNQFSL KLSSVTAADTAVYYCARLPRGDRDAFDIWGQ GTMVTVSSGGGGQGGGGQGGGGQDIVMTQSP SSLSASVGDRVTITCRASQGISNYLAWYQQK GKVPKLLIYAAS TLQSGVPSRFSGSGSGTDFTL TISSLQPEDFATYYCQSYSTPFTFGCGTKVEIK SGGGGQEVQLVESGGGLVQPGGSLKLSCAASG FTFNKYAMNWVRQAPGKGM EWVARIRSKYN NYATYYADAVKDRFTISRDDSKNTLYLQMN</p>

				LKTEDTAVYYCVRAGNFGSSYISYFAYWGQGT LVTVSSGGGGQGGGGQGGGGQQTVVVTQEPSL TVSPGGTVTITCGSSTGAVTSGNYPNWIQKKPG QAPRGLIGGTKFLAPGTPARFSGSLEGGKAALT LSGVQPEDEAEYYCVLYYSNRWVFGSGTKLT VL
251.	CH3 15-E11 CC x I2L x G4 x scFc x G4 x MS 15-B12 CC x I2L_GQ - Full Sequence	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRSDDTA VYYCATRYFYVMDYWGQGT LVTVSSGGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQKPKGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYC VQY AQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGT LVTVSSGGGGGQGG GGGGGGGQQTVVVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGTKFL APGTPARFSGSLEGGKAALT LSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGCPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGGQGGG GQGGGGGQGGGGQGGGGGQGGGGQCPPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEEPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGQVQLQE SGPGLVKPSETLSLTCTVSGGSISSSSYFWGWR QPPGKCLEWIGNIYYSGSSNYPNLSKSRVTISV DTSKNQFSKLSSVTAADTAVYYCARLPRGDR DAFDIWGQGTMTVTVSSGGGGQGGGGQGGGG QDIVMTQSPSSLSASVGDRVTITCRASQGISNY LAWYQKPKGKVPKLLIYAASLQSGVPSRFSG SGSGTDFTLTISSLQPEDFATYYCQQS YSTPFTF GCGTKVEIKSGGGGQEVQLVESGGGLVQPGGS LKLSCAASGFTFNKYAMNWVRQAPGKGMW VARIRSKYNNYATYYADAVKDRFTISRDDSKN TLYLQMNNLKTEDTAVYYCVRAGNFGSSYISY FAYWGQGT LVTVSSGGGGQGGGGQGGGGQQT TVVTQEPSLTVSPGGTVTITCGSSTGAVTSGNY PNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGS LEGGKAALT LSGVQPEDEAEYYCVLYYSNRW VFGSGTKLTVL
252.	CH3 15-E11 CC x I2L x G4 x scFc x G4 x MS 15-B12	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRSDDTA

	<p>CC x I2L - Full Sequence</p>			<p>VYYCATRYFYVMDYWGQGLTVTVSSGGGGSGGGSGGGGSDIQMTQSPSSLSASVGDRVTITCRASQDISNYLNWYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCVQYAQFPLTFGCGTKVEIKSGGGGSEVQLVESGGGLVQPGGSLKLSAASGFTFNKYAMNWVRQAPGKMEWVARIRSKYNNYATYYADAVKDRFTISRDDSKNTLYLQMNNLKTEDTAVYYCVRAGNFGSSYISYFAYWGQGLTVTVSSGGGGSGGGGSGSQT VVTQEPSLTVSPGGTVTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGSLEGGKAALTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYTQKSLSLSPGKGGGGSGGGSGGGGSDKHTHTCPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYTQKSLSLSPGKGGGGQVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSYFWGWIRQPPGKCLEWIGNIYYSGSSNYPNLSKSRVTISVDTSKNQFSLKLSSVTAADTAVYYCARLPRGDRDAFDIWGQGMVTVSSGGGGSGGGGSGGGGSDIVMTQSPSSLSASVGDRVTITCRASQGISNYLAWYQQKPGKVPKLLIYAASSTLQSGVPSRFSGSGSGTDFTLTISSLQPEDFATYYCQSSYSTPFTFGCGTKVEIKSGGGGSEVQLVESGGGLVQPGGSLKLSAASGFTFNKYAMNWVRQAPGKMEWVARIRSKYNNYATYYADAVKDRFTISRDDSKNTLYLQMNNLKTEDTAVYYCVRAGNFGSSYISYFAYWGQGLTVTVSSGGGGSGGGGSGGSGSQT VVTQEPSLTVSPGGTVTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGSLEGGKAALTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL</p>
<p>253.</p>	<p>CH3 15-E11 CC x I2L x G4S3 x scFc x G4S3 x MS 15-B12 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGSGGGSGGGGSDIQMTQSPSSLSASVGDRVTITCRASQDISNYLNWYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCVQYAQFPLTFGCGTKVEIKSGGGGSEVQLVESGGGLVQPGGSLKLSAASGFTFNKYAMNWVRQAPGKMEWVARIRSKYNNYATYYADAVKDRFTISRDDSKNTLYLQMNNLKTEDTAVYYCVRAGNFGSSYISYFAYWGQGLTVTVSSGGGGSGGGGSGGSGSQT VVTQEPSLTVSPGGTVTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGSLEGGKAALTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL</p>

				<p>GSGGGGSQTVVTQEPSLTVSPGGTVTITCGSST GAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAP GTPARFSGSLEGGKAALTLSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVLGGGGSGGGSSGG GGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNQKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLDSDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGSGGGSSGGGGSGGGSSGGGG SGGGSSDKTHTCPPCPAPELLGGPSVFLFPPK KDTLMISRTPEVTCVVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSTYRCVSVLTVLH QDWLNQKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSDGSFFLYS KLTVDKSRWQQGNVFSCSVMHEALHNHYTQK SLSLSPGKGGGGSGGGSSGGGGSSQVQLQESGP GLVKPSETLSLTCTVSGGSISSSSYFWGWIRQP GKCLEWIGNIYYSGSSNYPNPSLKSRTISVDTS KNQFSLKLSSVTAADTAVYYCARLPRGDRDAF DIWGQGTMTVTVSSGGGGSGGGSSGGGGSDIV MTQSPSSLSASVGDRVTITCRASQGISNYLAWY QQKPGKVPKLLIYAASLTLQSGVPSRFSGSGSGT DFTLTISSLQPEDFATYYCQQSYSTPFTFGCGTK VEIKSGGGGSEVQLVESGGGLVQPGGSLKLSL AASGFTFNKYAMNWVRQAPGKGMWVARIR SKYNNYATYYADAVKDRFTISRDDSKNTLYLQ MNNLKTEDTAVYYCVRAGNFGSSYISYFAYW GQGLVTVVSSGGGGSGGGSSGGGGSSQTVVTQ EPSLTVSPGGTVTITCGSSTGAVTSGNYPNWIQ KKPGQAPRGLIGGTKFLAPGTPARFSGSLEGGK AALTLSGVQPEDEAEYYCVLYYSNRWVFGSG TKLTVL</p>
254.	CH3 15-E11 CC x I2M2 x G4 x scfc x G4 x MS 15-B12 CC x I2M2 - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIAYGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVVSSGGGGSS GGGGSGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSAASGFTFNKYAINWVREA PGKGLEWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTAYLQMNNLKTEDTAVYYCVRNAN FGTSYISYFAYWGQGLTVVSSGGGGSGGGSS GGGGSSQTVVTQEPSLTVSPGGTVTLTICGSSTG AVTSGNYPNWVQKKPGQAPRGLIGGTKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGSGTKLTVLGGGGDKTHTCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VDVSHEDPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNQKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM</p>

				<p>TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSGDGFFLYSKLTVDKSRWQQGNV FSCSVMHEALHNHYTQKSLSLSPGKGGGGSGG GGSGGGGSGGGGSGGGGSGGGGSDKTHTCP CPAPPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VVDVSHEDPEVKFNWYVDGVEVHNAKTKPCE EQYGSTYRCVSVLTVLHQDWLNGKEYKCKVNS NKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSGDGFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGGQ VQLQESGPGLVKPSETLSLTCTVSGGSISSSSYF WGWRQPPGKCLEWIGNIYYSGSSNYPNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRTVITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLVTVSSGGGGSGGGGSGGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLGSGVQPEDEAEYCVLWYS NRWVFGSGTKLTVL</p>
255.	CH3 15-E11 CC x I2M2 x (G4Q)3x scFmod x (G4Q)3 x MS 15- B12 CC x I2M2 - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLVTVSSGGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRTVIT CRASQDISNYLNWYQQKPGKVPKLLIYYTSRL HSGVPSRFSGSGTDFTLTISSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSCAASGFTFNKYAINWVRE APGKLEWVARIRSKYNNYATYYADAVKDRF TISRDDSKNTAYLQMNNLKTEDTAVYYCVRN ANFGTSYISYFAYWGQGLVTVSSGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTLTCGS STGAVTSGNYPNWVQKPKPGQAPRGLIGGTKFL APGTPARFSGSLLGGKAALTLGSGVQPEDEAEY YCVLWYSNRWVFGSGTKLTVLGGGGQGGGG QGGGGQCPCPAPPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEEPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSGDGFFLYSKLTV DKSRWQQGNVFSCSVMHEALHNHYTQKSLSL SPGKGGGGQGGGGQGGGGQGGGGQGGGGQGG GGGQCPCPAPPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEEPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE</p>

				<p>SNGQPENNYKTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSVCSVMHEALHNHYTQKSLSLSPG KGGGGQGGGGQGGGGQVQLQESGPGLVKPS ETLSLTCTVSGGSISSSSYFWGWIRQPPGKCLE WIGNIYSGSSNYPNPSLKSRTISVDTSKNQFSL KLSSVTAADTAVYYCARLPRGDRDAFDIWGQ GTMVTVSSGGGGQGGGGQGGGGQDIVMTQSP SSLSASVGDRVTITCRASQGISNYLAWYQKQP GKVPKLLIYAATLQSGVPSRFSGSGSGTDFTL TISSLQPEDFATYYCQSYSTPFTFGCGTKVEIK SGGGQEVQLVESGGGLVQPGGSLKLSAASG FTFNKYAINWVREAPGKGLEWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTAYLQMNNL KTEDTAVYYCVRNANFGTYSYISYFAYWGQGT LVTVSSGGGGQGGGGQGGGGQQTVVVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQKQP GQAPRGLIGGKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGSGTKL TVL</p>
256.	<p>CH3 15-E11 CC x I2M2 x G4S3 x scFc x G4S3 x MS 15-B12 CC x I2M2 - Full Sequence</p>	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWRVQAPGQCLEWMGNIAYGKGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRSDDTA VYYCATRYFYVMDYWGQGLVTVSSGGGGG GGGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQKPKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSAASGFTFNKYAINWVREA PGKGLEWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTAYLQMNNLKTEDTAVYYCVRNAN FGTYSYISYFAYWGQGLVTVSSGGGGSGGGG GGGGSQTVVTQEPSLTVSPGGTVTLTCGSSTG AVTSGNYPNWVQKPKPGQAPRGLIGGKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGSGTKLTVLGGGGSGGGGSGGG GSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTL MISRTPEVTCVVVDVSHEDPEVKFNWYVDGV EVHNAKTKPCEEQYGSTYRCVSVLTVLHQDW LNGKEYKCKVSNKALPAPIEKTISKAKGQPREP QVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAV EWESNGQPENNYKTPPVLDSDGSFFLYSKL VDKSRWQQGNVFSVCSVMHEALHNHYTQKSL LSPGKGGGGSGGGGSGGGGSGGGGSGGGGSG GGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTPPVLDSDGSFFLYSK LTVDKSRWQQGNVFSVCSVMHEALHNHYTQK LSLSPGKGGGGSGGGGSGGGGSGVQLQESGP LVKPSSETLSLTCTVSGGSISSSSYFWGWIRQPPG KCLEWIGNIYSGSSNYPNPSLKSRTISVDTSK NQFSLKLSSVTAADTAVYYCARLPRGDRDAFD IWGQGTMTVTVSSGGGGSGGGGSGGGGSDIVM TQSPSSLSASVGDRVTITCRASQGISNYLAWYQ</p>

				<p>QKPGKVPKLLIYAASSTLQSGVPSRFSGSGSGTD FTLTISLQPEDFATYYCQQSYSTPFTFGCGTKV EIKSGGGGSEVQLVESGGGLVQPGGSLKLSA ASGFTFNKYAINWVREAPGKGLEWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTAYLQMN NLKTEDTAVYYCVRNANFGTSYISYFAYWGQ GTLVTVSSGGGGSGGGGSGGGGSQT VVTQEPS LTVSPGGTVTLTCGSSTGAVTSGNYPNWVQKK PGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAA LTLSGVQPEDEAEYYCVLWYSNRWVFGSGTK LTVL</p>
257.	<p>CH3 15-E11 CC x I2M2 x G4 x scFc x G4 x MS 15- B12 CC x I2M2 _GQ - Full Sequence</p>	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKV SCKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGT LTVVSSGGGGQ GGGGQGGGQDIQMTQSPSSLSASVGDRVITIT CRASQDISNYLNWYQQKPGKVPKLLIYYSRL HSGVPSRFSGSGSGTDFLTISLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAINWVRE APGKGLEWVARIRSKYNNYATYYADAVKDRF TISRDDSKNTAYLQMNNLKTEDTAVYYCVRN ANFGTSYISYFAYWGQGT LTVVSSGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTLTCGS STGAVTSGNYPNWVQKKPGQAPRGLIGGTKFL APGTPARFSGSLLGGKAA LTLSGVQPEDEAEY YCVLWYSNRWVFGSGTKLTVLGGGGCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEEPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS SCVMHEALHNHYTQKSLSLSPGKGGGGQGG GGQGGGGQGGGGQGGGGQGGGGQCPPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL QESGPGLVKPSSETLSLTCTVSGGSISSSSYFWG WIRQPPGKCLEWIGNIYSGSSNYNPSLKSRTV ISVDTSKNQFSLKLSSVTAADTAVYYCARLPRG DRDAFDIWGQGTMTVTVSSGGGGQGGGGQGG GGQDIVMTQSPSSLSASVGDRVITITCRASQGIS NYLAWYQQKPGKVPKLLIYAASSTLQSGVPSRF SGSGSGTDFLTISLQPEDFATYYCQQSYSTPF TFGCGTKVEIKSGGGGQEVQLVESGGGLVQPG GSLKLSAASGFTFNKYAINWVREAPGKGLEW VARIRSKYNNYATYYADAVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRNANFGTSYIS YFAYWGQGT LTVVSSGGGGQGGGGQGGGGQ QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWVQKKPGQAPRGLIGGTKFLAPGTPARFS</p>

				GSLGGKAALTLSGVQPEDEAEYYCVLWYSNRWVFGSGTKLTVL
258.	CH3 15-E11 CCx I2C 44/100cc x scFc x MS 15-B12 CC x I2C 44/100cc0 - Full Sequence	artificial	Aa	<p> QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGS GGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSAASGFTFNKYAMNWVRQ APGKCLEWVARIRSKYNNYATYYADSVKDRF TISRDDSKNTAYLQMNNLKTEDTAVYYCVRH GNFGNSYISYWAYWGQGLTVTVSSGGGSGG GSGGGGGSQT VVTQEPSLTVSPGGTVTLTCGSS TGAVTSGNYPNWVQKPGQAPRGLIGGTKFL APGTPARFSGSLGGKAALTLSGVQPEDEAEY YCVLWYSNRWVFGCGTKLTVLGGGGSGGGGS GGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPK DTLMISRTPEVTCVVDVSHEDPEVKFNWYVD GVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQ DWLNGKEYKCKVSNKALPAPIEKTISKAKGQP REPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPVLDSDGSFFLYS KLTVDKSRWQQGNVFSCSVMHEALHNHYTQK SLSLSPGKGGGGSGGGGSGGGGSGGGGSGGG GSGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSDGSFFL YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGVQLQES GPGLVKPSSETLSLTCTVSGGSSISSSYFWGWIR QPPGKCLEWIGNIYYSGSSNYPNPSLKSRTISV DTSKNQFSLKLSVTAADTAVYYCARLPRGDR DAFDIWGQGTMTVTVSSGGGGSGGGGSGGGGS DIVMTQSPSSLSASVGDRVTITCRASQGISNYL AWYQQKPGKVPKLLIYAAS TLQSGVPSRFSGS GSGTDFTLTISSLQPEDFATYYCQQSYSTPFTFG CGTKVEIKSGGGGSEVQLVESGGGLVQPGGSL KLSAASGFTFNKYAMNWVRQAPGKCLEWV ARIRSKYNNYATYYADSVKDRFTISRDDSKNT AYLQMNNLKTEDTAVYYCVRHGNFGNSYISY WAYWGQGLTVTVSSGGGSGGGGSGGGGSGG TVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGNY PNWVQKPGQAPRGLIGGTKFLAPGTPARFSG SLLGGKAALTLSGVQPEDEAEYYCVLWYSNR WVFGCGTKLTVL </p>
259.	CH3 24-D7 CC x I2L x G4S3 x scFc x G4S3 x MS 15-B12 CC x I2L - Full Sequence	artificial	Aa	<p> QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIH SKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGS GGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH </p>

				<p>SGVPSRFSGSGSGTDFTLTISLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSCAASGFTFNKYAMNWVRQ APGKGMWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSGGGGSGGG GSGGGGSQTVVVTQEPSLTVSPGGTVTITCGSST GAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAP GTPARFSGSLEGGKAALTLSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVLGGGGSGGGGSGG GGSDKTHTCPAPPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNQKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLDSDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGSGGGGSGGGGSGGGGSGGGG SGGGGSDKTHTCPAPPELLGGPSVFLFPPK KDTLMISRTPEVTCVVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSTYRCVSVLTVLH QDWLNQKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSDGSFFLYS KLTVDKSRWQQGNVFSCSVMHEALHNHYTQK LSLSPGKGGGGSGGGGSGGGGSGVQLQESGP GLVKPSETLSLTCTVSGGSISSSSYFWGWIRQPP GKCLEWIGNIYYSGSSNYNPSLKSRTISVDTS KNQFSLKLSVTAADTAVYYCARLPRGDRDAF DIWGQGMVTVSSGGGGSGGGGSGGGGSDIV MTQSPSSLSASVGDRVTITCRASQGISNYLAWY QQKPGKVPKLLIYAASLQSGVPSRFSGSGSGT DFTLTISLQPEDFATYYCQQSYSTPFTFGCGTK VEIKSGGGGSEVQLVESGGGLVQPGGSLKLSC AASGFTFNKYAMNWVRQAPGKGMWVARIR SKYNNYATYYADAVKDRFTISRDDSKNTLYLQ MNNLKTEDTAVYYCVRAGNFGSSYISYFAYW GQGLTVTVSSGGGGSGGGGSGGGGSGTQVVVTQ EPSLTVSPGGTVTITCGSSTGAVTSGNYPNWIQ KKPGQAPRGLIGGTKFLAPGTPARFSGSLEGG AALTLSGVQPEDEAEYYCVLYYSNRWVFGSG TKLTVL</p>
260.	CH3 24-D7 CC x I2L x G4 x scFc x G4 x MS15-B12 CC x I2L _GQ - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGQ GGGGQGGGQDIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYSRL HSGVPSRFSGSGSGTDFTLTISLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSCAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGLTVTVSSGGGGQGG GGQGGGGQQTVVVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGTKFL</p>

				<p>APGTPARFSGSLEGGKAALTLSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGCPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQGGG GQGGGGQGGGGQGGGGQGGGGQCPPCAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEEPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGQVQLQE SGPGLVKPSETLSLTCTVSGGSISSSSYFWGWIR QPPGKCLEWIGNIYYSGSSNYPNPSLKSRTISV DTSKNQFSLKLSVTAADTAVYYCARLPRGDR DAFDIWGQGTMTVTVSSGGGGQGGGGQGGGG QDIVMTQSPSSLSASVGDRVTITCRASQGISNY LAWYQQKPGKVPKLLIYAASLQSGVPSRFSG SGSGTDFTLTISSLPEDFATYYCQQSSTPFTF GCGTKVEIKSGGGGQEVQLVESGGGLVQPGGS LKLSCAASGFTFNKYAMNWRVQAPGKGMW VARIRSKYNNYATYYADAVKDRFTISRDDSKN TLYLQMNNLKTEDTAVYYCVRAGNFGSSYISY FAYWGQGTLLTVTVSSGGGGQGGGGQGGGGQ TVVTQEPSTLTVSPGGTVTITCGSSTGAVTSGNY PNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGS LEGGKAALTLSGVQPEDEAEYYCVLYYSNRW VFGSGTKLTVL</p>
261.	CH3 24-D7 CC x I2M2 x G4 x scfc x G4 x MS 15-B12 CC x I2M2 - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWRVQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRLLSDDTA VYYCATRYFYVMDYWGQGTLLTVTVSSGGGGS GGGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFTLTISSLPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSCAASGFTFNKYAINWVREA PGKGLEWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTAYLQMNNLKTEDTAVYYCVRNAN FGTSYISYFAYWGQGTLLTVTVSSGGGGSGGGGS GGGGSQTVVTQEPSTLTVSPGGTVTLTCSSTG AVTSGNYPNWVQKKPGQAPRGLIGGTKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGSGTKLTVLGGGGDKTHTCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VDVSHEDPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPEN YKTTTPVLDSDGSFFLYSKLTVDKSRWQQGNV FSCVMHEALHNHYTQKSLSLSPGKGGGGSGG</p>

				<p>GGSGGGGSGGGGSGGGGSGGGGSDKTHTCPP CPAPPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VVDVSHEDPEVKFNWYVDGVEVHNAKTKPCE EQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK NKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSGDSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGQ VQLQESGPGLVKPSETLSLTCTVSGGSISSSSYF WGWIRQPPGKCLEWIGNIYYSGSSNYPNLSKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLGSGVQPEDEAEYCVLWYS NRWVFGSGTKLTVL</p>
262.	CH3 24-D7 CC x G4 x scFc x G4 x MS 15-B12 CC x I2M2_GQ - Full Sequence	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRVIT CRASQDISNYLNWYQQKPGKVPKLLIYYSRL HSGVPSRFSGSGSGTDFLTITSSSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSCAASGFTFNKYAINWVRE APGKGLEWVARIRSKYNNYATYYADAVKDRF TISRDDSKNTAYLQMNNLKTEDTAVYYCVRN ANFGTSYISYFAYWGQGLTVTVSSGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTLTCGS STGAVTSGNYPNWVQKPKPGQAPRGLIGGTKFL APGTPARFSGSLLGGKAALTLGSGVQPEDEAEY YCVLWYSNRWVFGSGTKLTVLGGGGCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEEPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTTTPVLDSGDSFFLYSKLTVDKSRWQQGNV SCSVMHEALHNHYTQKSLSLSPGKGGGGQGG GGQGGGGQGGGGQGGGGQGGGGQCPPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPVLDSGDSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL QESGPGLVKPSETLSLTCTVSGGSISSSSYFWG</p>

				<p>WIRQPPGKCLEWIGNIYYSGSSNYPNPSLKSRTV ISVDTSKNQFSLKLSSVTAADTAVYYCARLPRG DRDAFDIWGQGMVTVSSGGGGQGGGGQGG GGQDIVMTQSPSSLSASVGDRVTITCRASQGIS NYLAWYQQKPGKVPKLLIYAASTLQSGVPSRF SGSGSGTDFTLTISLQPEDFATYYCQQSYSTPF TFGCGTKVEIKSGGGGQEVQLVESGGGLVQPG GSLKLSAASGFTFNKYAINWVREAPGKGLEW VARIRSKYNNYATYYADAVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRNANFGTSYIS YFAYWGQGLTVTVSSGGGGQGGGGQGGGGQ QTVVTVQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWWQKKPGQAPRGLIGGKFLAPGTPARFS GSLGKAALTLGSGVQPEDEAEYCVLWYSN RWVFGSGTKLTVL</p>
<p>263.</p>	<p>CH3 24-D7 CC x I2L x (G4Q)3x scFmod x (G4Q)3 x MS 15- B12 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYSRL HSGVPSRFSGSGTDFTLTISLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGLTVTVSSGGGGQGG GGGGGGQTVVTVQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGKFL APGTPARFSGSLEGGKAALTLGSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGQGGGGQ GGGGQCPPCPAPPELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEEPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSDGSFFLYSKLTV DKSRWQQGNVFCSSVMHEALHNHYTQKSLSL SPGKGGGGQGGGGQGGGGQGGGGQGGGGQGG GGGQCPPCPAPPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEEPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSSVMHEALHNHYTQKSLSLSPG KGGGGQGGGGQGGGGQVQLQESGPGLVKPS ETLSLTCTVSGGSISSSSYFWGWIRQPPGKCLE WIGNIYYSGSSNYPNPSLKSRTVISVDTSKNQFSL KLSSVTAADTAVYYCARLPRGDRDAFDIWGQ GTMVTVSSGGGGQGGGGQGGGGQDIVMTQSP SSLSASVGDRVTITCRASQGISNYLAWYQQK GKVPKLLIYAASTLQSGVPSRFSGSGSGTDFTL TISSLQPEDFATYYCQQSYSTPFTFGCGTKVEIK SGGGGQEVQLVESGGGLVQPGGSLKLSAASG FTFNKYAMNWVRQAPGKGMWVARIRSKYN</p>

				<p>NYATYYADAVKDRFTISRDDSKNTLYLQMNN LKTEDTAVYYCVRAGNFGSSYISYFAYWGQGT LVTVSSGGGGQGGGGQGGGGQQTVVVTQEPSL TVSPGGTVTITCGSSTGAVTSGNYPNWIQKPG QAPRGLIGGTKFLAPGTPARFSGSLEGGKAALT LSGVQPEDEAEYYCVLYYSNRWVFGSGTKLT VL</p>
264.	<p>CH3 24-D7 CC x I2L x G4 x scFc x G4 x MS 15-B12 CC x I2L - Full Sequence</p>	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRLSDDTA VYYCATRYFYVMDYWGGQGLTVTVSSGGGGG GGGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFLTISLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSAASGFTFNKYAMNWVRQ APGKGMWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSGGGGSGGG GSGGGGSQTVVTQEPSLTVSPGGTVTITCGSST GAVTSGNYPNWIQKPGQAPRGLIGGTKFLAP GTPARFSGSLEGGKAALTLSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVLGGGGDKTHTCPP CPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VVDVSHEDPEVKFNWYVDGVEVHNAKTKPCE EQYGSTYRCVSVLTVLHQDWLNGKEYKCKV NKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSGDFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGG GGGSGGGGSGGGGSGGGGSGGGGSDKTHTCP PCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTC VVDVSHEDPEVKFNWYVDGVEVHNAKTKPC EEQYGSTYRCVSVLTVLHQDWLNGKEYKCKV SNKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSGDFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGQ VQLQESGPGLVKPSSETLSLTCTVSGGSISSSSYF GWIRQPPGKCLEWIGNIYYSGSSNYPNLSKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGGSGGGG SGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSQTVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVL</p>
265.	<p>CH3 24-D7 CC x I2M2 x (G4Q)3x</p>	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIHKAHGTNY</p>

	<p>scF_{cm}od x (G4Q)₃ x MS 15- B12 CC x I2M2 - Full Sequence</p>			<p>NQKFQGRVTMTVDTSSSTAYMELSRRLRSDDTA VYYCATRYFYVMDYWGQGLVTVSSGGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSCAASGFTFNKYAINWVRE APGKGLEWVARIRSKYNNYATYYADAVKDRF TISRDDSKNTAYLQMNNLKTEDTAVYYCVRN ANFGTSYISYFAYWGQGLVTVSSGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTLTCGS STGAVTSGNYPNWVQKKPGQAPRGLIGGTKFL APGTPARFSGSLLGGKAALTLSGVQPEDEAEY YCVLWYSNRWVFGSGTKLTVLGGGGQGGGG QGGGGQCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEEPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSDGSFFLYSKLTV DKSRWQQGNVFCSCVMHEALHNHYTQKSLSL SPGKGGGGQGGGGQGGGGQGGGGQGGGGQGG GGGQCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEEPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG KGGGGQGGGGQGGGGQVQLQESGPGLVKPS ETLSLTCTVSGGSISSSYFWGWIRQPPGKCLE WIGNIYYSGSSNYPNPSLKSRTISVDTSKNQFSL KLSSVTAADTAVYYCARLPRGDRDAFDIWGQ GTMVTVSSGGGGQGGGGQGGGGQDIVMTQSP SSLSASVGDRVTITCRASQGISNYLAWYQQK GKVPKLLIYAASLQSGVPSRFSGSGSGTDFTL TISSLQPEDFATYYCQQSYPFTFGCGTKVEIK SGGGGQEVQLVESGGGLVQPGGSLKLSCAASG FTFNKYAINWVREAPGKGLEWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTAYLQMNNL KTEDTAVYYCVRNANFGTSYISYFAYWGQGL LTVSSGGGGQGGGGQGGGGQQT VVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQKKP GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGSGTKL TVL</p>
<p>266.</p>	<p>CH3 24-D7 CC x I2M2 x G4S3 x scFc x G4S3 x MS 15-B12 CC x I2M2 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLVQSGAEVKKPGASVKVSKCASGYFTFN YWMNWVRQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRRLRSDDTA VYYCATRYFYVMDYWGQGLVTVSSGGGGGS GGGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSCAASGFTFNKYAINWVREA PGKGLEWVARIRSKYNNYATYYADAVKDRFTI</p>

				<p>SRDDSKNTAYLQMNNLKTEDTAVYYCVRNAN FGTSYISYFAYWGQGLTVTVSSGGGGSGGGGS GGGGSQTVVTQEPSLTVSPGGTVTLTCGSSTG AVTSGNYPNWVQKKPGQAPRGLIGGTKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGSGTKLTVLGGGGSGGGGSGGG GSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTL MISRTPEVTCVVVDVSHEDPEVKFNWYVDGV EVHNAKTKPCEEQYGSTYRCVSVLTVLHQDW LNGKEYKCKVSNKALPAPIEKTISKAKGQPREP QVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAV EWESNGQPENNYKTTPVLDSGDFFLYSKLT VDKSRWQQGNVFCFSVMHEALHNHYTQKSLS LSPGKGGGGSGGGGSGGGGSGGGGSGGGGSG GGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPVLDSGDFFLYSK LTVDKSRWQQGNVFCFSVMHEALHNHYTQKS LSLSPGKGGGGSGGGGSGGGGSGVQLQESGPG LVKPSETLSLTCTVSGGSISSSSYFWGWIRQPPG KCLEWIGNIYYSGSSNYNPSLKSRTVISVDTSK NQFSLKLSSVTAADTAVYYCARLPRGDRDAFD IWGQGTMTVTVSSGGGGSGGGGSGGGGSDIVM TQSPSSLSASVGDRVTITCRASQGISNYLAWYQ QKPGKVPKLLIYAASLQSGVPSRFSGSGSGTD FTLTISSLQPEDFATYYCQQSYSTPFTFGCGTKV EIKSGGGGSEVQLVESGGGLVQPGGSLKLSCA ASGFTFNKYAINWVREAPGKGLEWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTAYLQMN NLKTEDTAVYYCVRNANFGTSYISYFAYWGQ GTLTVTVSSGGGGSGGGGSGGGGSGGGGSGGGG LTVSPGGTVTLTCGSSTGAVTSGNYPNWVQKK PGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAA LTLSGVQPEDEAEYYCVLWYSNRWVFGSGTK LTVL</p>
267.	<p>CH3 24-D7 CCx 6H10.09x (G4S)3x scFcx (G4S)3x MS R4L CCx 6H10.09 - Full Sequence</p>	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSKKASGYTFTN YWMNWVRQAPGQCLEWMGNIHKAHGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGGS GGGGSGGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYTSRLH SGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSCAASGFTFNKYAMNWVRQ APGKGMWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSGGGGSGGG GSGGGGSGGGGSDIQMTQSPSSLSASVGDRVTITC GAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAP GTPARFSGSLEGGKAALTLSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVLGGGGSGGGGSGGG GGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG</p>

				<p>VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLDSGDSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGSGGGGSGGGGSGGGGSGGGG SGGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPK KDTLMISRTPEVTCVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSTYRCVSVLTVLH QDWLNKKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSGDSFFLYS KLTVDKSRWQQGNVFSCSVMHEALHNHYTQK LSLSPGKGGGSGGGGSGGGGSGVQLVQSGA EVKKPGASVKVSCKASGYTFTGYYIHWRQA PGQCLEWMGWINPNSGGTNYAQKFQGRVTMT RDTISISTAYMELSRLSDDTAVYYCARVEAVA GREYYYFSGMDVWGQGTTVTVSSGGGGSGGG GSGGGGSSYELTQPPSVSVSPGQTASITCSGEK LGDKYVYWYQQKPGQSPVLVIYQSTKRPSGVP ERFSGSNSGNTATLTISGTQAMDEADYYCQAY HASTAVFGCGTKLTVLSGGGGSEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKGMEWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTLYLQMNNLKTEDTAVYYCVRAGN FGSSYISYFAYWGQGTTLTVVSSGGGGSGGGGS GGGGSQTVVTQEPSLTVSPGGTVTITCGSSTGA VTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGT PARFSGSLEGGKAALTLSGVQPEDEAEYYCVL YYSNRWVFGSGTKLTVL</p>
268.	<p>CH3 R164L CC x I2Ccc(44/100)x (G4S)3x scFcx (G4S)3x MS R4L CCx I2Ccc(44/100) - Full Sequence</p>	artificial	aa	<p>EVQLLESGLVQPGGSLVLSCAASGFTFSSY WMYWVRQAPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDNSKNTLYLQMNSLRAEDSAVY YCARWDYTHFDVWGQGTTVTVSSGGGGSGG GGSGGGSEIVMTQSPATLSVSPGERATLSCRA SSVSVMHWYQQKPGQAPRLLIYGTSNLSGV PVRFGSGSGTEFTLTISRLQSEDAVYYCQQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGTTLTVVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGTVTITCGSSTGA VTSGNYPNWWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLGGGGSGGGGSGGG GSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTL MISRTPEVTCVVDVSHEDPEVKFNWYVDGVE VHNAKTKPCEEQYGSTYRCVSVLTVLHQDW LNGKEYKCKVSNKALPAPIEKTISKAKGQPREP QVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAV EWESNGQPENNYKTTPPVLDSGDSFFLYSKLT VDKSRWQQGNVFSCSVMHEALHNHYTQKSLS LSPGKGGGSGGGGSGGGGSGGGGSGGGGSG GGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVDVSHEDPEVKFNWYVDG</p>

				<p>VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLDSGDSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGSGGGGSGGGGSQVQLVQSGA EVKKPGASVKVSCKASGYTFTGYYIHWVRQA PGQCLEWMGWINPNSGGTNYAQKFQGRVTMT RDTISISTAYMELSRRLSDDTAVYYCARVEAVA GREYYYFSGMDVWGQGTITVTVSSGGGSGGGG GSGGGGSSYELTQPPSVSVSPGQTASITCSGEK LGDKYVYWYQQKPGQSPVLVIYQSTKRPSGVP ERFSGSNSGNTATLTISGTQAMDEADYYCQAY HASTAVFGCGTKLTVLSGGGGSEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKCLEWVARIRSKYNNYATYYADSVKDRFTIS RDDSKNTAYLQMNNLKTEDTAVYYCVRHGNF GNSYISYWAYWGQGLTVTVSSGGGSGGGGGS GGGGSQTVVTQEPSLTVSPGGTITLTCGSSTG AVTSGNYPNWVQKPGQAPRGLIGGTKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGCGTKLTVL</p>
269.	<p>CH3 R164L CCx 6H10.09x (G4S)3x scFcx (G4S)3x MS R4L CCx 6H10.09 - Full Sequence</p>	artificial	aa	<p>EVQLLESGLVQPGGSLVRLSCAASGFTFSSY WMYWVRQAPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDNSKNTLYLQMNSLRAEDSAVY YCARWDYTHFDVWGQGTITVTVSSGGGSGGG GGSGGGGSEIVMTQSPATLSVSPGERATLSCRA SSSVSYMHYQQKPGQAPRLLIYGTSNLASGV PVRFGSGSGTEFTLTISRLQSEDAVYYCQQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KMEWVARIRSKYNNYATYYADAVKDRFTIS RDDSKNTLYLQMNNLKTEDTAVYYCVRAGNF GSSYISYFAYWGQGLTVTVSSGGGSGGGGSG GGGGSQTVVTQEPSLTVSPGGTITITCGSSTGAV TSGNYPNWIQKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAALTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGSGGGGSGGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDNLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSGDSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGSGGGGSGGGGSGGGGSGGGGSGGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDNLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSGDSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGSGGGGSGGGGSGGGGSGGGGSGGGGGS ASVKVSCKASGYTFTGYYIHWVRQAPGQCLE WMGWINPNSGGTNYAQKFQGRVTMTRDTSIS</p>

				<p>TAYMELSRRLSDDTAVYYCARVEAVAGREYY YFSGMDVWGQGTTVTSSGGGGSGGGGSGGG GSSYELTQPPSVSVSPGQTASITCSGEKLGDKY VYWYQQKPGQSPVLVIYQSTKRPSGVPERFSG SNSGNTATLTISGTQAMDEADYYCQAYHASTA VFGCGTKLTVLSGGGGSEVQLVESGGGLVQPG GSLKLSAASGFTFNKYAMNWVRQAPGKGM WVARIRSKYNNYATYYADAVKDRFTISRDDSK NTLYLQMNNLKTEDTAVYYCVRAGNFGSSYIS YFAYWGQGTTLTVSSGGGGSGGGGSGGGGSSQ TVVTQEPLSLTVSPGGTVTITCGSSTGAVTSGNY PNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGS LEGGKAALTLSGVQPEDEAEYYCVLYYSNRW VFGSGTKLTVL</p>
<p>270.</p>	<p>CH3 R164L CCx I2Ccc(44/100)x (G4)x scFc x (G4) x MS R4L CCx I2Ccc(44/100) - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>EVQLLESAGGGLVQPGGSRVLSAASGFTFSSY WMYWVRQAPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDNSKNTLYLQMNLSRAEDSAVY YCARWDYTHFDVWGQGTTVTSSGGGGSGG GGGGGGSEIVMTQSPATLSVSPGERATLSCRA SSSVSYMHWYQQKPGQAPRLLIYGTSNLASGV PVRFSGSGSGTEFTLTISRLEQSEDVAVYYCQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGTTLTVSSGGGGSGGGGSG GGGSTVVTQEPLSLTVSPGGTVTLTCGSSTGA VTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLGGGGDKTHTCPPCP APELLGGPSVFLFPPKPKDTLMISRTPEVTCV VDVSHEDPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSDGSFFLYSKLTVDKSRWQQGNV FSCSVMHEALHNHYTQKSLSLSPGKGGGGSGG GGGGGGGGGGGGGGGGGGGGGGGGSDKHTCPP CPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VDVSHEDPEVKFNWYVDGVEVHNAKTKPCE EQYGSTYRCVSVLTVLHQDWLNGKEYKCKVS NKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSDGSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGQ VQLVQSGAEVKKPGASVKVSCKASGYTFTGY YIHWVRQAPGQCLEWMGWNPNSGGTNYAQ KFQGRVTMTRDTSISTAYMELSRRLSDDTAVY YCARVEAVAGREYYYFSGMDVWGQGTTVTSS SGGGGSGGGGGGGGGGGSSYELTQPPSVSVSPG QTASITCSGEKLGDKYVYWYQQKPGQSPVLVI YQSTKRPSGVPERFSGSNSGNTATLTISGTQAM DEADYYCQAYHASTAVFGCGTKLTVLSGGGG SEVQLVESGGGLVQPGGSLKLSAASGFTFNK YAMNWVRQAPGKCLEWVARIRSKYNNYATY YADSVKDRFTISRDDSKNTAYLQMNNLKTEDT</p>

				AVYYCVRHGNFGNSYISYWAYWGQGLTVTVS SGGGGSGGGGSGGGGSGGGGSGGGGSGGGG TVTLTCGSSTGAVTSGNYPNWVQQKPGQAPR GLIGGTKFLAPGTPARFSGSLLGGKAALTLSGV QPEDEAEYYCVLWYSNRWVFGCGTKLTVL
271.	CH3 R170R CC x I2C 44/100cc x scFc x MS R4L CC x I2C 44/100cc0 - Full Sequence	artificial	Aa	EVQLLESGLLVQPGGSLRLSCAASGFTFSSY WMHWVRQTPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDKSKNTLYLQMNSLRAEDTAVY YCARWDYSHFDVWGQGTTVTVSSGGGGSGG GGSGGGGSEIVMTQSPATLSVSPGERATLTCRA SSVSVMHWYQQKPGQAPRLLIYGTSNLVSGV PARFSGSGSGTEFTLTISSLQSEDFAVYYCQQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGLTVTVSSGGGGSGGGGSG GGGSGTQVVTQEPSLTVSPGGTTLTCGSSTGA VTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLGGGGSGGGGSGGG GSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTL MISRTPEVTCVVVDVSHEDPEVKFNWYVDGV EVHNAKTKPCEEQYGSTYRCVSVLTVLHQDW LNGKEYKCKVSNKALPAPIEKTISKAKGQPREP QVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAV EWESNGQPENNYKTTTPVLDSDGSFFLYSKLT VDKSRWQQGNVFSCSVMHEALHNHYTQKSLS LSPGKGGGGSGGGGSGGGGSGGGGSGGGGSG GGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGSGGGGSGGGGSGVQLVQSGA EVKKPGASVKVSCKASGYTFTGYYIHWRQA PGQCLEWMGWINPNSGGTNYAQKFQGRVTMT RDTISISTAYMELSLRSDDTAVYYCARVEAVA GREYYYFSGMDVWGQGTTVTVSSGGGGSGGG GSGGGGSSYELTQPPSVSVSPGQTASITCSGEK LGDKYVYWYQQKPGQSPVLVIYQSTKRPSGVP ERFSGSNSGNTATLTISGTQAMDEADYYCQAY HASTAVFGCGTKLTVLSGGGGSEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKCLEWVARIRSKYNNYATYYADSVKDRFTISR RDDSKNTAYLQMNNLKTEDTAVYYCVRHGNF GNSYISYWAYWGQGLTVTVSSGGGGSGGGGSG GGGSGTQVVTQEPSLTVSPGGTTLTCGSSTG AVTSGNYPNWVQQKPGQAPRGLIGGTKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGCGTKLTVL
272.	MS 01-G11 CCx 6H10.09x (G4S)3x scFc	artificial	Aa	QVQLVESGGGLVQPGGSLRLSCAASGFTFSDY YMTWIRQAPGKCLEWLSYISSSGSTIYYAEAV KGRFTISRDNKNSLFLQMNSLRAEDTAVYYC

	(G4S)3x CH3 005-D5 CCx 6H10.09 - Full Sequence			<p>ARDRNSHFDYWGGQTLVTVSSGGGGSGGGGS GGGGSDIMTQSPSSVSASVGDRVTITCRASQGI RTWLAWYQQKPGKAPKLLIYGASGLQSGVPS RFSGSGSGTDFTLTISSLQPEDFATYYCQQAESF PRTFGCGTKVEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTITCGSSTGAVTS GNYPNWIQKKPGQAPRGLIGGKFLAPGTPAR FSGSLEGGKAALTLSGVQPEDEAEYYCVLYYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSEVQLLES GGGLVQPG GSLRLSCAASGFSFSSYPINWVRQAPGKCLEW VGVIWTGGGTNYASSVKGRFTISRDN SKNTVY LQMNSLRAEDTAVYYCAKSRGVYDFKGRGA MDYWGQGLTVTVSSGGGGSGGGGSGGGGSDI VMTQSPDSLAVSLGERATINCKSSQSLLYSSNQ KNYFAWYQQKPGQPPLLIYWASTRESGVPDR FSGSGSGTDFTLTISSLQAEDVAVYYCQYYYSY PYTFGCGTKLEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTITCGSSTGAVTS GNYPNWIQKKPGQAPRGLIGGKFLAPGTPAR FSGSLEGGKAALTLSGVQPEDEAEYYCVLYYS NRWVFGSGTKLTVL</p>
273.	MS 15-B12 CC x I2L x (G4Q)3 x scFc x (G4Q)3 x CH3 22- A12 CC x I2L - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAAS TLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG</p>

				<p>CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTPPVLDSDGSFFLYSKLTVDKSRW QQGNVFSCSVMHEALHNHYTQKSLSLSPGKG GGGQGGGGQGGGGQGGGGQGGGGQGGGGQ CPPCPAPPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEEPEVKFNWYVDGVEVHNAKTK PCEEQYGSTYRCVSVLTVLHQDWLNGKEYKC KVSNKALPAPIEKTISKAKGQPREPQVYTLPPS REEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ PENNYKTPPVLDSDGSFFLYSKLTVDKSRWQ QQGNVFSCSVMHEALHNHYTQKSLSLSPGKGG GGQGGGGQGGGGQVQLVQSGAEVKKPGAS VKVSCKASGYTFTNYYWMNWVRQAPGQCLEW MGNIAYGVKGTNYNQQKFQGRVTMTVDTSSST AYMELSRLRSDDTAVYYCATRYFYVMDYWG QGTLVTVSSGGGGQGGGGQGGGGQDIQMTQS PSSLSASVGDRTITCRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGQEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKGMWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ GTLVTVSSGGGGQGGGGQGGGGQQTIVTQEP SLTVSPGGTVTITCGSSTGAVTSGNYPNWIQKK PGQAPRGLIGGTKFLAPGTPARFSGSLEGGKAA LTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKL TVL</p>
275.	MS 15-B12 CC x I2L x G4 x scFc xG4 x CH3 26-E5 CC x I2L_GQ - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTISLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTLVTVSSGGGGQGGGGQGG GGGQQTIVTQEPSTVSPGGTVTITCGSSTGAV TSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAA LTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGCPPCPAPPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEE PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTPPVLD SDGSFFLYSKLTVDKSRWQQGNVFSCSVMHE ALHNHYTQKSLSLSPGKGGGGQGGGGQGGGG QGGGGQGGGGQGGGGQCPPCPAPPELLGGPSV FLFPPKPKDTLMISRTPEVTCVVVDVSHEEPEV KFNWYVDGVEVHNAKTKPCEEQYGSTYRCVS VLTVLHQDWLNGKEYCKVSNKALPAPIEKTI</p>

				NQKNYFAWYQQKPGQPPKLLIYWASTRESGV PDRFSGSGSFTDFTLTISSSLQAEDVAVYYCQQY YSYPYTFGCGTKLEIKSGGGGQEVQLVESGGG LVQPGGSLKLSAASGFTFNKYAMNWVRQAP GKGMWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTLYLQMNNLKTEDTAVYYCVRAGN FGSSYISYFAYWGQGLTVTVSSGGGGQGGGGQ GGGGQQT VVTQEPLTVSPGGT VTITCGSSTGA VTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGT PARFSGSLEGGKAAL T LSGVQPEDEAEYYCVL YYSNRWVFGSGTKLTVL
277.	MS 15-B12 CC x I2L x G4 x scFc xG4 x CH3 15- E11 CC x I2L _GQ - Full Sequence	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSFTDFTLTISSSLQPEDFATYYCQSSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGQGGGGQ GGGQQT VVTQEPLTVSPGGT VTITCGSSTGAV TSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAAL T LSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGCPCPAPPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEE PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHE ALHNHYTQKSLSLSPGKGGGGQGGGGQGGGG QGGGGQGGGGQGGGGQCPAPPELLGGPSV FLFPPKPKDTLMISRTPEVTCVVVDVSHEEPEV KFNWYVDGVEVHNAKTKPCEEQYGSTYRCVS VLTVLHQDWLNGKEYKCKVSNKALPAPIEKTI SKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTPPVLD DGSFFLYSKLTVDKSRWQQGNVFCFSVMHEA LHNHYTQKSLSLSPGKGGGGQVQLVQSGAEV KKPGASVKVSCKASGYTFTNYYWMNWVRQAP GQCLEWMGNIA YGVKGTNYNQKFQGRVTMT VDTSSSTAYMELSRLRSDDTAVYYCATRYFYV MDYWGQGLTVTVSSGGGGQGGGGQGGGGQD IQMTQSPSSLSASVGDRVTITCRASQDISNYLN WYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGS GTDFTLTISSSLQPEDVATYYCVQYAQFPLTFGC GTKVEIKSGGGGQEVQLVESGGGLVQPGGSLK LSAASGFTFNKYAMNWVRQAPGKGMWVVA RIRSKYNNYATYYADAVKDRFTISRDDSKNTL YLQMNNLKTEDTAVYYCVRAGNFGSSYISYFA YWGQGLTVTVSSGGGGQGGGGQGGGGQQT V VTQEPLTVSPGGT VTITCGSSTGAVTSGNYPN WIQKKPGQAPRGLIGGTKFLAPGTPARFSGSLE

				GGKAALTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL
278.	MS 15-B12 CC x I2L x G4S3 x scFc x G4S3 x CH3 26-E5 CC x I2L - Full Sequence	artificial	Aa	<p> QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGGTDFLTITSSLPEDFATYYCQSSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGLSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNLLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLVTVSSGGGGSGGGGSGG GGSQTVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKPGQAPRGLIGGTFKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTPPVLDSDGSFFLYSKLTVDK RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTPPVLDSDGSFFLYSKLTVDK RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTSYWMHWVRQAPGQCLE WMGVIRTSTSYTIYNQKFKGRVTMTRDTSTST VYMESSLRSEDVAVYYCARSGPGYFDVWGQ GTMTVTVSSGGGGSGGGGSGGGGSDIQLTQSPS FLSASVGDRTITCRASENIYSYLAWYQQKPG KAPKLLIYNAKTLAEGVPSRFSGSGSGTEFTLTI SSLQPEDFATYYCQHNYGTPYTFGCGTKLEIKS GGGGSEVQLVESGGGLVQPGLSLKLSCAASGFT FNKYAMNWVRQAPGKMEWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTLYLQMNLL KTEDTAVYYCVRAGNFGSSYISYFAYWGQGLV TVTVSSGGGGSGGGGSGGGGSGTQVVTQEPSLTV SPGGTVTITCGSSTGAVTSGNYPNWIQKPGQ APRGLIGGTFKFLAPGTPARFSGSLEGGKAALTL SGVQPEDEAEYYCVLYYSNRWVFGSGTKLTV L </p>
279.	MS 15-B12 CC x I2Lx G4S3 x scFc x G4S3 x CH3 24-D7 CC x I2L - Full Sequence	artificial	aa	<p> QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP </p>

				<p>SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSQTVVTQEPLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIHSKAHGTNYNQKFQGRVTMTVDTSS STAYMELSRLSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKMEWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ GTLTVTVSSGGGGSGGGGSGGGGSGTQVVTQEPS LTVSPGGTVTITCGSSTGAVTSGNYPNWIQKKP GQAPRGLIGGTKFLAPGTPARFSGSLEGGKAAL TLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL</p>
280.	MS 15-B12 CC x I2M2 x G4 x scfc x G4 x CH3 15-E11 CC xI2M2 - Full Sequence	artificial	aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGGG GSQTVVTQEPLTVSPGGTVTITCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR</p>

				<p>FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGDKTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGSGGGGS GGGGSGGGSGGGSGGGSGGGSDKTHTCPPCPAP ELGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL VQSGAEVKKPGASVKVCKASGYTFTNYWMN WVRQAPGQCLEWMGNIAYGKGTNYNQKFQ GRVTMTVDTSSSTAYMELSRRLSDDTAVYYC ATRYFYVMDYWGQGLTVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSLSASVGDRVTITCRASQD ISNYLNWYQQKPGKVPKLLIYYTSRLHSGVPSR FSGSGGTDFTLTISSLQPEDVATYYCVQYAQF PLTFGCGTKVEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSCAASGFTFNKYAINWVREAPGKGL EWWARIRSKYNNYATYYADAVKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRNANFGTSY ISYFAYWGQGLTVTVSSGGGGSGGGSGGGGS QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWVQKKPGQAPRGLIGGTKFLAPGTPARFS GSLGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGSGTKLTVL</p>
281.	MS 15-B12 CC x I2M2 x (G4Q)3x scFcmo d x (G4Q)3 x CH3 22- A12 CC x I2M2_GQ - Full Sequence	artificial	aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGGTDFTLTISSLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGQGGGGQGGG GQQT VVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGQGGGGQGGGGQC PPCPAPELGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHPEPEVKFNWYVDGVEVHNAKTKP CEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDSDGSFFLYSKLTVDKSRWQQG NVFSCSVMHEALHNHYTQKSLSLSPGKGGG</p>

				<p>QGGGGQGGGGQGGGGQGGGGQGGGGQCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVV VDVSHEEPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSGDGSFFLYSKLTVDKSRWQQGNV FSCSVMHEALHNHYTQKSLSLSPGKGGGGQ GGGQGGGGQVQLVQSGAEVVKPGASVKVSC KASGYTFTSSWMNWRVQAPGQCLEWMGRIYT GTGETKYSKGFQGRVTITRDTASTAYMELSSL TSEDVAVYYCARQRDYGALYAMDYWGQGT LTVSSGGGGQGGGGQGGGGQDIQLTQSPSFLS ASVGDRVTITCRASDDIYSYLAWYQQKPKGAP KLLVYNAKTLAEGVPSRFSGSGSGTEFTLTISL QPEDFATYYCQNHDRTPFTFGCGTKVDIKSGG GGQEVQLVESGGGLVQPGGSLKLSCAASGFTF NKYAINWVREAPGKGLEWVARIRSKYNNYAT YYADAVKDRFTISRDDSKNTAYLQMNNLKTE DTAVYYCVRNANFGTSYISYFAYWGQGT LVTVSSGGGGQGGGGQGGGGQQT VVTQEPSLTVS PGGTVTLTCGSSTGAVTSGNYPNWVQKPKGQ APRGLIGGTKFLAPGTPARFSGSLLGGKAALTL SGVQPEDEAEYYCVLWYSNRWVFGSGTKLTV L</p>
282.	MS 15-B12 CC x I2M2 x (G4Q)3x scFemod x (G4Q3) x CH3 15-E11 CC x I2M2 - Full Sequence	artificial	aa	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPKGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTISSLQPEDFATYYCQSSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGK LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGT LTVTVSSGGGGQGGGGQGGG GQQT VVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPKGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTL SGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGQGGGGQGGGGQC PPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHEEPEVKFNWYVDGVEVHNAKTKP CEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDSGDGSFFLYSKLTVDKSRWQQG NVFSCSVMHEALHNHYTQKSLSLSPGKGGGG QGGGGQGGGGQGGGGQGGGGQGGGGQCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVV VDVSHEEPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSGDGSFFLYSKLTVDKSRWQQGNV FSCSVMHEALHNHYTQKSLSLSPGKGGGGQ</p>

				<p>GGGQGGGGQQVQLVQSGAEVKKPGASVKVSC KASGYTFTNYWMNWVRQAPGQCLEWMGNIA YGVKGTNYNQKFQGRVTMTVDTSSSTAYMEL SRLRSDDTAVYYCATRYFYVMDYWGQGLVT VSSGGGGQGGGGQGGGGQDIQMTQSPSSLSAS VGDRVTITCRASQDISNYLNWYQQKPGKVPKL LIYYTSRLHSGVPSRFSGSGSGTDFTLTISSLQPE DVATYYCVQYAQFPLTFGCGTKVEIKSGGGGQ EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AINWVREAPGKGLEWVARIRSKYNNYATYYA DAVKDRFTISRDDSKNTAYLQMNNLKTEDTA VYYCVRNANFGTSYISYFAYWGQGLVTVSSG GGGQGGGGQGGGGQQT VVTQEP SLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQKKPGQAPRG LIGGTKFLAPGTPARFSGSLLGGKAALTLSGVQ PEDEAEYYCVLWYSNRWVFGSGTKLTVL</p>
<p>283.</p>	<p>MS 15-B12 CC x I2M2 x G4 x scFc x G4 x CH3 005- D5 CC x I2M2_GQ - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGK LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLVTVSSGGGGQGGGGQGGGG GQQT VVTQEP SLTVSPGGT VTLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGCPCPAPELLGGPS VFLFPPKPKDTLMISRTPEVTCVVVDVSHEEPE VKFNWYVDGVEVHNAKTKPCEEQYGSTYRCV SVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQGNVFCSSVMHEA LHNHYTQKSLSLSPGKGGGGQGGGGQGGGGQ GGGGQGGGGQGGGGQCPCPAPELLGGPSVFL FPPKPKDTLMISRTPEVTCVVVDVSHEEPEVKF NWYVDGVEVHNAKTKPCEEQYGSTYRCVSVL TVLHQDWLNGKEYKCKVSNKALPAPIEKTISK AKGQPREPQVYTLPPSREEMTKNQVSLTCLVK GFYPSDIAVEWESNGQPENNYKTTTPVLDSG SFFLYSKLTVDKSRWQQGNVFCSSVMHEALH NHYTQKSLSLSPGKGGGGEVQLLES GGGLVQP GGSLRLSCAASGFSFSSYPINWVRQAPGKCLE WVGVIWTGGGTNYASSVKGRFTISRDN SKNTV YLQMNSLRAEDTAVYYCAKSRGVYDFKGRGA MDYWGQGLVTVSSGGGGQGGGGQGGGGQD IVMTQSPDSLAVSLGERATINCKSSQSLLYSSN QKNYFAWYQQKPGQP KLLIYWASTRESGVP DRFSGSGSGTDFTLTISSLQAEDVAVYYCQQY YSYPYTFGCGTKLEIKSGGGGQEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAINWVREAPG</p>

				KGLEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRNANFG TSYISYFAYWGQGTLLTVSSGGGGQGGGGQ GGGQQTVVVTQEPLSLTVSPGGTVTLTCGSSTGA VTSGNYPNWVQKKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGSGTKLTVL
284.	MS 15-B12 CC x I2M2 x G4 x scFc x G4 x CH3 22- A12 CC x I2M2_GQ - Full Sequence	artificial	aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQKPKGKVPKLLIYAASLTQSGVP SRFSGSGSGLTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGTLLTVSSGGGGQGGGGQGGG GQQTVVVTQEPLSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGCPCPAPELLGGPS VFLFPPKPKDTLMISRTPEVTCVVVDVSHEEPE VKFNWYVDGVEVHNAKTKPCEEQYGSTYRCV SVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQGNVFSCSVMHEA LHNHYTQKSLSLSPGKGGGGQGGGGQGGGGQ GGGGQGGGGQGGGGQCPPCPAPELLGGPSVFL FPPKPKDTLMISRTPEVTCVVVDVSHEEPEVKF NWYVDGVEVHNAKTKPCEEQYGSTYRCVSVL TVLHQDWLNGKEYKCKVSNKALPAPIEKTISK AKGQPREPQVYTLPPSREEMTKNQVSLTCLVK GFYPSDIAVEWESNGQPENNYKTTTPVLDSG SFFLYSKLTVDKSRWQQGNVFSCSVMHEALH NHYTQKSLSLSPGKGGGGQVQLVQSGAEVVK PGASVKVCKASGYTFTSSWMNWVRQAPGQC LEWMGRIYTGGETKYSKGFQGRVITITRDTSA STAYMELSSLTSEDVAVYYCARQRDYGALYA MDYWGQGTLLTVSSGGGGQGGGGQGGGGQD IQLTQSPSFLSASVGDRVTITCRASDDIYSYLA WYQKPKGKAPKLLVYNAKTLAEGVPSRFSGSGS GTEFTLITSSLPEDFATYYCQNHDRTPFTFGC GTKVDIKSGGGGQEVQLVESGGGLVQPGGSLK LSAASGFTFNKYAINWVREAPGKGLEWVARI RSKYNNYATYYADAVKDRFTISRDDSKNTAYL QMNNLKTEDTAVYYCVRNANFGTSYISYFAY WGQGTLLTVSSGGGGQGGGGQGGGGQQTVV TQEPLSLTVSPGGTVTLTCGSSTGAVTSGNYPN WVQKKPGQAPRGLIGGTKFLAPGTPARFSGSL LGGKAALTLSGVQPEDEAEYYCVLWYSNRWV FGSGTKLTVL
285.	MS 15-B12 CC x I2M2 x G4 x scFc	artificial	aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS

	<p>x G4 x CH3 26-E5 CC x I2M2_GQ - Full Sequence</p>			<p>RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGTTLTVTVSSGGGGQGGGGQGGG GQQT VVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWWVQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGCPCPAPELLGGPS VFLFPPKPKDTLMISRTPEVTCVVVDVSHEEPE VKFNWYVDGVEVHNAKTKPCEEQYGSTYRCV SVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQGNVFCFSVMHEA LHNHYTQKSLSLSPGKGGGGQGGGGQGGGGQ GGGGQGGGGQGGGGQCPPCPAPELLGGPSVFL FPPKPKDTLMISRTPEVTCVVVDVSHEEPEVKF NWYVDGVEVHNAKTKPCEEQYGSTYRCVSVL TVLHQDWLNGKEYKCKVSNKALPAPIEKTISK AKGQPREPQVYTLPPSREEMTKNQVSLTCLVK GFYPSDIAVEWESNGQPENNYKTTTPVLDSDG SFFLYSKLTVDKSRWQQGNVFCFSVMHEALH NHYTQKSLSLSPGKGGGGQVQLVQSGAEVKK PGASVKVCKASGYTFTSYWMHWVRQAPGQC LEWMGVIRTSTSYTIYNQKFKGRVTMTRDTST STVYMESSLRSED TAVYYCARSGPGYFDVW GQGTMTVTVSSGGGGQGGGGQGGGGQDIQLTQ SPSFLSASVGDRVTITCRASENIYSYLAWYQQK PGKAPKLLIYNAKTLAEGVPSRFSGSGSGTEFT LTISSLPEDFATYYCQHNRYGTPYTFGCGTKLE IKSGGGGQEVQLVESGGGLVQPGGSLKLSCAA SGFTFNKYAINWVREAPGKGLEWVARIRSKYN NYATYYADAVKDRFTISRDDSKNTAYLQMN LKTEDTAVYYCVRNANFGTSYISYFAYWGQ TLTVTVSSGGGGQGGGGQGGGGQQT VVTQEPS LTVSPGGTVTLTCGSSTGAVTSGNYPNWWVQK PGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAA LTLSGVQPEDEAEYYCVLWYSNRWVFGSGTK LTVL</p>
<p>286.</p>	<p>MS 15-B12 CC x I2M2 x G4S3 x scFc x G4S3 x CH3 15-E11 CC x I2M2 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS</p>

				<p>YISYFAYWGQGLVTVSSGGGGSGGGGSGGG GSQTVVTQEPLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYYWMMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFGQGRVTMTVDTSS STAYMELSRLRSDDTAVYYCATRYFYVMDYW GQGLVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAINWVREAPGKGLEWVARIRSKYN NYATYYADAVKDRFTISRDDSKNTAYLQMNN LKTEDTAVYYCVRNANFGTYSYISYFAYWGQ TLVTVSSGGGGSGGGGSGGGGSGQTVVTQEPL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQKKP GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGSGTKL TVL</p>
287.	MS 15-B12 CC x I2Ccc(44/100)x (G4S)3x scFex (G4S)3x CH3 14- D1 CCx I2Ccc(44/100) - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSSGSSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK CLEWVARIRSKYNNYATYYADSVKDRFTISR DSKNTAYLQMNNLKTEDTAVYYCVRHGNFGN SYISYWAYWGQGLVTVSSGGGGSGGGGSGG GGQTVVTQEPLTVSPGGTVTLTCGSSTGAVT SGNYPNWVQKPGQAPRGLIGGTKFLAPGTPA RFGSLLGGKAALTLSGVQPEDEAEYYCVLWY SNRWVFGCGTKLTVLGGGGSGGGGSGGGGSD KHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK</p>

				<p>EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYFTFTSYWMHWVRQAPGQCLE WMGVITYTSGSYTIYNQKFQGRVTMTRDTSTST AYMELSSLRSEDNAVYYCARSGPGYFDVWGQ GTMVTVSSGGGGSGGGGSGGGGSDIQLTQSPS FLSASVGDRTITCRASGNIHNYLAWYQQKPG KAPKLLIYNAKTLAEGVPSRFSGSGSGTEFTLKI SSLQPEDFATYYCQHFAWTPYTFGCGTKLEIKS GGGGSEVQLVESGGGLVQPGGSLKLSAASGF TFNKYAMNWVRQAPGKCLEWVARIRSKYNN YATYYADSVKDRFTISRDDSKNTAYLQMNNL KTEDNAVYYCVRHGNGNSYISYWAYWGQGT LVTVSSGGGGSGGGGSGGGGSGTQVVTQEPSLT VSPGGTVTLTCSSTGAVTSGNYPNWVQKPG QAPRGLIGGTKFLAPGTPARFSGSLLGGKAALT LSGVQPEDEAEYYCVLWYSNRWVFGCGTKLT VL</p>
288.	<p>MS 15-B12 CC x I2L x (G4Q)3 x scfc x (G4Q)3 x CH3 005-D5 CC x I2L - Full Sequence</p>	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADNAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFLTITSSSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDNAVYYCVRAGNFG SSYISYFAYWGQGTLVTVSSGGGGQGGGGQ GGGQQTQVVTQEPSLTVSPGGTVTITCSSTGAV TSGNYPNWIQKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAALTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGQGGGGQGGGG QCPCPAPELLGGPSVFLFPPKPKDTLMISRTP EVTCTVVDVSHHEEPEVKFNWYVDGVEVHNAKT KPCEEQYGSTYRCVSVLTVLHQDWLNGKEYK CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTTPPVLDSDGSFFLYSKLTVDKSRW QQGNVFSCSVMHEALHNHYTQKSLSLSPGK GGGQGGGGQGGGGQGGGGQGGGGQGGGGQ CPCPAPELLGGPSVFLFPPKPKDTLMISRTP EVTCTVVDVSHHEEPEVKFNWYVDGVEVHNAKT PCEEQYGSTYRCVSVLTVLHQDWLNGKEYK</p>

				<p>KVSNKALPAPIEKTISKAKGQPREPQVYTLPPS REEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ PENNYKTPPVLDSDGSFFLYSKLTVDKSRWQ QGNVFSCSVMHEALHNHYTQKSLSLSPGKGG GGQGGGGQGGGGQEVQLLESQGGGLVQPGGSL RLSCAASGFSFSSYPINWVRQAPGKCLEWVGI WTGGGTNYASSVKGRFTISRDNKNTVYLQM NSLRAEDTAVYYCAKSRGVYDFKGRGAMDY WGQGTLLTVSSGGGGQGGGGQGGGGQDIVM TQSPDSLAVSLGERATINCKSSQSLLYSSNQKN YFAWYQQKPGQPPKLLIYWASTRESGVPDRFS GSGSGTDFLTITSSLQAEDVAVYYCQQYYSYP YTFGCGTKLEIKSGGGGQEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWVRQAPGK MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNLLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGTLLTVSSGGGGQGGGGQGG GGQQTIVTQEPSLTVSPGGTITITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLGSGVQPEDEAEYYCVLY SNRWVFGSGTKLTVL</p>
<p>289.</p>	<p>MS 15-B12 CC x I2L x (G4Q)3 x scfc x (G4Q)3 x CH3 26-E5 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGTDFLTITSSLPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNLLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTLLTVSSGGGGQGGGGQGG GGQQTIVTQEPSLTVSPGGTITITCGSSTGAV TSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAALTLGSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGQGGGGQGGGG QCPCPAPELLGGPSVFLFPPKPKDTLMISRTPE VTCVVVDVSHEEPEVKFNWYVDGVEVHNAKT KPCCEEQYGSTYRCVSVLTVLHQDWLNGKEYK CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTPPVLDSDGSFFLYSKLTVDKSRW QQGNVFSCSVMHEALHNHYTQKSLSLSPGKGG GGQGGGGQGGGGQGVQLVQSGAEVKKPGAS VKVSCKASGYTFSTSYWMHWVRQAPGQCLEW MGVIRTSTSYTIYNQKFKGRVTMTRDSTSTV YMELSSLRSEDTAVYYCARSGPGYFDVWGQG</p>

				<p>TMVTVSSGGGGQGGGGQGGGGQDDIQLTQSPSF LSASVGDRVTITCRASENIYSYLAWYQQKPGK APKLLIYNAKTLAEGVPSRFSGSGSGTEFTLTIS SLQPEDFATYYCQHNYGTPYTFGCGTKLEIKSG GGGQEVQLVESGGGLVQPGGSLKLSCAASGFT FNKYAMNWVRQAPGKGMEWVARIRSKYNNY ATYYADAVKDRFTISRDDSKNTLYLQMNNLK TEDTAVYYCVRAGNFGSSYISYFAYWGQGLV TVSSGGGGQGGGGQGGGGQQT VVTQEPSLTV SPGGT VTITCGSSTGAVTSGNYPNWIQKPGQ APRGLIGGTKFLAPGTPARFSGSLEGGKAALTL SGVQPEDEAEYYCVLYYSNRWVFGSGTKLTV L</p>
<p>290.</p>	<p>MS 15-B12 CC x I2L x (G4Q)3x scFcmo d x (G4Q3) x CH3 24- D7 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLQESGPGLVKPSETLSLTCTVSSGSSSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFLTITSSSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLVTVSSGGGGQGGGGQGG GGGQQT VVTQEPSLTVSPGGT VTITCGSSTGAV TSGNYPNWIQKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAALTL SGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGQGGGGQGGGG QCPCPAPELLGGPSVFLFPPKPKDTLMISRTPE VTCVVVDVSHEEPEVKFNWYVDGVEVHNAKT KPCEEQYGSTYRCVSVLTVLHQDWLNGKEYK CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTTTPVLDS DGSFFLYSKLTVDKSRW QQGNVFSCSVMHEALHNHYTQKSLSLSPGKG GGGQGGGGQGGGGQGGGGQGGGGQGGGGQ CPCPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEEPEVKFNWYVDGVEVHNAKT PCEEQYGSTYRCVSVLTVLHQDWLNGKEYK KVS NKALPAPIEKTISKAKGQPREPQVYTLPPS REEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ PENNYKTTTPVLDS DGSFFLYSKLTVDKSRWQ QQGNVFSCSVMHEALHNHYTQKSLSLSPGKG GGQGGGGQGGGGQVQLVQSGAEVKKPGAS VKVSCKASGYTFTNYWMNWVRQAPGQCLEW MGNHNSKAHGTNYNQKFQGRVTMTVDTSSST AYMELSR LRSDDTAVYYCATRYFYVMDYWG QGT LTVTVSSGGGGQGGGGQGGGGQDDIQLTQSP PSSLSASVGDRVTITCRASQDISNYLNWYQQK GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGQEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKGMEWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ</p>

				GTLVTVSSGGGGQGGGGQGGGGQQTVVVTQEP SLTVSPGGTVTITCGSSTGAVTSGNYPNWIQKK PGQAPRGLIGGTKFLAPGTPARFSGSLEGGKAA LTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKL TVL
291.	MS 15-B12 CC x I2L x G4 x scFc x G4 x CH3 15-E11 CCx I2L - Full Sequence	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYPNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGGQTVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYG STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYKT TPPVLDSGDSFFLYSKLTVDKSRWQQGNVFC SVMHEALHNHYTQKLSLSPGKGGGGSGGGG SGGGGSGGGGSGGGGSGGGGSDKTHTCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTTTPVLDSDGSFFLYSKLTVDKSRWQQGNV SCSVMHEALHNHYTQKLSLSPGKGGGGQVQ LVQSGAEVKKPGASVKVCKASGYTFTNYWM NWVRQAPGQCLEWMGNIAYGKGTNYNQKF QGRVTMTVDTSSSTAYMELSRRLSDDTAVYY CATRYFYVMDYWGQGLTVTVSSGGGGSGGG GSGGGGSDIQMTQSPSSLSASVGDRVTITCRAS QDISNYLNWYQQKPGKVPKLLIYYSRLHSGV PSRFSGSGSGTDFLTITSSLPEDVATYYCVQY AQFPLTFGCGTKVEIKSGGGGSEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKGMEWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTLYLQMNNLKTEDTAVYYCVRAGN FGSSYISYFAYWGQGLTVTVSSGGGGSGGGG GGGGQTVVTQEPSLTVSPGGTVTITCGSSTGA VTSNYPNWIQKKPGQAPRGLIGGTKFLAPGT PARFSGSLEGGKAAALTLSGVQPEDEAEYYCVL YYSNRWVFGSGTKLTVL
292.	MS 15-B12 CC x I2L x G4 x scFc x G4 x CH3 22-A12 CC x I2L_GQ -	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYPNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG

	Full Sequence			<p>QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGQGGGGQ GGGQQT VVTQEPSLTVSPGGTVTITCGSSTGAV TSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAALTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGCPCPAPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSH EEPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCSSVMHE ALHNHYTQKSLSLSPGKGGGGQGGGGQGGGG QGGGGQGGGGQGGGGQCPPCPAPELLGGPSV FLFPPKPKDTLMISRTPEVTCVVVDVSH EEPVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCSSVMHE ALHNHYTQKSLSLSPGKGGGGQVQLVQSGAEV VKPGASVKVCKASGYTFTSSWMNWVRQAPG QCLEWMGRIYTGGETKYSKGFQGRVTITRDT SASTAYMELSSLTSEDTAVYYCARQRDYGALY AMDYWGQGLTVTVSSGGGGQGGGGQGGGGQ DIQLTQSPSFLSASVGDRVTITCRASDDIYSYLA WYQQKPGKAPKLLVYNAKTLAEGVPSRFSGS GSGTEFTLTISSLQPEDFATYYCQNHDRTPFTFG CGTKVDIKSGGGGQEVQLVESGGGLVQP GGSLLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGQGGGGQ GGGQQT VVTQEPSLTVSPGGTVTITCGSSTGAV TSGNYPNWIQKKPGQAPRGLIGGTKFLAPGTP ARFSGSLEGGKAALTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGCPCPAPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSH EEPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCSSVMHE ALHNHYTQKSLSLSPGKGGGGQVQLVQSGAEV VKPGASVKVCKASGYTFTSSWMNWVRQAPG QCLEWMGRIYTGGETKYSKGFQGRVTITRDT SASTAYMELSSLTSEDTAVYYCARQRDYGALY AMDYWGQGLTVTVSSGGGGQGGGGQGGGGQ DIQLTQSPSFLSASVGDRVTITCRASDDIYSYLA WYQQKPGKAPKLLVYNAKTLAEGVPSRFSGS GSGTEFTLTISSLQPEDFATYYCQNHDRTPFTFG CGTKVDIKSGGGGQEVQLVESGGGLVQP GGSLLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGQGGGGQ GGGQQT VVTQEPSLTVSPGGTVTITCGSSTGAV</p>
293.	MS 15-B12 CC x I2L x G4 x scFc x G4 x CH3 24-D7 CC x I2L_GQ - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGQGGGGQ GGGQQT VVTQEPSLTVSPGGTVTITCGSSTGAV</p>

				<p>TSGNYPNWIQKKPGQAPRGLIGGKFLAPGTP ARFSGSLEGGKAALTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGCPCPAPELLGG PSVFLFPPKPKDTLMISRTP EVT CVVVDVSHEE PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCSCVMHE ALHNHYTQKSLSLSPGKGGGGQGGGGQGGGG QGGGGQGGGGQGGGGQCPPCPAPELLGGPSV FLFPPKPKDTLMISRTP EVT CVVVDVSHEEPEV KFNWYVDGVEVHNAKTKPCEEQYGSTYRCVS VLTVLHQDWLNGKEYKCKVSNKALPAPIEKTI SKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTPPVLDS DGSFFLYSKLTVDKSRWQQGNVFCSCVMHEA LHNHYTQKSLSLSPGKGGGGQVQLVQSGAEV KKPGASVKVCKASGYFTFTNYWMNWVRQAP GQCLEWMGNIHKAHGTNYNQKFQGRVTMT VDTSSSTAYMELSRLRSDDTAVYYCATRYFYV MDYWGGGTLTVVSSGGGGQGGGGQGGGGQD IQMTQSPSSLSASVGDRVTITCRASQDISNYLN WYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGS GTDFTLTISSLQPEDVATYYCVQYAQFPLTFGC GTKVEIKSGGGGQEVQLVESGGGLVQPGGSLK LSCAASGFTFNKYAMNWVRQAPGKGMWVA RIRSKYNNYATYYADAVKDRFTISRDDSKNTL YLQMNNLKTEDTAVYYCVRAGNFGSSYISYFA YWGQGTLLTVVSSGGGGQGGGGQGGGGQQT VTQEPSTVSPGGTVTITCGSSTGAVTSGNYPN WIQKKPGQAPRGLIGGKFLAPGTPARFSGSLE GGKAALTLSGVQPEDEAEYYCVLYYSNRWVF GSGTKLTVL</p>
294.	MS 15-B12 CC x I2L x G4S3 x scFc x G4S3 x CH3 005-D5 CC x I2L - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQQSY TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTLLTVVSSGGGGSGGGGSGG GGSQTVVTQEPSTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLY YSNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS</p>

				<p>NHYTQKSLSLSPGKGGGGQVQLVQSGAEVKK PGASVKVSCKASGYTFTNYWMNWVRQAPGQ CLEWMGNIHSKAHGTNYNQKFQGRVTMTVDT SSSTAYMELSRLRSDDTAVYYCATRYFYVMD YWGQGTLLTVVSSGGGGQGGGGQGGGGQDIQ MTQSPSSLSASVGDRVTITCRASQDISNYLNWY QQKPGKVPKLLIYYTSRLHSGVPSRFSGSGSGT DFTLTISLQPEDVATYYCVQYAFPLTFGCGT KVEIKSGGGGQEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAINWVREAPGKGLEWVARIRS KYNNYATYYADAVKDRFTISRDDSKNTAYLQ MNLKTEDTAVYYCVRNANFGTSYISYFAYW GQGTLTVVSSGGGGQGGGGQGGGGQQT VVT QEPSLTVSPGGTVTLTCGSSTGAVTSGNYPNW VQKPGQAPRGLIGGTKFLAPGTPARFSGSLLG GKAALTLGSGVQPEDEAEYYCVLWYSNRWVFG SGTKLTVL</p>
<p>296.</p>	<p>MS 15-B12 CC x I2M2 x (G4Q)3x scFcmo d x (G4Q)3 x CH3 005-D5 CC x I2M2_GQ - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESG PGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAAS TLQSGVP SRFSGSGSGTDFTLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGK LEWVARIRSKYN NYATYYADAVKDRFTISRDD SKNTAYLQMNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGTLTVVSSGGGGQGGGGQGGG GQQT VVTQEPSLTVSPGGTVTLTCGSSTGAVT GNYPNWVQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLGSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGQGGGGQGGGGQC PPCAPPELLGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHEEPEVKFNWYVDGVEVHNAKTKP CEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQG NVFSCSVMEALHNHYTQKSLSLSPGKGGGG QGGGGQGGGGQGGGGQGGGGQGGGGQCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVV VDVSHEEPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDS DGSFFLYSKLTVDKSRWQQGNV FSCSVMEALHNHYTQKSLSLSPGKGGGGQ GGGGQGGGGQEVQLLES GGGLVQPGGSLRLSC AASGFSFSSYPINWVRQAPGKCLEWVGVITG GGTNYASSVKGRFTISRDN SKNTVYLQMNSLR AEDTAVYYCAKSRGVYDFKGRGAMDYWGQG TLTVVSSGGGGQGGGGQGGGGQDIVMTQSPD SLAVSLGERATINCKSSQSLLYSSNQKNYFAW YQQKPGQPPKLLIYWASTRESGVPDRFSGSGG TDFTLTISLQAEDVAVYYCQYYSYPTFGC</p>

			<p>GTKLEIKSGGGGQEVQLVESGGGLVQPGGSLK LSCAASGFTFNKYAINWVREAPGKGLEWVARI RSKYNNYATYYADAVKDRFTISRDDSKNTAYL QMNNLKTEDTAVYYCVRNANFGTSYISYFAY WGQGTLLTVSSGGGGQGGGGQGGGGQQTVV TQEPSLTVSPGGTVTLTCGSSTGAVTSGNYPN WVQKKPGQAPRGLIGGTKFLAPGTPARFSGSL LGGKAALTLSGVQPEDEAEYYCVLWYSNRWW FSGGTKLTVL</p>
<p>297.</p>	<p>MS 15-B12 CC x I2M2 x (G4Q)3x scFcmo d x (G4Q)3 x CH3 26- E5 CCx I2M2_GQ - Full Sequence</p>	<p>artificial</p>	<p>aa</p> <p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKPKLLIYAASSTLQSGVP SRFSGSGGTDFLTITSSLPEDFATYYCQOSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGT SYISYFAYWGQGTLLTVSSGGGGQGGGGQGGG GQQTVVTVTQEPSLTVSPGGTVTLTCGSSTGAVT SGNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGQGGGGQGGGGQC PPCAPELLGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHEEPEVKFNWYVDGVEVHNAKTKP CEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDSGDSFFLYSKLTVDKSRWQQG NVFSCVMHEALHNHYTQKSLSLSPGKGGGGG QGGGGQGGGGQGGGGQGGGGQGGGGQCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVV VDVSHHEEPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSGDSFFLYSKLTVDKSRWQQGNV FSCVMHEALHNHYTQKSLSLSPGKGGGGGQ GGGQGGGGQGVQLVQSGAEVKKPGASVKVSC KASGYTFTSYWMHWVRQAPGQCLEWMGVIR TSTSYTIYNQKFKGRVTMTRDTSTSTVYME LSSLRSEDTAVYYCARSFGPYFDVWGQGTMTV TVSSGGGGQGGGGQGGGGQDIQLTQSPSFLS ASVGDRTITCRASENIYSYLAWYQQKPGKAP KLLIYNAKTLAEGVPSRFSGSGSTEFTLTISS LPEDFATYYCQHNRYGTPYTFGCGTKLEIKS GGGGQEVQLVESGGGLVQPGGSLKLSCAASG FTFNKYAINWVREAPGKGLEWVARIRSKYNNY ATYYADAVKDRFTISRDDSKNTAYLQMNNL KTEDTAVYYCVRNANFGTSYISYFAYWGQGT LLTVSSGGGGQGGGGQGGGGQQTVVTVTQEP SLTVSPGGTVTLTCGSSTGAVTSGNYPNWVQ KPGQAPRGLIGGTKFLAPGTPARFSGSLLGG KAALTLSGVQPEDEAEYYCVLWYSNRWVFG SGTCLTVL</p>

<p>298.</p>	<p>MS 15-B12 CCx 6H10.09x (G4)x scFcx (G4)x CH3 14-D1 CCx 6H10.09 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFTLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTMTVTVSSGGGGSGGGGSGG GGSTVVTQEPSTVSPGGTITITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGKFLAPGTPA RFSGSLEGGKAALTLGSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYG STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYKT TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFC SVMHEALHNHYTQKSLSLSPGKGGGGSGGGG SGGGGSGGGGSGGGGSGGGGSDKTHTCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTPPVLDSDGSFFLYSKLTVDKSRWQQGNV SCSVMHEALHNHYTQKSLSLSPGKGGGGQVQ LVQSGAEVKKPGASVKVSCKASGYTFTSYWM HWVRQAPGQCLEWMGVIYTSGSYTIYNQKFQ GRVTMTRDTSTSTAYMELSSLRSEDVAVYYCA RSGPGYFDVWGQGTMTVTVSSGGGGSGGGGSG GGSDIQLTQSPSFLSASVGDRVTITCRASGNIH NYLAWYQQKPGKAPKLLIYNAKTLAEGVPSRF SGSGSGTEFTLKISLQPEDFATYYCQHFATP YTFGCGTKLEIKSGGGGSEVQLVESGGGLVQP GGSLKLSAASGFTFNKYAMNWVRQAPGK MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGTMTVTVSSGGGGSGGGGSGG GSQTVVTQEPSTVSPGGTITITCGSSTGAVTS GNYPNWIQKKPGQAPRGLIGGKFLAPGTPAR FSGSLEGGKAALTLGSGVQPEDEAEYYCVLYS NRWVFGSGTKLTVL</p>
<p>299.</p>	<p>MS 15-B12 CCx 6H10.09x (G4S)3x scFcx (G4S)3x CH3 14- D1 CCx 6H10.09 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFTLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK</p>

				<p>GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVVSSGGGGSGGGGSGG GGSQTVVTQEPLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKHTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS ASVKVSCKASGYTFTSYWMHWVRQAPGQCLE WMGVIYTSGSYTIYNQKFQGRVTMTRDTSTST AYMELSSLRSEDVAVYYCARSGPGYFDVWGQ GTMVTVSSGGGGSGGGGSGGGGSDIQLTQSPS FLSASVGDRVTITCRASGNIHNYLAWYQQKPG KAPKLLIYNAKTLAEGVPSRFSGSGSGTEFTLKI SSLQPEDFATYYCQHFAWTPYTFGCGTKLEIKS GGGGSEVQLVESGGGLVQPGGSLKLSAASGF TFNKYAMNWVRQAPGKGMWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTLYLQMNNL KTEDTAVYYCVRAGNFGSSYISYFAYWGQGL TVVSSGGGGSGGGGSGGGGSGGGGSGGGGSG SPGGTVTITCGSSTGAVTSGNYPNWIQKKPGQ APRGLIGGTKFLAPGTPARFSGSLEGGKAALTL SGVQPEDEAEYYCVLYYSNRWVFGSGTKLTV L</p>
300.	MS 15-B12 CCx I2M2 x G4 x scFc x G4 x CH3 15- E11 CC x I2M2 _ GQ - Full Sequence	artificial	aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYPNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFLTITSSLQPEDFATYYCQSSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGK LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVVSSGGGGQGGGGQGGG GQQT VVTQEPLTVSPGGT VTLTCGSSTGAVTS GNYPNWWVQKKPGQAPRGLIGGTKFLAPGTPAR FGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGCPCPPELLGGPS VFLFPPKPKDTLMISRTPVTCVVVDVSHPEE</p>

				<p>VKFNWYVDGVEVHNAKTKPCEEQYGSTYRCV SVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTPVLDS DGSFFLYSKLTVDKSRWQQGNVFSCSVMHEA LHNHYTQKSLSLSPGKGGGGQGGGGQGGGGQ GGGGQGGGGQGGGGQCPCPAPELLGGPSVFL FPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKF NWXVVDGVEVHNAKTKPCEEQYGSTYRCVSVL TVLHQDWLNGKEYKCKVSNKALPAPIEKTISK AKGQPREPQVYTLPPSREEMTKNQVSLTCLVK GFYPSDIAVEWESNGQPENNYKTTPVLDSDG SFFLYSKLTVDKSRWQQGNVFSCSVMHEALH NHYTQKSLSLSPGKGGGGQVQLVQSGAEVKK PGASVKVSKASGYFTFTNYWMNWVRQAPGQ CLEWMGNIAYGVKGTNYNQKFGQGRVTMTVD TSSSTAYMELSLRSDDTAVYYCATRYFYVMD YWGQGLTVTVSSGGGGQGGGGQGGGGQDIQ MTQSPSSLSASVGDRVTITCRASQDISNYLNWY QQKPGKVPKLLIYYTSRLHSGVPSRFSGSGSGT DFTLTISSLQPEDVATYYCVQYAQFPLTFGCGT KVEIKSGGGGQEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAINWVREAPGKGLEWVARIRS KYNRYATYYADAVKDRFTISRDDSKNTAYLQ MNNLKTEDTAVYYCVRNANFGTSYISYFAYW GQGTLTVTVSSGGGGQGGGGQGGGGQQT VVT QEPSLTVSPGGTVTLTCSSTGAVTSGNYPNW VQKPKGQAPRGLIGGKFLAPGTPARFSGSLLG GKAALTLGSGVQPEDEAEYYCVLWYSNRWVFG SGTKLTVL</p>
301.	MS 15-B12 CC x I2L x G4S3 x scFc x G4S3 x CH3 15-E11 CC x I2 - Full Sequence	artificial	aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNRYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTLTVTVSSGGGGSGGGGSGG GGSTVVTQEPSLTVSPGGTVTITCSSTGAVT SGNYPNWIQKPKGQAPRGLIGGKFLAPGTPA RFSGSLEGGKAALTLGSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVH AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPVLDSDG SFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH</p>

				<p>NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCSCVMHEALHNHYTQKSLSLSPG KGGGGSGGGSGGGGSQVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFQGRVTMTVDTSS STAYMELSRLSDDTAVYYCATRYFYVMDYW GQGTLVTVSSGGGGSGGGSGGGGSDIQTQS PSSLSASVGDRVITICRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKGMWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ GTLTVTVSSGGGGSGGGSGGGGSQT VVTQEPS LTVSPGGT VITICGSSTGAVTSGNYPNWIQKKP GQAPRGLIGGTKFLAPGTPARFSGSLEGGKAAL TLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLT VL</p>
<p>302.</p>	<p>MS 15-B12 CC x I2L x G4 x scFc x G4 x CH3 005-D5 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVITICRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMN NLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTLVTVSSGGGGSGGGGSGG GGSTVVTQEPSLTVSPGGT VITICGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQY STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYKT TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFC SCVMHEALHNHYTQKSLSLSPGKGGGGSGGGG SGGGGSGGGGSGGGGSGGGGSDKTHTCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTTPPVLDSDGSFFLYSKLTVDKSRWQQGNV FCVMHEALHNHYTQKSLSLSPGKGGGGGEVQL LES GGGLVQPGGSLRLS CAASGFSFSSYPINWV RQAPGKCLEWVGVIWTGGGTNYASSVKGRFTI SRDNSKNTVYLQMN SLRAEDTAVYYCAKSRG</p>

				<p>VYDFKGRGAMDYWGQGTTLVTVSSGGGGSGG GGSGGGSDIVMTQSPDSLAVSLGERATINCKS SQSLLYSSNQKNYFAWYQQKPGQPPKLLIYWA STRESGVPDRFSGSGGTDFTLTISSLQAEDVA VYYCQQYYSYPYTFGCGTKLEIKSGGGGSEVQ LVESGGGLVQPGGSLKLSAASGFTFNKYAMN WVRQAPGKGMWVVARIRSKYNNYATYYADA VKDRFTISRDDSKNTLYLQMNNLKTEDTAVYY CVRAGNFGSSYISYFAYWGQGTTLVTVSSGGGG SGGGSGGGGSQTVVVTQEPSLTVSPGGTVTITC GSSTGAVTSGNYPNWIQKPGQAPRGLIGGTK FLAPGTPARFSGSLEGGKAALTLGVQPEDEAE YYCVLYYSNRWVFGSGTKLTVL</p>
<p>303.</p>	<p>MS 15-B12 CC x I2L x G4 x scFc x G4 x CH3 22-A12 CC x I2L - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGTTLVTVSSGGGGSGGGGSGG GGGQTVVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYG STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYKT TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFC SVMHEALHNHYTQKSLSLSPGKGGGGSGGGG SGGGGSGGGGSGGGGSGGGGSDKTHTCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTPPVLDSDGSFFLYSKLTVDKSRWQQGNV SCSVMHEALHNHYTQKSLSLSPGKGGGGQVQ LVQSGAEVVKPGASVKVSKASGYTFTSSWM NWVRQAPGQCLEWMGRIYTGGETKYSKGFQ GRVTITRDTSASTAYMELSSLTSEDYAVYYCAR QRDYGALYAMDYWGQGTTLVTVSSGGGGSGG GGSGGGGSDIQLTQSPSFLSASVGDRVTITCRA SDDIYSYLAWYQQKPGKAPKLLVYNAKTLAE GVPSRFSGSGGTDFTLTISSLQPEDFATYYCQN HDRTPFTFGCGTKVDIKSGGGGSEVQLVESGG GLVQPGGSLKLSAASGFTFNKYAMNWVRQA PGKGMWVVARIRSKYNNYATYYADAVKDRFT ISRDDSKNTLYLQMNNLKTEDTAVYYCVRAG NFGSSYISYFAYWGQGTTLVTVSSGGGGSGGGG</p>

				SGGGGSQTVVTQEPSLTVSPGGTVTITCGSSTG AVTSGNYPNWIQKKPGQAPRGLIGGTKFLAPG TPARFSGSLEGGKAALTLSGVQPEDEAEYYCV LYYSNRWVFGSGTKLTVL
304.	MS 15-B12 CC x I2L x G4 x scFc x G4 x CH3 24-D7 CC x I2L - Full Sequence	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYPNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSQTVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYG STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYKT TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFC SVMHEALHNHYTQKSLSLSPGKGGGGSGGGG SGGGGSGGGGSGGGGSGGGGSDKTHTCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTTTPVLDSDGSFFLYSKLTVDKSRWQQGNV SCSVMHEALHNHYTQKSLSLSPGKGGGGQVQ LVQSGAEVKKPGASVKVSCKASGYFTFTNYWM NWVRQAPGQCLEWMGNIHKAHGTNYNQKF QGRVTMTVDTSSSTAYMELSRRLSDDTAVYY CATRYFYVMDYWGQGLTVTVSSGGGGSGGGG GSGGGGSDIQMTQSPSSLSASVGDRVTITCRAS QDISNYLNWYQQKPGKVPKLLIYYSRLHSGV PSRFSGSGSGTDFLTITSSLPEDVATYYCVQY AQFPLTFGCGTKVEIKSGGGGSEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKGMWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTLYLQMNNLKTEDTAVYYCVRAGN FGSSYISYFAYWGQGLTVTVSSGGGGSGGGG GGGGSQTVVTQEPSLTVSPGGTVTITCGSSTGA VTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGT PARFSGSLEGGKAALTLSGVQPEDEAEYYCVL YYSNRWVFGSGTKLTVL
305.	MS 15-B12 CC x I2L x G4S3 x scFc x G4S3 x CH3 22- A12 CC x I2L - Full Sequence	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYPNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ

				<p>GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSQT VVTQEPLTVSPGGT VTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVVKPG ASVKVSCKASGYTFTSSWMNWVRQAPGQCLE WMGRIYTGTGETKYSKGFQGRVTITRDTAST AYMELSSLTSED TAVYYCARQRDYGALYAMD YWGQGLTVTVSSGGGGSGGGGSGGGGSDIQL TQSPSFLSASVGDRVTITCRASDDIYSYLAWYQ QKPGKAPKLLVYNAKTLAEGVPSRFSGSGSGT EFTLTISSLQPEDFATYYCQNHDRTPFTFGCGT KVDIKSGGGGSEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAMNWVRQAPGKMEWVARI RSKYNNYATYYADAVKDRFTISRDDSKNTLYL QMNNLKTEDTAVYYCVRAGNFGSSYISYFAY WGQGLTVTVSSGGGGSGGGGSGGGGSGT VVT QEPLTVSPGGT VTITCGSSTGAVTSGNYPNWI QKKPGQAPRGLIGGTKFLAPGTPARFSGSLEGG KAALTLSGVQPEDEAEYYCVLYYSNRWVFGS GTKLTVL</p>
306.	MS 15-B12 CC x I2Lx G4 x scFc x G4 x CH3 26-E5 CC x I2L - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSQT VVTQEPLTVSPGGT VTITCGSSTGAVT</p>

			<p>SGNYPNWIQKKPGQAPRGLIGGTFKFLAPGTPA RFGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYG STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYKT TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFC SVMHEALHNHYTQKSLSLSPGKGGGGSGGGG SGGGSGGGGGSGGGGGSGGGGSDKTHTCPPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEDEPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTTTPPVLDSDGSFFLYSKLTVDKSRWQQGNV SCSVMHEALHNHYTQKSLSLSPGKGGGGQVQ LVQSGAEVKKPGASVKVCKASGYTFTSYWM HWVRQAPGQCLEWMGVIRTSTSYTIYNQKFK GRVTMTRDTSTSTVYMESSLRSEDVAVYYCA RSGPGYFDVWGQGTMTVTVSSGGGGSGGGGGG GGGSDIQLTQSPSFLSASVGDRVTITCRASENIY SYLAWYQQKPGKAPKLLIYNAKTLAEGVPSRF SGSGSGTEFTLTISLQPEDFATYYCQHNYGTP YTFGCGTKLEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGLTVTVSSGGGGSGGGGGSGGG GSQTVVTQEPSLTVSPGGTVTITCGSSTGAVTS GNYPNWIQKKPGQAPRGLIGGTFKFLAPGTPAR FSGSLEGGKAALTLSGVQPEDEAEYYCVLYYS NRWVFGSGTKLTVL</p>
307.	MS 15-B12 CC x I2M2 x G4 x scfc x G4 x CH3 22- A12 CC x I2M2 - Full Sequence	artificial	<p>Aa QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGGTDFTLTISLQPEDFATYYCQSSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGGSGGG GSQTVVTQEPSLTVSPGGTVLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTFKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGDKTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFCSS</p>

				<p>VMHEALHNHYTQKSLSLSPGKGGGGSGGGGS GGGGSGGGSGGGSGGGGSDKTHTCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL VQSGAEVVKPGASVKVSCKASGYTFTSSWMN WVRQAPGQCLEWMGRIYTGGETKYSKGFQ RVTITRDTSASTAYMELSSLTSEDVAVYYCARQ RDYGALYAMDYWGQGLTVTVSSGGGGSGGG GSGGGGSDIQLTQSPSFLSASVGDRVTITCRAS DDIYSYLAWYQQKPKGAPKLLVYNAKTLAEG VPSRFSGSGSGTEFTLTISLQPEDFATYYCQNH DRTPFTFGCGTKVDIKSGGGGSEVQLVESGGG LVQPGGSLKLSKAASGFTFNKYAINWVREAPG KGLEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRNANFG TSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVT SGNYPNWVQKPKGQAPRGLIGGTKFLAPGTPA RFSGLLGGKAALTLGSGVQPEDEAEYYCVLWY SNRWVFGSGTKLTVL</p>
308.	MS 15-B12 CC x I2M2 x G4 x scfc x G4 x CH3 24-D7 CC x I2M2 - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPKGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFLTISLQPEDFATYYCQSSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSKAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPKGQAPRGLIGGTKFLAPGTPAR FSGLLGGKAALTLGSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGDKTHTCPAPAP LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGSGGGGS GGGGSGGGSGGGSGGGGSDKTHTCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL</p>

				<p>VQSGAEVKKPGASVKVSCASGYTFTNYWMN WVRQAPGQCLEWMGNIHKAHGTNYNQKFQ GRVTMTVDTSSTAYMELSRRLSDDTAVYYC ATRYFYVMDYWGQGLVTVSSGGGGSGGGGS GGGSDIQMTQSPSSLSASVGDRVITICRASQD ISNYLNWYQQKPGKVPKLLIYYTSRLHSGVPSR FSGSGSGTDFLTISLQPEDVATYYCVQYAQF PLTFGCGTKVEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAINWVREAPGKGL EWVARIRSKYNNYATYYADAVKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRNANFGTSY ISYFAYWGQGLVTVSSGGGGSGGGSGGGGS QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWWVQKPGQAPRGLIGGTFKFLAPGTPARFS GSLGKAALTLGSGVQPEDEAEYYCVLWYSN RWVFGSGTKLTVL</p>
<p>309.</p>	<p>MS 15-B12 CC x I2M2 x G4 x scfc x G4 x CH3 005- D5 CC x I2M2 - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGGSGGGG SGGGSDIVMTQSPSSLSASVGDRVITICRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGTDFLTISLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLVTVSSGGGGSGGGSGGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWWVQKPGQAPRGLIGGTFKFLAPGTPAR FSGSLGKAALTLGSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGDKTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVDVVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGSGGGGS GGGGSGGGSGGGSGGGGSDKHTHTCPPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGGEVQL LESGGGLVQPGGSLRLS AASGFSFSSYPINWV RQAPGKCLEWGVIVTGGGTNYASSVKGRFTI SRDNSKNTVYLQMNSLRAEDTAVYYCAKSRG VYDFKGRGAMDYWGQGLVTVSSGGGGSGG GGSGGGSDIVMTQSPDSLAVSLGERATINCKS SQSLLYSSNQKNYFAWYQQKPGQPPKLLIYWA STRESGVPDRFSGSGSGTDFLTISLQAEDVA VYYCQQYYSYPYTFGCGTKLEIKSGGGGSEVQ LVESGGGLVQPGGSLKLSAASGFTFNKYAIN</p>

				<p>WVREAPGKGLEWVARIRSKYNNYATYYADA VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRNANFGTSYISYFAYWGQGLTVTVSSGGG GSGGGGSGGGGSQTVVTQEPLTVSPGGTVTL TCGSSTGAVTSGNYPNWVQKPKGQAPRGLIGG TKFLAPGTPARFSGSLLGGKAALTLSGVQPEDE AEYYCVLWYSNRWVFGSGTKLTVL</p>
310.	<p>MS 15-B12 CC x I2M2 x G4 x scfc x G4 x CH3 26-E5 CC x I2M2 - Full Sequence</p>	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGSDIVMTQSPSSLSASVGDRVITICRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGLTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGLSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPKGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGDKTHTCPPCPAP LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGSGGGG GGGGSGGGGSGGGGSGGGGSDKHTHTCPPCP APPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL VQSGAEVKKPGASVKVCKASGYTFTSYWMH WVRQAPGQCLEWMGVIRTSTSYTIYNQKFKG RVTMTRDTSTSTVYMESSLRSEDVAVYYCAR SGPGYFDVWGQGTMTVTVSSGGGGSGGGGSGG GGSDIQLTQSPSFLSASVGDRVITICRASENIYS YLAWYQQKPGKAPKLLIYNAKTLAEGVPSRFS GSGSGTEFTLTITSSLPEDFATYYCQHNHYGTPY TFGCGTKLEIKSGGGGSEVQLVESGGGLVQPG GSLKLSAASGFTFNKYAINWVREAPGKGLEW VARIRSKYNNYATYYADAVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRNANFGTSYIS YFAYWGQGLTVTVSSGGGGSGGGGSGGGGSGG TVVTQEPLTVSPGGTVTLTCGSSTGAVTSGNY PNWVQKPKGQAPRGLIGGTKFLAPGTPARFSG SLLGGKAALTLSGVQPEDEAEYYCVLWYSNR WVFGSGTKLTVL</p>
311.	<p>MS 15-B12 CC x I2M2 x G4S3 x</p>	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS</p>

	<p>scFc x G4S3 x CH3 22-A12 CC x I2M2 - Full Sequence</p>			<p>RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNKLTEDTAVYYCVRNANFGTS YISYFAYWGQGTLLTVVSSGGGGSGGGGSGGG GSQTVVTQEPSTVSPGGTVTLTCGSSTGAVTS GNYPNWWVQKPGQAPRGLIGGKFLAPGTPAR FSGSLLGGKAALTLGSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELGGPSVFLFPPKPKDTLMISRT PEVTCVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELGGPSVFLFPPKPKDTLMIS RTPEVTCVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVVKPG ASVKVSCKASGYTFTSSWMNWRQAPGQCLE WMGRIYTGGETKYSKGFQGRVTITRDTAST AYMELSSLTSEDVAVYYCARQRDYGALYAMD YWGQGTLLTVVSSGGGGSGGGGSGGGGSDIQL TQSPSFLSASVGDRVTITCRASDDIYSYLAWYQ QKPGKAPKLLVYNAKTLAEGVPSRFSGSGSGT EFTLITSSLPEDFATYYCQNHDRTPFTFGCGT KVDIKSGGGGSEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAINWVREAPGKGLEWVARIRS KYNNYATYYADAVKDRFTISRDDSKNTAYLQ MNKLTEDTAVYYCVRNANFGTSYISYFAYW GQGTLLTVVSSGGGGSGGGGSGGGGSGTQVVTQ EPSTVSPGGTVTLTCGSSTGAVTSGNYPNWW QKPGQAPRGLIGGKFLAPGTPARFSGSLLGG KAALTLGSGVQPEDEAEYYCVLWYSNRWVFGS GTKLTVL</p>
<p>312.</p>	<p>MS 15-B12 CC x I2M2 x G4S3 x scFc x G4S3 x CH3 005-D5 CC x I2M2 - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD</p>

				<p>SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPVLDSGDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPVLDSGDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSEVQLLESGGGLVQPG GSLRLSCAASGFSFSSYPINWVRQAPGKCLEW VGVWTGGGTNYASSVKGRFTISRDNKNTVY LQMNSLRAEDTAVYYCAKSRGVYDFKGRGA MDYWGQGLTVTVSSGGGGSGGGGSGGGGSDI VMTQSPDSLAVSLGERATINCKSSQSLLYSSNQ KNYFAWYQKPGQPPKLLIYWASTRESGVPDR FSGSGSGTDFTLTISLQAEDVAVYYCQQYYSY PYTFGCGTKLEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAINWVREAPGKGL EWVARIRSKYNNYATYYADAVKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRNANFGTSY ISYFAYWGQGLTVTVSSGGGGSGGGGSGGGGS QTVVTQEPLTVSPGGTVTLTCGSSTGAVTSGN YPNWVQKPGQAPRGLIGGTKFLAPGTPARFS GSLGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGSGTKLTVL</p>
313.	MS 15-B12 CC x I2M2 x G4S3 x scFc x G4S3 x CH3 26-E5 CC x I2M2 - Full Sequence	artificial	aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFTLTISLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA</p>

				<p>KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTSYWMHWVRQAPGQCLE WMGVIRTSTSYTIYNQKFKGRVTMTRDTSTST VYMELSSLRSEDNAVYYCARSQPGYFDVWGQ GTMVTVSSGGGGSGGGGSGGGGSDIQLTQSPS FLSASVGDRTITCRASENIYSYLAWYQQKPG KAPKLLIYNAKTLAEGVPSRFSGSGSGTEFTLTI SSLQPEDFATYYCQHNYGTPYTFGCGTKLEIKS GGGGSEVQLVESGGGLVQPGGSLKLSAASGF TFNKYAINWVREAPGKGLEWVARIRSKYNNY ATYYADAVKDRFTISRDDSKNTAYLQMNNLK TEDTAVYYCVRNANFGTSYISYFAYWGQGT LTVSSGGGGSGGGGSGGGGSGTQVVTQEPSLTV SPGGTVTLTCGSSTGAVTSGNYPNWVQKKPGQ APRGLIGGTKFLAPGTPARFSGSLLGGKAALTL SGVQPEDEAEYYCVLWYSNRWVFGSGTKLTV L</p>
314.	<p>MS 15-B12 CC x I2M2 xG4S3 x scFc x G4S3 - CH3 24-D7CC x I2M2 - Full Sequence</p>	artificial	aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGGTDFLTITSSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGT LTVSSGGGGSGGGGSGGG GSQTQVVTQEPSLTVSPGGTVTLTCGSSTGAVT SGNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTL SGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH</p>

				<p>NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSQVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIHSKAHGTNYNQKFQGRVTMTVDTSS STAYMELSRLSDDTAVYYCATRYFYVMDYW GQGTLVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVITICRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAINWVREAPGKGLEWVARIRSKYN NYATYYADAVKDRFTISRDDSKNTAYLQMNN LKTEDTAVYYCVRNANFGTSYISYFAYWGQG TLVTVSSGGGGSGGGGSGGGGSQTVVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQKKP GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGSGTKL TVL</p>
<p>315.</p>	<p>MS 15-B12 CCx I2C 44/100cc x scFc x CH3 15- E11 CC x I2C4/100cc0 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVITICRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK CLEWVARIRSKYNNYATYYADSVKDRFTISR DSKNTAYLQMNNLKTEDTAVYYCVRHGNFGN SYISYWAYWGQGTLVTVSSGGGGSGGGGSGG GGSTVVTQEPSLTVSPGGTVTLTCGSSTGAVT SGNYPNWVQKPGQAPRGLIGGTKFLAPGTPA RFGSLLGGKAALTLSGVQPEDEAEYYCVLWY SNRWVFGCGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSQVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFQGRVTMTVDTSS</p>

				<p>STAYMELSRRLSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKCLEWVARIRSKYN NYATYYADSVKDRFTISRDDSKNTAYLQMN LKTEDTAVYYCVRHGNFGNSYISYWAYWGQ TLTVTVSSGGGGSGGGGSGGGGSGTQVVTQEP SLTVSPGGTVTLTCGSSTGAVTSGNYPNWV QQKPGQAPRGLIGGTKFLAPGTPARFSGSLL GGKAALTLSGVQPEDEAEYYCVLWYSNRWV FGCGTKLTVL</p>
<p>316.</p>	<p>MS 25-E3 CCx 6H10.09x (G4)x scFcx (G4)x CH3 22-A12 CCx 6H10.09 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWWWIRQPPGKCLEWIGSIYYSGSTYYNPSLKS RVTISVDTSKNQFSLKLNSVTAADTAVYYCAR LPRGDRMTFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSEIVLTQSPGTLSPGERATLSCRASQS VSSSYLAWYQQKPGQAPRLLIYGASSRATGIPD RFSGSGSGTDFTLTISRLEPEDFAVYYCQQYGS SPFTFGCGTKLEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSCAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTQVVTQEPSTVSPGGTVTITCGSSTGAVTS GNYPNWIQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLEGGKAALTLSGVQPEDEAEYYCVLYYS NRWVFGSGTKLTVLGGGGDKTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVDVVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQVQL VQSGAEVVKPGASVKVSCKASGYTFTSSWMN WVRQAPGQCLEWMGRIYTGGETKYSKGFQ RVTITRDTASTAYMELSSLTSEDVAVYYCARQ RDYGALYAMDYWGQGLTVTVSSGGGGSGGGG SGGGGSDIQLTQSPSFLSASVGDRVTITCRAS DDIYSYLAWYQQKPGKAPKLLVYNAKTLAEG VPSRFSGSGSGTEFTLTISSLQPEDFATYYCQNH DRTPFTFGCGTKVDIKSGGGGSEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKGMWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTLYLQMNNLKTEDTAVYYCVRAGN</p>

				FGSSYISYFAYWGQGLTVTVSSGGGGSGGGGS GGGGSQTVVTQEPSLTVSPGGTVTITCGSSTGA VTSGNYPNWIQKKPGQAPRGLIGGTKFLAPGT PARFSGSLEGGKAALTLSGVQPEDEAEYYCVL YYSNRWVFGSGTKLTVL
317.	MS 25-E3 CCx 6H10.09x (G4S)3x scFcx (G4S)3x CH3 22- A12 CCx 6H10.09 - Full Sequence	artificial	aa	QVQLQESGPGLVKPSSETLSLTCTVSSGGSISSSSY FVWWIRQPPGKCLEWIGSIYYSGSTYYNPSLKS RVTISVDTSKNQFSLKLNsvTAADTAVYYCAR LPRGDRMTFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSEIVLTQSPGTLSPGERATLSCRASQS VSSYLAWYQKPGQAPRLLIYGASSRATGIPD RFSGSGSGTDFTLTISRLEPEDFAVYYCQQYGS SPFTFGCGTKLEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTITCGSSTGAVTS GNYPNWIQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLEGGKAALTLSGVQPEDEAEYYCVL YYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVVKPG ASVKVSCKASGYTFTSSWMNWVRQAPGQCLE WMGRIYTGTGETKYSKGFQGRVTITRDTAST AYMELSSLTSED TAVYYCARQRDYGALYAMD YWGQGLTVTVSSGGGGSGGGGSGGGGSDIQL TQSPSFLSASVGDRVTITCRASDDIYSYLAWYQ QKPGKAPKLLVYNAKTLAEGVPSRFSGSGSGT EFTLTISSLQPEDFATYYCQNHDRTPFTFGCGT KVDIKSGGGGSEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAMNWVRQAPGKGMWVARI RSKYNNYATYYADAVKDRFTISRDDSNTLYL QMNNLKTEDTAVYYCVRAGNFGSSYISYFAY WGQGLTVTVSSGGGGSGGGGSGGGGSGQTVVT QEPSLTVSPGGTVTITCGSSTGAVTSGNYPNWI QKKPGQAPRGLIGGTKFLAPGTPARFSGSLEGG KAALTLSGVQPEDEAEYYCVL YYSNRWVFGS GTKLTVL
318.	MS 46-A3 CC x I2Ccc(44/100)x (G4S)3x scFcx	artificial	aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMGWVRQAPGKCLEWVAVISYHGSNKYYAD AVKGRFTISRDN SKNTLYLQMNSLRAEDTAVY

	(G4S)3x CH3 005-D5 CCx I2Ccc(44/100) - Full Sequence			<p>YCAREGAHFGSGSYYPYYYYAMDVWGQGT TVTSSGGGGSGGGGGSGGGGSEIVTQSPGTL SPGERATLSCRASQSVSSSYLAWYQQKPGQAP RLLIYGASIRATGIPDRFSGSGSGTDFTLTISRLE PEDFAVYYCQQTGSSPIFTFGCGTKVEIKSGGG GSEVQLVESGGGLVQPGGSLKLSAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGLVTV SSGGGGSGGGGGSGGGGSQTVVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQQKPGQAP RGLIGGTKFLAPGTPARFSGSLLGGKAALTL VQPEDEAEYYCVLWYSNRWVFGCGTKLTVL GGGSGGGGGSGGGGSDKTHTCPPCPAPPELLGGP SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDP EVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC VSVLTVLHQDWLNGKEYKCKVSNKALPAPIEK TISKAKGQPREPQVYTLPPSREEMTKNQVSLT LVKGFYPSDIAVEWESNGQPENNYKTTTPVLD SDGSFFLYSKLTVDKSRWQQGNVFCFSVMHEA LHNHYTQKSLSLSPGKGGGGSGGGGGSGGGG GGGGSGGGGGSGGGGSDKTHTCPPCPAPPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHED PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTTPVLD DSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHE ALHNHYTQKSLSLSPGKGGGGSGGGGGSGGGG SEVQLLESVGGGLVQPGGSLRLS AASGFSFSSY PINWVRQAPGKCLEWVGIWVGGGTNYASSV KGRFTISRDNKNTVYLQMNSLRAEDTAVYYC AKSRGVYDFKGRGAMDYWGQGLVTVSSGG GGSGGGGGSGGGGSDIVMTQSPDSLAVSLGERA TINCKSSQSLLYSSNQKNYFAWYQQKPGQPPK LLIYWASTRESGVPDRFSGSGSGTDFTLTISSLQ AEDVAVYYCQQYYSYPYTFGCGTKLEIKSGGG GSEVQLVESGGGLVQPGGSLKLSAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGLVTV SSGGGGSGGGGGSGGGGSQTVVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQQKPGQAP RGLIGGTKFLAPGTPARFSGSLLGGKAALTL VQPEDEAEYYCVLWYSNRWVFGCGTKLTVL</p>
319.	MS R4L CC x I2C 44/100cc x scFc x CH3 R170R CC x I2C4/100cc0 - Full Sequence	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTG YYIHVVRQAPGQCLEWMGWINPNSGGTNYA QKFQGRVTMTRDTSISTAYMELSRRLSDDTAV YYCARVEAVAGREYYYFSGMDVWGQGTTVT VSSGGGGSGGGGGSGGGGSSYELTQPPSVSVP GQTASITCSGEKLGDKYVYVYQQKPGQSPVL VIYQSTKRPSGVPERFSGSNSGNTATLTISGTQA MDEADYYCQAYHASTAVFGCGTKLTVLSGGG GSEVQLVESGGGLVQPGGSLKLSAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED</p>

				<p>TAVYYCVRHGNFGNSYISYWAYWGQGLTVTV SSGGGGSGGGGSGGGGSQTVVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQQKPGQAP RGLIGGTKFLAPGTPARFSGSLLGGKAALTLG VQPEDEAEYYCVLWYSNRWVFGCGTKLTVLG GGGSGGGGSGGGGSDKTHTCPPCPAPPELLGGP SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDP EVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC VSVLTVLHQDWLNGKEYKCKVSNKALPAPIEK TISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTPPVLDS DGSFFLYSKLTVDKSRWQQGNVFSCSVMHEA LHNHYTQKSLSLSPGKGGGGSGGGGSGGGGS GGGGSGGGGSGGGGSDKTHTCPPCPAPPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHED PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHE ALHNHYTQKSLSLSPGKGGGGSGGGGSGGGG SEVQLLESGLLVQPGGSLRLSCAASGFTFSSY WMHWVRQTPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDKSKNTLYLQMNSLRAEDTAVY YCARWDYSHFDVWGQGTTVTVSSGGGGSGG GGGGGGSEIVMTQSPATLSVSPGERATLTCRA SSSVSYMHWYQQKPGQAPRLLIYGTSLVSGV PARFSGSGSGTEFTLTISSLQSEDFAVYYCQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGLTVTVSSGGGGSGGGGSG GGGGSGGGGSGGGGSDKTHTCPPCPAPPELLGG PAVTSGNYPNWVQQKPGQAPRGLIGGTKFLAPG TPARFSGSLLGGKAALTLGSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVL</p>
<p>320.</p>	<p>MS R4L CC x I2Ccc(44/100)x (G4S)3x scFex (G4S)3x CH3 08- A11 CCx I2Ccc(44/100) - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTG YYIHVWRQAPGQCLEWMGWINPNSGGTNYA QKFQGRVTMTRDTSISTAYMELSRLSDDTAV YYCARVEAVAGREYYYFSGMDVWGQGTTVT VSSGGGGSGGGGSGGGGSSYELTQPPSVSVP GQTASITCSGEKLGDKYVYVYQQKPGQSPVL VIYQSTKRPSGVPERFSGSNSGNTATLTISGTQA MDEADYYCQAYHASTAVFGCGTKLTVLSGGG GSEVQLVESGGGLVQPGGSLKLSCAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGLTVTV SSGGGGSGGGGSGGGGSGGGGSGGGGSGGGG SGGGSGGGGSGGGGSDKTHTCPPCPAPPELLGGP SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDP EVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC</p>

				<p>VSVLTVLHQDWLNGKEYKCKVSNKALPAPIEK TISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHEALHNHYTQKSLSLSPGKGGGGSGGGSGGGSGGGSGGGSGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHEALHNHYTQKSLSLSPGKGGGGSGGGSGGGSGGGSEVQLLESGLVQPGGSLRLSCAASGFTFSSYWMHWVRQTTPGKCLEWVSKIDPSDDYTNYNQKVKGRFTISIDKSKNTLYLQMNSLRAEDTAVYYCARWDYNYFDVWGQGTITVTVSSGGGGSGGGGGGGSEIVMTQSPATLSVSPGERATLTCRASSVSVMHWYQQKPGQAPRLLIYGTSNLSGVPARFSGSGTEFTLTISSLQSEDFAVYYCQQWSSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGLVQPGGSLKLSCAASGFTFNKYAMNWRQAPGKCLEWVARIRSKYNNYATYYADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAVYYCVRHGNFGNSYISYWAYWGQGTITVTVSSGGGGSGGGGGSGGGGQTIVVTQEPSLTVSPGGTVTLTLCGSSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAALTLGSGVQPEDEAEYYCVLWYSNRWVFGCGTKLTVL</p>
321.	MS R4L CC x I2Ccc(44/100)x (G4S)3x scFex (G4S)3x CH3 R164L CCx I2Ccc(44/100) - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTGYYIHWVRQAPGQCLEWMGWINPNSGGTNYAQKFQGRVTMTRDTSISTAYMELSRLSDDTAVYYCARVEAVAGREYYYFSGMDVWGQGTITVTVSSGGGGSGGGSGGGSSYELTQPPSVSVPGQTASITCSGEKLGDKYVYVYQQKPGQSPVLVIYQSTKRPSGVPERFSGSNSGNTATLTISGTQAMDEADYYCQAYHASTAVFGCGTKLTVLSGGGGSEVQLVESGGGLVQPGGSLKLSCAASGFTFNKYAMNWRQAPGKCLEWVARIRSKYNNYATYYADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAVYYCVRHGNFGNSYISYWAYWGQGTITVTVSSGGGGSGGGSGGGGQTIVVTQEPSLTVSPGGTVTLTLCGSSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAALTLGSGVQPEDEAEYYCVLWYSNRWVFGCGTKLTVLGGGGGGGGSGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHEALHNHYTQKSLSLSPGKGGGGSGGGSGGGSGGGSEVQLLESGLVQPGGSLKLSCAASGFTFNKYAMNWRQAPGKCLEWVARIRSKYNNYATYYADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAVYYCVRHGNFGNSYISYWAYWGQGTITVTVSSGGGGSGGGSGGGGQTIVVTQEPSLTVSPGGTVTLTLCGSSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAALTLGSGVQPEDEAEYYCVLWYSNRWVFGCGTKLTVLGGGGGGGGSGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC</p>

				<p>CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHE ALHNHYTQKSLSLSPGKGGGGSGGGGSGGGG SEVQLLESGLVQPGGSLRLSCAASGFTFSSY WMYWRQAPGKCLEWVSKIDPSDDYTNYNQ KVKGRFTISIDNSKNTLYLQMNSLRAEDSAVY YCARWDYTHFDVWGQGTITVTVSSGGGGSGG GGSGGGSEIVMTQSPATLSVSPGERATLSCRA SSVSVMHWYQQKPGQAPRLLIYGTSNLAAGV PVRFGSGSGTEFTLTISRLQSEDAVYYCQQW SSYPLTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGTITVTVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGTVTLTCGSSTGA VTSGNYPNWVQQKPGQAPRGLIGGKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVL</p>
<p>322.</p>	<p>MS R4L CCx I2Ccc(44/100)x (G4)x scFc x (G4) x CH3 08-A11 CCx I2Ccc(44/100) - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTG YYIHWRQAPGQCLEWMGWINPNSGGTNYA QKFQGRVTMTRDTSISTAYMELSRLRSDDTAV YYCARVEAVAGREYYYFSGMDVWGQGTITV VSSGGGGSGGGGSGGGGSSYELTQPPSVSVP GQTASITCSGEKLGDKYVYVYQQKPGQSPVL VIYQSTKRPSGVPERFSGSNSGNTATLTISGTQA MDEADYYCQAYHASTAVFGCGTKLTVLSSGG GSEVQLVESGGGLVQPGGSLKLSCAASGFTFN KYAMNWRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGTITV SSGGGGSGGGGSGGGGSQTVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQQKPGQAP RGLIGGKFLAPGTPARFSGSLLGGKAALTLSG VQPEDEAEYYCVLWYSNRWVFGCGTKLTVLGG GGDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNGKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLDSDGSFFLYSK LTVDKSRWQQGNVFCFSVMHEALHNHYTQKS LSLSPGKGGGGSGGGGSGGGGSGGGGSGGGG SGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK KDTLMISRTPEVTCVVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSTYRCVSVLTVLH QDWLNGKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSDGSFFLYS KLTVDKSRWQQGNVFCFSVMHEALHNHYTQK LSLSPGKGGGGSEVQLLESGLVQPGGSLRLS CAASGFTFSSYVMHWVRQTPGKCLEWVSKID PSDDYTNYNQKVKGRFTISIDKSKNTLYLQMN SLRAEDTAVYYCARWDYNYFDVWGQGTITV</p>

				VSSGGGGSGGGGSGGGGSEIVMTQSPATLSVS PGERATLTCRASSSVSYMHWYQQKPGQAPRLL IYGTSNLVSGVPARFSGSGSGTEFTLTISLQSE DFAVYYCQQWSSYPLTFGCGTKVEIKSGGGGS EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKCLEWVARIRSKYNNYATYY ADSVKDRFTISRDDSKNTAYLQMNNLKTEDA VYYCVRHGNFGNSYISYWAYWGQGLTVTVSS GGGGSGGGGSGGGGSQTVVTQEPSLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQKPGQAPRG LIGGTKFLAPGTPARFSGSLLGGKAALTLGSGVQ PEDEAEYYCVLWYSNRWVFGCGTKLTVL
323.	MS15-B12 CC x I2M2 x (G4Q)3x scF _{cm} od x (G4Q)3 x CH3 24- D7 CC x I2M2 - Full Sequence	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGG QGGGGQDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFTLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGQEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDA VYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGQGGGGQGGG GQQTVVTQEPSLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLGSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGQGGGGQGGGGQC PPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHEEPEVKFNWYVDGVEVHNAKTKP CEEQYGSTYRCVSVLTVLHQQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDSGDSFFLYSKLTVDKSRWQQG NVFSCSVMHEALHNHYTQKSLSLSPGKGGGG QGGGGQGGGGQGGGGQGGGGQGGGGQC PPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVV VDVSHEEPEVKFNWYVDGVEVHNAKTKPCEE QYGSTYRCVSVLTVLHQQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSGDSFFLYSKLTVDKSRWQQGNV FSCSVMHEALHNHYTQKSLSLSPGKGGGGQ GGGQGGGGQVQLVQSGAEVKKPGASVKVSC KASGYTFTNYWMNWVRQAPGQCLEWMGNIH SKAHGTNYNQKFQGRVTMTVDTSSSTAYMEL SRLRSDDTAVYYCATRYFYVMDYWGQGLTVT VSSGGGGQGGGGQGGGGQDIQMTQSPSSLSAS VGDRVTITCRASQDISNYLNWYQQKPGKVPKL LIYYTSRLHSGVPSRFSGSGSGTDFTLTISLQPE DVATYYCVQYAQFPLTFGCGTKVEIKSGGGGQ EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AINWVREAPGKGLEWVARIRSKYNNYATYYA DAVKDRFTISRDDSKNTAYLQMNNLKTEDA VYYCVRNANFGTSYISYFAYWGQGLTVTVSSG GGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT

				VTLTCGSSTGAVTSGNYPNWVQKKPGQAPRG LIGGTKFLAPGTPARFSGSLLGGKAALTLSGVQ PEDEAEYYCVLWYSNRWVFGSGTKLTVL
324.	heFc(A) x(G4)x MS 15-B12 CCx 6H10.09	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYDTTPPVLDSDGSFFLYSCLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG KGGGGQVQLQESGPGLVKPSETLSLTCTVSGG SISSSSYFWGWIRQPPGKCLEWIGNIYYSGSSN YNPSLKSRTISVDTSKNQFSLKLSVTAADTA VYYCARLPRGDRDAFDIWGQGMVTVSSGGG GSGGGGSGGGSDIVMTQSPSSLSASVGDRVTI TCRASQGISNYLAWYQQKPGKVPKLLIYAAS LQSGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQSYSTPFTFGCGTKVEIKSGGGGSEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGLTVTVSSGGGGSGG GGSGGGGSQTVVTQEPSLTVSPGGTVTITCGSS TGAVTSGNYPNWIQKKPGQAPRGLIGGTKFLA PCTPARFSGSLEGGKAALTLSGVQPEDEAEYY CVLYYSNRWVFGSGTKLTVL
325.	CH3 15-E11 CC 6H10.09 x (G4)x heFc(B)	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGVKGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGG GGGGSGGGSDIQMTQSPSSLSASVGDRVTITC RASQDISNYLNWYQQKPGKVPKLLIYYTSRLH SGVPSRFSGSGSGTDFTLTISSLQPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSEVQLVESG GGLVQPGGSLKLSAASGFTFNKYAMNWVRQ APGKGMWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVR GNFGSSYISYFAYWGQGLTVTVSSGGGGSGGG GSGGGGSQTVVTQEPSLTVSPGGTVTITCGSST GAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAP GTPARFSGSLEGGKAALTLSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVLGGGGDKTHTCPP CPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VVDVSHEDPEVKFNWYVDGVEVHNAKTKPCE EQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSRKE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKLSLSLSPGK
326.	heFc(B) x (G4)x CH3 15-E11 CCx 6H10.09	artificial	Aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDK SRWQQGNVFSCSVMHEALHNHYTQKLSLSLSP

				GKGGGGQVQLVQSGAEVKKPGASVKVSCKAS GYTFTNYWMNWVRQAPGQCLEWMGNIAYGV KGTNYNQKFQGRVTMTVDTSSSTAYMELSRL RSDDTAVYYCATRYFYVMDYWGQGLVTVSS GGGGSGGGGSGGGGSDIQMTQSPSSLSASVGD RVTITCRASQDISNYLNWYQQKPGKVPKLLIY YTSRLHSGVPSRFSGSGSGTDFTLTISSLQPEDV ATYYCVQYAQFPLTFGCGTKVEIKSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKGMWVARIRSKYNNYATYYAD AVKDRFTISRDDSKNTLYLQMNNLKTEDTAVY YCVRAGNFGSSYISYFAYWGQGLVTVSSGGG GSGGGGSGGGGSQT VVTQEPSLTVSPGGTVTIT CGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGT KFLAPGTPARFSGSLEGGKAALTLSGVQPEDEA EYYCVLYYSNRWVFGSGTKLTVL
327.	MS 15-B12 CCx 6H10.09x (G4)x heFc(A)	artificial	Aa	QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLVTVSSGGGGSGGGGSGG GGSQT VVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGT KFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGDKTHTCPPCPAPE LLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYG STYRCVSVLTVLHQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTKN QVSLTCLVKGFYPSDIAVEWESNGQPENNYDT TPPVLDSDGSFFLYSDLTVDKSRWQQGNVFC SVMHEALHNHYTQDSLSPGK
328.	H MS 15-B12 x H 6H10.09 x (G4S)3 x heFc(A) x (G4S)3 x H CH3 15-E11 x H 6H10.09	artificial	Aa	QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSGGGGSEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAMNWVRQAPGKGMWVARI RSKYNNYATYYADAVKDRFTISRDDSKNTLYL QMNNLKTEDTAVYYCVRAGNFGSSYISYFAY WGQGLVTVSSGGGGSGGGGSGGGGSDKTH CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEDPEVKFNWYVDGVEVHNAKT KPCEEQYGSTYRCVSVLTVLHQDWLNGKEYK CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYDTTPPVLDSDGSFFLYSDLTVDKSRW QQGNVFCSSVMHEALHNHYTQDSLSPGK GGGSGGGGSGGGGSQVQLVQSGAEVKKPGAS

				VKVSCKASGYTFTNYWMNWVRQAPGQCLEW MGNIAYGVKGTNYNQKFQGRVTMTVDTSSST AYMELSRLRSDDTAVYYCATRYFYVMDYWG QGTLVTVSSGGGGSGGGGSGGGGSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKGMWVARIRSKYNNYATYYAD AVKDRFTISRDDSKNTLYLQMNCLKTEDITAVY YCVRAGNFGSSYISYFAYWGGTLTVSS
329.	L MS 15-B12 x L 6H10.09 x (G4S)3 x heFc(B) x (G4S)3 x L CH3 15-E11 x L 6H10.09	artificial	Aa	DIVMTQSPSSLSASVGDRVITICRASQGISNYL AWYQQKPKGKVPKLLIYAASLTQSGVPSRFSGS GSGTDFLTITSSLPEDFATYYCQQSYSTPFTFG CGTKVEIKSGGGGSGGGGSGGGGSGGGGSSQT VVTQEPSLTVSPGGTVTITCGSSTGAVTSGNYP NWIQKKPGQAPRGLIGGTFKFLAPGTPARFSGSL EGGKAALTLSGVQPEDEAEYYCVLYYSNRWV FGSGTKLTVLGGGGSGGGGSGGGGSDKTHTCP PCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTC VVVDVSHEDPEVKFNWYVDGVEVHNAKTKPC EEQYGSTYRCVSVLTVLHQDWLNGKEYKCKV SNKALPAPIEKTISKAKGQPREPVYTLPPSRKE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGSG GGGSGGGGSDIQMTQSPSSLSASVGDRVITICR ASQDISNYLNWYQQKPKGKVPKLLIYYTSRLHS GVPSRFSGSGSGTDFLTITSSLPEDVATYYCV QYAQFPLTFGCGTKVEIKSGGGGSGGGGSGGG GSGGGGSSQT VVTQEPSLTVSPGGTVTITCGSST GAVTSGNYPNWIQKKPGQAPRGLIGGTFKFLAP GTPARFSGSLEGGKAALTLSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVL
330.	IgG1 subtype hinge	artificial	aa	DKTHTCP
331.	IgG2 subtype hinge	artificial	aa	ERKCCVEC
332.	IgG3 subtype hinge	artificial	aa	ELKTPLDTHTCP
333.	IgG3 subtype hinge	artificial	aa	ELKTPLGDTTHTCP
334.	EpCAM 5-10 LH - HCDR1	artificial	aa	NYWLG
335.	EpCAM 5-10 LH - HCDR2	artificial	aa	DIFPGSGNIHYNEKFKG
336.	EpCAM 5-10 LH - HCDR3	artificial	aa	LRNWDEPMDY
337.	EpCAM 5-10 LH - LCDR1	artificial	aa	KSSQSLLNSGNQKNYLT
338.	EpCAM 5-10 LH - LCDR2	artificial	Aa	WASTRES
339.	EpCAM 5-10 LH - LCDR3	artificial	Aa	QNDYSYPLT
340.	EpCAM 5-10 LH - VH	artificial	Aa	EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF K GKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVTVSS

341.	EpCAM 5-10 LH - VL	artificial	Aa	ELVMTQSPSSLT VTAGEKVTMSCKSSQSL LNS GNQKNYLTWYQQKPGQP KLLIYWASTRESG VPDRFTGSGSGTDFTL TISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIK
342.	EPCAM 5-10 x scFc x H2 x I2Ccc x I2Ccc - Full Sequence	artificial	Aa	ELVMTQSPSSLT VTAGEKVTMSCKSSQSL LNS GNQKNYLTWYQQKPGQP KLLIYWASTRESG VPDRFTGSGSGTDFTL TISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGT SVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSS LTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVTVSSGGGGDKT HTPPCPAPELLGGPSVFL FPPKPKDTLMIS RTP EVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVL TVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTP EVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVL TVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGEVQLLES GGGVVPGRSLRLS CAASGF TFSSYGMGWVRQAPGKGLEWVAVISYDGSNK YYADSVKGRFTISRDN SKNTLYLQMNSLRAED TAVYYCAREGAHFGSGSY YPLYYYYAMDVW GQGT TVTVSSGGGGSGGGGSGGGGSELT LTQS PGTLSPGERATLSCRASQSVSSSYLAWYQQK PGQAPRLLIYGASIRATGIPDRFSGSGSGTDFTL TISRLEPEDFAVYYCQQY GSSPIFTFGPGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLS CAAS GFTFNKYAMNWVRQAPGKCLEWVARIRSKYN NYATYYADSVKDRFTISRDDSKNTAYLQMN N LKTEDTAVYYCVRHGNFGNSYISYWAYWGQG TLVTVSSGGGGSGGGGSGGGGSQT VVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQQK P GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGCGTKL TVLGGGGSGGGGSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAMNWVRQAPGKC LEWVARIRSKYNNYATYYADSVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRHGNFGNS YISYWAYWGQGLTVTVSSGGGGSGGGGSGGG GSQT VVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYS NRWVFGCGTKLTVL
343.	EPCAM 5-10 x H2 x scFc x I2Ccc x I2Ccc - Full Sequence	artificial	Aa	ELVMTQSPSSLT VTAGEKVTMSCKSSQSL LNS GNQKNYLTWYQQKPGQP KLLIYWASTRESG VPDRFTGSGSGTDFTL TISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS

				<p>EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTTVTVSSGGGGSEV QLLESGGGVVQPGRSLRLSCAASGFTFSSYGM GWVRQAPGKGLEWVAVISYDGSNKYYADSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AREGAHFGSGSYYPYLYYYAMDVWGQGTTV TVSSGGGGSGGGGSGGGGSELTLTQSPGTLSL PGERATLSCRASQSVSSYLAWYQQKPGQAPR LLIYGASIRATGIPDRFSGSGGTDFTLTISRLEP EDFAVYYCQQYGSSPIFTFGPGTKVEIKSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGEVQLVESGGGLVQPGGSLKLSAASGF TFNKYAMNWVRQAPGKCLEWVARIRSKYNN YATYYADSVKDRFTISRDDSKNTAYLQMNNL KTEDTAVYYCVRHGNFGNSYISYWAYWGQGT LTVVSSGGGGSGGGGSGGGGSGTQVVTQEPSLT VSPGGTVTLTSGSSTGAVTSGNYPNWVQQKPG QAPRGLIGGTKFLAPGTPARFSGSLLGGKAALT LSGVQPEDEAEYYCVLWYSNRWVFGGTKLT VLGGGGSGGGGSGGGGSEVQLVESGGGLVQP GGSLKLSAASGFTFNKYAMNWVRQAPGKCL EWWARIRSKYNNYATYYADSVKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRHGNFGNSY ISYWAYWGQGTTLTVVSSGGGGSGGGGSGGGG SQTQVVTQEPSLTVSPGGTVTLTSGSSTGAVTSG NYPNWVQQKPGQAPRGLIGGTKFLAPGTPARF SGSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGGTKLTVL</p>
344.	EPCAM 5-10 x H2 x I2Ccc x scFc x I2Ccc - Full Sequence	artificial	Aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGGTDFTLTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTTVTVSSGGGGSEV QLLESGGGVVQPGRSLRLSCAASGFTFSSYGM GWVRQAPGKGLEWVAVISYDGSNKYYADSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AREGAHFGSGSYYPYLYYYAMDVWGQGTTV</p>

				<p>TVSSGGGGSGGGGSGGGGSELTLTQSPGTL SLS PGERATLSCRASQSVSSSYLAWYQQKPGQAPR LLIYGASIRATGIPDRFSGSGSDFTLTISRLEP EDFAVYYCQQYGSSPIFTFGPGTKVEIKSGGGG SEVQLVESGGGLVQPGGSLKLSCAASGFTFNK YAMNWVRQAPGKCLEWVARIRSKYNNYATY YADSVKDRFTISRDDSKNTAYLQMNNLKTEDT AVYYCVRHGNFGNSYISYWAYWGQGLTVTS SGGGSGGGGSGGGGSGTQVVTQEPSLTVSPGG TVTLTCGSSTGAVTSGNYPNWVQKPGQAPR GLIGGTKFLAPGTPARFSGSLLGGKAALTLGSGV QPEDEAEYYCVLWYSNRWVFGCGTKLTVLGG GGDKTHTCPPCPAPELLGGPSVFLFPPKPKDTL MISRTPEVTCVVVDVSHEDPEVKFNWYVDGV EVHNAKTKPCEEQYGSTYRCVSVLTVLHQDW LNGKEYKCKVSNKALPAPIEKTISKAKGQPREP QVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAV EWESNGQPENNYKTTTPVLDSDG SFFLYSKLT VDKSRWQQGNVFSCSVMHEALHNHYTQKSLS LSPGKGGGSGGGGSGGGGSGGGGSGGGGSG GGGSDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNGKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSDG SFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAMNWVRQAPGKCLEWVARIR SKYNNYATYYADSVKDRFTISRDDSKNTAYLQ MNLKTEDTAVYYCVRHGNFGNSYISYWAY WGQGLTVTVSSGGGSGGGGSGGGGSGTQVVT QEPSLTVSPGGTTLTCGSSTGAVTSGNYPNW VQKPGQAPRGLIGGTKFLAPGTPARFSGSLLG GKAALTLGSGVQPEDEAEYYCVLWYSNRWVFG CGTKLTVL</p>
345.	EPCAM 5-10 x H2 x I2Ccc x I2Ccc x scFc - Full Sequence	artificial	Aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSL LNS GNQKNYL TWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGSGGGGSGGGG EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVTVSSGGGGSEV QLLESGGGVVQGRSLRLSCAASGFTFSSYGM GWVRQAPGKGLEWVAVISYDGSNKYYADSV KGRFTISRDN SKNTLYLQMNSLRAEDTAVYYC AREGAHFGSGSYYP LYYYYAMDVWGQGT TTV TVSSGGGSGGGGSGGGGSELTLTQSPGTL SLS PGERATLSCRASQSVSSSYLAWYQQKPGQAPR LLIYGASIRATGIPDRFSGSGSDFTLTISRLEP EDFAVYYCQQYGSSPIFTFGPGTKVEIKSGGGG SEVQLVESGGGLVQPGGSLKLSCAASGFTFNK YAMNWVRQAPGKCLEWVARIRSKYNNYATY YADSVKDRFTISRDDSKNTAYLQMNNLKTEDT AVYYCVRHGNFGNSYISYWAYWGQGLTVTS</p>

				<p>SGGGGSGGGGSGGGGSQTVVTQEPSLTVSPGG TVTLTCGSSTGAVTSGNYPNWVQKPGQAPR GLIGGTKFLAPGTPARFSGSLLGGKAALTLSGV QPEDEAEYYCVLWYSNRWVFGCGTKLTVLGG GGSGGGGSGGGGSEVQLVESGGGLVQPGGSL KLSAASGFTFNKYAMNWVRQAPGKCLEWV ARIRSKYNNYATYYADSVKDRFTISRDDSKNT AYLQMNNLKTEDTAVYYCVRHGNFGNSYISY WAYWGQGLTVTVSSGGGGSGGGGSGGGGSQ TVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGNY PNWVQKPGQAPRGLIGGTKFLAPGTPARFSG SLLGGKAALTLSGVQPEDEAEYYCVLWYSNR WVFGCGTKLTVLGGGDKTHTCPPCPAPPELLG GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH DPEVKFNWYVDGVEVHNAKTKPCEEQYGSTY RCVSVLTVLHQDWLNGKEYKCKVSNKALPAPI EKTISKAKGQPREPQVYTLPPSREEMTKNQVSL TCLVKGFYPSDIAVEWESNGQPENNYKTTPPV LDSGDSFFLYSKLTVDKSRWQQGNVFSCSVMH EALHNHYTQKSLSLSPGKGGGGSGGGGSGGG GSGGGGSGGGGSGGGGSDKTHTCPPCPAPELL GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH EDPEVKFNWYVDGVEVHNAKTKPCEEQYGST YRCVSVLTVLHQDWLNGKEYKCKVSNKALPA PIEKTISKAKGQPREPQVYTLPPSREEMTKNQV SLTCLVKGFYPSDIAVEWESNGQPENNYKTTPP VLDSGDSFFLYSKLTVDKSRWQQGNVFSCSVM HEALHNHYTQKSLSLSPGK</p>
346.	EPCAM 5-10 x I2C x scFc x I2C x H2 - Full Sequence	artificial	Aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYLTWYQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF K GKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTITVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKGLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLTVTVSSGG GGSGGGGSGGGGSQTVVTQEPSLTVSPGGTVT LTCGSSTGAVTSGNYPNWVQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTLSGVQPED EAEYYCVLWYSNRWVFGGKTKLTVLGGGGSG GGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLF PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPPVLDSDGSF FLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSGGGGS GGGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLF PPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA</p>

				<p>KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTPPVLDSDGSF FLYSKLTVDKSRWQQGNVFCVSMHEALNH YTQKSLSLSPGKGGGGSGGGGGSGGGGSEVQLV ESGGGLVQPGGSLKLSCAASGFTFNKYAMNW VRQAPGKGLEWVARIRSKYNNYATYYADSVK DRFTISRDDSKNTAYLQMNNLKTEDTAVYYCV RHGNFGNSYISYWAYWGQGLTVTVSSGGGGS GGGSGGGGSQTVVVTQEPSLTVSPGGTVTLTC GSSTGAVTSGNYPNWVQQKPGQAPRGLIGGTK FLAPGTPARFSGSLLGGKAALTLGSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGSEVQL LESGGGVVQPGRSLRLSCAASGFTFSSYGMGW VRQAPGKGLEWVAVISYDGSNKYYADSVKGR FTISRDNKNTLYLQMNSLRAEDTAVYYCARE GAHFGSGSYYPYLYYYAMDVWGQGTTVTVSS GGGSGGGSGGGGSELTLTQSPGTLSPGER ATLSCRASQSVSSSYLAWYQQKPGQAPRLLIY GASIRATGIPDRFSGSGSGTDFTLTISRLEPEDFA VYYCQQYGSSPIFTFGPGTKVEIK</p>
347.	EPCAM 5-10 x I2Ccc x H2 x I2Ccc - Full Sequence	artificial	aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSCAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLTVTVSSGG GSGGGGGSGGGGSQTVVVTQEPSLTVSPGGTVT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTLGSGVQPED EAEYYCVLWYSNRWVFGCGTKLTVLGGGGSE VQLLESGGGVVQPGRSLRLSCAASGFTFSSYG MGWVRQAPGKGLEWVAVISYDGSNKYYADS VKGRFTISRDNKNTLYLQMNSLRAEDTAVYY CAREGAHFGSGSYYPYLYYYAMDVWGQGT TVTVSSGGGGSGGGGGSGGGGSELTLTQSPGTL LSPGERATLSRASQSVSSSYLAWYQQKPGQA PRLLIYGASIRATGIPDRFSGSGSGTDFTLTISR LEPEDFAVYYCQQYGSSPIFTFGPGTKVEIKSGG GGSEVQLVESGGGLVQPGGSLKLSCAASGFTF NKYAMNWVRQAPGKCLEWVARIRSKYNNYA TYYADSVKDRFTISRDDSKNTAYLQMNNLKTE DTAVYYCVRHGNFGNSYISYWAYWGQGLTVT VSSGGGGSGGGGGSGGGGSQTVVVTQEPSLTVSP GGTVTLTCGSSTGAVTSGNYPNWVQQKPGQA PRGLIGGTKFLAPGTPARFSGSLLGGKAALTL GSVQPEDEAEYYCVLWYSNRWVFGCGTKLTVL</p>
348.	EPCAM 5-10 x I2Ccc x scFc x H2 x I2Ccc - Full Sequence	artificial	aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGGSGGGGS</p>

				<p>EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLTVTVSSGG GGSGGGSGGGGSQT VVTQEPSLTVSPGGT V T LTCGSSTGAVTSGNYPNWVQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTL SGVQPED EAEYYCVLWYSNRWVFGCGTKLTVLGGGGSG GGSGGGGSDKTHTCPPCPAPPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTTPVLDS DGSF FLYSKLTVDKSRWQQGNV FSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSGGGGS GGGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLF PPKPKDTLMISRTPEVTCVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTTPVLDS DGSF FLYSKLTVDKSRWQQGNV FSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSEVQLL ESGGGVVQPGRSLRLSCAASGFTFSSYGMGWV RQAPGKGLEWVAVISYDGSNKYYADSVKGRF TISRDN SKNTLYLQMNSLRAEDTAVYYCAREG AHFGSGSYYP LYYYYAMDVWGQGT TTVTVSSG GGSGGGGSGGGGSELTLTQSPGTLSLSPGERA TLSCRASQSVSSSYLAWYQKPGQAPRLLIYG ASIRATGIPDRFSGSGSDFTLTISRLEPEDFAV YYCQQYGS SPIFTFGPGTKVEIKSGGGGSEVQL VESGGGLVQPGGSLKLSAASGFTFNKYAMN WVRQAPGKCLEWVARIRSKYNNYATYYADSV KDRFTISRDDSKNTAYLQMNNLKTEDTAVYYC VRHGNFGNSYISYWAYWGQGLTVTVSSGGGG SGGGSGGGGSQT VVTQEPSLTVSPGGT V TLT CGSSTGAVTSGNYPNWVQKPGQAPRGLIGGT KFLAPGTPARFSGSLLGGKAALTL SGVQPEDEA EYYCVLWYSNRWVFGCGTKLTVL</p>
349.	EPCAM 5-10 x I2Ccc x scFc x I2Ccc x H2 - Full Sequence	artificial	aa	<p>ELVMTQSPSSLT VTAGEKVTMSCKSSQSLLNS GNQKNYLTWYQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFLT LTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLTVTVSSGG</p>

				<p>GGSGGGGSGGGGSQTVVTQEPSLTVSPGGTVT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTLSGVQPED EAEYYCVLWYSNRWVFGCGTKLTVLGGGGSG GGGSGGGGSDKTHTCPPCPAPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPPVLDSDGSF FLYSKLTVDKSRWQQGNVFSVMHEALHNH YTQKLSLSLSPGKGGGGSGGGGSGGGGSGGGGS GGGSGGGGSDKTHTCPPCPAPELLGGPSVFLF PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPPVLDSDGSF FLYSKLTVDKSRWQQGNVFSVMHEALHNH YTQKLSLSLSPGKGGGGSGGGGSGGGGSEVQLV ESGGGLVQPGGSLKLSCAASGFTFNKYAMNW VRQAPGKCLEWVARIRSKYNNYATYYADSVK DRFTISRDDSKNTAYLQMNNLKTEDTAVYYCV RHGNFGNSYISYWAYWGQGLTVTVSSGGGGG GGGSGGGGSQTVVTQEPSLTVSPGGTVTLTC GSSTGAVTSGNYPNWVQQKPGQAPRGLIGTK FLAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGSEVQL LESGGGVVQPGRSLRLSCAASGFTFSSYGMGW VRQAPGKGLEWVAVISYDGSNKYYADSVKGR FTISRDNKNTLYLQMNSLRAEDTAVYYCARE GAHFGSGSYYPYLYYYAMDVWGQGTTVTVSS GGGSGGGGSGGGGSELTTLTQSPGTLSPGER ATLSRASQSVSSSYLAWYQQKPGQAPRLIY GASIRATGIPDRFSGSGSGTDFTLTISRLEPEDFA VYYCQQYGSSPIFTFGPGTKVEIK</p>
<p>350.</p>	<p>EPCAM 5-10 x I2Cccx (G4S)10 x H2 x I2Ccc - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>ELVMTQSPSSLTVTAGEKVTMCKSSQSLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSCAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLTVTVSSGG GGGSGGGGSGGGGSQTVVTQEPSLTVSPGGTVT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTLSGVQPED EAEYYCVLWYSNRWVFGCGTKLTVLGGGGSG GGGSGGGGSGGGGSGGGGSGGGGSGGGGSGG GGGSGGGGSGGGGSEVQLLES GGGVVQPGRSLR LSCAASGFTFSSYGMGWVRQAPGKGLEWVA VISYDGSNKYYADSVKGRFTISRDNKNTLYLQ</p>

				<p>MNSLRAEDTAVYYCAREGAHFGSGSYYP YYAMDVWGQTTVTVSSGGGGSGGGGGSGGG GSELTLTQSPGTLSPGERATLSCRASQSVSS YLAWYQQKPGQAPRLLIYGASIRATGIPDRFSG SSGTDFTLTISRLEPEDFAVYYCQQYGSPIFT FGPGTKVEIKSGGGGSEVQLVESGGGLVQPGG SLKLSAASGFTFNKYAMNWRQAPGKCLEW VARIRSKYNNYATYYADSVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRHGNFGNSYIS YWAYWGQGLVTVSSGGGGSGGGGGSGGGGS QTVVTQEPLTVSPGGTVTLTCGSSTGAVTSGN YPNWVQQKPGQAPRGLIGGTKFLAPGTPARFS GSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGCGTKLTVL</p>
351.	<p>EPCAM 5-10 x I2Cccx G4Sx PD1xG4S x H2 x I2Ccc - Full Sequence</p>	artificial	aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQTTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLVTVSSGG GSGGGGGSGGGGSQTVVTQEPLTVSPGGTVT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTLSGVQPED EAEYYCVLWYSNRWVFGCGTKLTVLGGGGSL DSPDRPWNPTFSPALLVTEGDNATFTCSFSN TSEFVLNWRMSPSNQTDKLAAPEDRSQPG QDCRFRVTQLPNGRDFHMSVVRARRNDSGT YLCGAI LAPKAQIKESLRAELRVTERRAEVPTA HPSPPRPAGQFQGGGGSEVQLLESGGGVVQP GRSLRLS AASGFTFSSYGMGWVRQAPGKGLE WVAVISYDGSNKYYADSVKGRFTISRDN SKNT LYLQMNSLRAEDTAVYYCAREGAHFGSGSY PLYYYYAMDVWGQTTVTVSSGGGGSGGGG SGGGGSELTLTQSPGTLSPGERATLSCRASQ SVSSSYLAWYQQKPGQAPRLLIYGASIRATGIP DRFSGSGSGTDFTLTISRLEPEDFAVYYCQQY SSPIFTFGPGGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSAASGFTFNKYAMNWRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGLVTVSSGGGGSGGGGGSG GGGSQTVVTQEPLTVSPGGTVTLTCGSSTGA VTSNYPNWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVL</p>
352.	<p>EpCAM 5- 10_x(EAAAK)10 _x I2Ccc_xG4_xscFc _xG4_xMSLN_H</p>	artificial	aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY</p>

	<p>_x(EAAAK)10_x I2Ccc - Full Sequence</p>			<p>WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTTVTSSGEAAAKE AAAKEAAAKEAAAKEAAAKEAAAKEAAAKE AAAKEAAAKEAAAKEVQLVESGGGLVQPGGS LKLSCAASGFTFNKYAMNWVRQAPGKCLEWV ARIRSKYNNYATYYADSVKDRFTISRDDSKNT AYLQMNNLKTEDTAVYYCVRHGNFGNSYISY WAYWGQGTTLTVSSGGGGSGGGGSGGGGSQ TVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGNY PNWVQQKPGQAPRGLIGGTKFLAPGTPARFSG SLLGGKAALTLSGVQPEDEAEYYCVLWYSNR WVFGCGTKLTVLGGGGDKTHTCPPCPAPPELLG GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH DPEVKFNWYVDGVEVHNAKTKPCEEQYGSTY RCVSVLTVLHQDWLNGKEYKCKVSNKALPAPI EKTISKAKGQPREPQVYTLPPSREEMTKNQVSL TCLVKGFYPSDIAVEWESNGQPENNYKTTPPV LSDSGSFFLYSKLTVDKSRWQQGNVFCVSMH EALHNHYTQKSLSLSPGKGGGGSGGGGSGGG GSGGGGSGGGGSGGGGSDKTHTCPPCPPELL GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH EDPEVKFNWYVDGVEVHNAKTKPCEEQYGST YRCVSVLTVLHQDWLNGKEYKCKVSNKALPA PIEKTISKAKGQPREPQVYTLPPSREEMTKNQV SLTCLVKGFYPSDIAVEWESNGQPENNYKTTPP VLDSGDSFFLYSKLTVDKSRWQQGNVFCVSM HEALHNHYTQKSLSLSPGKGGGGGEVQLLESGG GVVQPGRSLRLSCAASGFTFSSYGMGWVRQAP GKGLEWVAVISYDGSNKYYADSVKGRFTISR NSKNTLYLQMNSLRAEDTAVYYCAREGAHFG SGSYYPYLYYYAMDVWGQGTTVTSSGGGGG GGGGSGGGGSELTLTQSPGTLSPGERATLSC RASQSVSSSYLAWYQQKPGQAPRLLIYGASIRA TGIPDRFSGSGSGTDFTLTISRLEPEDFAVYYCQ QYGSSPIFTFGPGTKVEIKSGEAAAKEAAAKEA AAAKEAAAKEAAAKEAAAKEAAAKEAAAKEA AAAKEAAAKEVQLVESGGGLVQPGGSLKLSCA ASGFTFNKYAMNWVRQAPGKCLEWVARIRSK YNNYATYYADSVKDRFTISRDDSKNTAYLQM NNLKTEDTAVYYCVRHGNFGNSYISYWAYWG QGTTLTVSSGGGGSGGGGSGGGGSQTVVTQEP SLTVSPGGTVTLTCGSSTGAVTSGNYPNWVQQ KPGQAPRGLIGGTKFLAPGTPARFSGSLLGGKA ALTLSGVQPEDEAEYYCVLWYSNRWVFGCGT KLTVL</p>
353.	<p>EpCAM_5- 10_x(G4S)3_x I2Ccc_xG4_xscFc _xG4_xMSLN_H _x(G4S)3_x I2Ccc- Full Sequence</p>	artificial	aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTTVTSSGGGGSGG GGSGGGGSEVQLVESGGGLVQPGGSLKLSCAA SGFTFNKYAMNWVRQAPGKCLEWVARIRSKY</p>

				<p>NNYATYYADSVKDRFTISRDDSKNTAYLQMN NLKTEDTAVYYCVRHGNGNSYISYWAYWGQ GTLVTVSSGGGGSGGGSGGGGSQT VVTQEPS LTVSPGGTVTLTCGSSTGAVTSGNYPNWVQQK PGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAA LTLSGVQPEDEAEYYCVLWYSNRWVFGCGTK LTVLGGGGDKTHTCPPCPAPPELLGGPSVFLFPP KPKDTLMISRTPEVTCVVVDVSHEDPEVKFNW YVDGVEVHNAKTKPCEEQYGSTYRCVSVLTV LHQDWLNGKEYKCKVSNKALPAPIEKTISKAK GQPREPQVYTLPPSREEMTKNQVSLTCLVKGF YPSDIAVEWESNGQPENNYKTTTPVLDSGGSFF LYSKLTVDKSRWQQGNVFCFSVMHEALHNHY TQKLSLSLSPGKGGGGSGGGSGGGSGGGSGG GGSGGGGSDKHTHTCPPCPAPPELLGGPSVFLFPP KPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTV VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTTPVLDSGGSF FLYSKLTVDKSRWQQGNVFCFSVMHEALHNH YTQKLSLSLSPGKGGGGEVQLLESGGGVVQPGR SLRLSCAASGFTFSSYGMGWVRQAPGKGLEW VAVISYDGSNKYYADSVKGRFTISRDN SKNTL YLQMNSLRAEDTAVYYCAREGAHFGSGSYYP LYYYYAMDVWGQGT TTVTVSSGGGGSGGGGS GGGSELTLTQSPGTLSPGERATLSCRASQS VSSSYLAWYQQKPGQAPRLLIYGASIRATGIPD RFSGSGSGTDFTLTISRLEPEDFAVYYCQQYGS SPIFTFGPGTKVEIKSGGGSGGGSGGGGSEV QLVESGGGLVQPGGSLKLSCAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNGNSYISYWAYWGQGT LVTVSSGG GGSGGGSGGGGSQT VVTQEPSLTVSPGGT VT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAA LTLSGVQPED EAEYYCVLWYSNRWVFGCGTKLTVL</p>
354.	EpCAM_5-10_x I2Ccc_xscFc_xM SLN_H2_x I2Ccc - Full Sequence	artificial	aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLNS GNQKNYL TWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRP HGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSCAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNGNSYISYWAYWGQGT LVTVSSGG GGSGGGSGGGGSQT VVTQEPSLTVSPGGT VT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAA LTLSGVQPED EAEYYCVLWYSNRWVFGCGTKLTVL DKHTHTC PPCPAPPELLGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHEDPEVKFNWYVDGVEVHNAKTKP</p>

				<p>CEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQG NVFSCSVMEALHNHYTQKSLSLSPGKGGGGS GGGSGGGGSGGGGSGGGGSGGGGSDKTHTC PPCPAPPELLGGPSVFLFPPKPKDTLMISRTPEVT CVVVDVSHEDPEVKFNWYVDGVEVHNAKTKP CEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK VSNKALPAPIEKTISKAKGQPREPQVYTLPPSRE EMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQG NVFSCSVMEALHNHYTQKSLSLSPGKEVQLL ESGGGVVQPGRSLRLSCAASGFTFSSYGMGWV RQAPGKGLEWVAVISYDGSNKYYADSVKGRF TISRDNKNTLYLQMNSLRAEDTAVYYCAREG AHFGSGSYYP LYYYYAMDVWGQGT TTVVSSG GGGSGGGGSGGGGSELTLTQSPGTLSPGERA TLSCRASQSVSSSYLAWYQQKPGQAPRLLIYG ASIRATGIPDRFSGSGSGTDFTLTISRLEPEDFAV YYCQQYGSSPIFTFGPGTKVEIKSGGGGSEVQL VESGGGLVQPGGSLKLSCAASGFTFNKYAMN WVRQAPGKCLEWVARIRSKYNNYATYYADSV KDRFTISRDDSKNTAYLQMNNLKTEDTAVYYC VRHGNFGNSYISYWAYWGQGT LTVVSSGGGG SGGGGSGGGGSQT VVTQEPSLTVSPGGTVTLT CGSSTGAVTSGNYPNWVQKPGQAPRGLIGGT KFLAPGTPARFSGSLLGGKAALTLSGVQPEDEA EYYCVLWYSNRWVFGCGTKLTVL</p>
355.	EpCAM_x(G4S)1 0_x I2Ccc_xscFc_x I2Ccc_x(G4S)10_ x_MSLN_H2 - Full Sequence	artificial	Aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLLS GNQKNYL TWYQQKPGQP KLLIYWASTRESG VPDRFTGSGSGTDFTLTISSVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGGS EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF K GKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGT TTVVSSGGGGSGG GGSGGGGSGGGGSGGGGSGGGGSGGGGSGGG GSGGGGSGGGGSEVQLVESGGGLVQPGGSLKL SCAASGFTFNKYAMNWVRQAPGKCLEWVARI RSKYNNYATYYADSVKDRFTISRDDSKNTAYL QMNNLKTEDTAVYYCVRHGNFGNSYISYWAY WGQGT LTVVSSGGGGSGGGGSGGGGSQT VVT QEPSLTVSPGGTVTLT CGSSTGAVTSGNYPNW VQKPGQAPRGLIGGT KFLAPGTPARFSGSLLG GKAALTLSGVQPEDEAEYYCVLWYSNRWVFG CGTKLTVLGGGGDKTHTCPPCPAPPELLGGPSV FLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEV KFNWYVDGVEVHNAKTKPCEEQYGSTYRCVS V LTVLHQDWLNGKEYKCKVSNKALPAPIEKTI SKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTTPVLDS DGSFFLYSKLTVDKSRWQQGNVFSCSVMEAL LHNHYTQKSLSLSPGKGGGSGGGGSGGGGS GGGSGGGGSGGGGSDKTHTCPPCPAPPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHED</p>

				PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHE ALHNHYTQKSLSLSPGKGGGGEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGTLVTVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGTVTLTCSSTGA VTSGNYPNWVQKPGQAPRGLIGGTFKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLSGGGGSGGGGSGGG GSGGGGSGGGGSGGGGSGGGGSGGGGSGGGG SEVQLLESQGGVVPGRSLRLSCAASGFTFSSY GMGWVRQAPGKGLEWVAVISYDGSNKYYAD SVKGRFTISRDNKNTLYLQMNSLRAEDTAVY YCAREGAHFGSGSYPLYYYYAMDVWGQGT TVTSSGGGGSGGGGSGGGGSELTLTQSPGTL SLSPGERATLSCRASQSVSSYLAWYQQKPGQ APRLLIYGASIRATGIPDRFSGSGSGTDFTLISR LEPEDFAVYYCQQYGSSPIFTFGPGTKVEIK
356.	CD20_99-E5_CC - HCDR1	artificial	Aa	SYWMH
357.	CD20_99-E5_CC - HCDR2	artificial	Aa	YITPSTGYTEYNQKFKG
358.	CD20_99-E5_CC - HCDR3	artificial	Aa	VHDYDRAMEY
359.	CD20_99-E5_CC - LCDR1	artificial	Aa	KASQDINKYIA
360.	CD20_99-E5_CC - LCDR2	artificial	Aa	YTSTLQP
361.	CD20_99-E5_CC - LCDR3	artificial	Aa	LQYASYPFT
362.	CD20_99-E5_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTSY WMHWVRQAPGQCLEWIGYITPSTGYTEYNQK FKGRVTMTRDKSTSTVYMELSSLTSEDYAVYY CARVHDYDRAMEYWGQGTITVTVSS
363.	CD20_99-E5_CC - VL	artificial	Aa	DIQMTQSPSSLSASVGDRTITCKASQDINKYIA WYQQKPGKPKLLIYYTSTLQPGVPSRFSGSGS GTDFTFTISLQPEDATYYCLQYASYPFTFGCG TRLEIK
364.	CD20_99-E5_CC EI - VL	artificial	aa	EIQMTQSPSSLSASVGDRTITCKASQDINKYIA WYQQKPGKPKLLIYYTSTLQPGVPSRFSGSGS GTDFTFTISLQPEDATYYCLQYASYPFTFGCG TRLEIK
365.	CD22_28- B7N655S_CC - HCDR1	artificial	aa	SYGIS
366.	CD22_28- B7N655S_CC - HCDR2	artificial	aa	WISAYSGNAIYAQKLQG
367.	CD22_28- B7N655S_CC -	artificial	aa	DPDYYGSGSYSDY

	HCDR3			
368.	CD22_28- B7N655S_CC - LCDR1	artificial	aa	RASQSVSSNLA
369.	CD22_28- B7N655S_CC - LCDR2	artificial	aa	GASSRAT
370.	CD22_28- B7N655S_CC - LCDR3	artificial	aa	QQYHSWPLLT
371.	CD22_28- B7N655S_CC - VH	artificial	aa	QVQLVQSGAEVKKPGASVKV SCKASGYTFTSY GISWVRQAPGQCLEWMGWISAYSG NAIYAQK LQGRVTMTRDTSTSTAYMELRSLR SDDTAVYY CARDPDYYGSGSYSDYWGQGLVTVSS
372.	CD22_28- B7N655S_CC - VL	artificial	Aa	EIVLTQSPATLSVSPGERATLSC RASQSVSSNLA WYQQKPGQAPRLLIYGASSRATGIP ARFSGSGS GTEFTLTISSLQSEDFAVYYCQQY HSWPLLTFG CGTKVEIK
373.	CD22_28- B7N655SCC_x_I 2C_x_(G4S)3_x_s cFc_x_(G4S)3_x_ CD20_99- E5_CC_x_I2C - Full Sequence	artificial	Aa	QVQLVQSGAEVKKPGASVKV SCKASGYTFTSY GISWVRQAPGQCLEWMGWISAYSG NAIYAQK LQGRVTMTRDTSTSTAYMELRSLR SDDTAVYY CARDPDYYGSGSYSDYWGQGLVTVSS GGGG SGGGGSGGGGSEIVLTQSPATLSV SPGERATLS CRASQSVSSNLA WYQQKPGQAPRLLIYGASSR ATGIPARFSGSGSGTEFTLTISSLQ SEDFAVYYC QQYHSWPLLTFGCGTKVEIKSGGGG SEVQLVE SGGGLVQPGGSLKLS CAASGFTFNKYAMNWV RQAPGKGLEWVARIRSKYNNYATYY ADSVKD RFTISRDDSKNTAYLQMN NLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLVTVSS GGGGSG GGGSGGGGSQTVVVTQEP SLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQ QKPGQAPRGLIGGTF LAPGTPARFSGSLLG GKAALTLSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVL GGGGSGGG GSGGGGSDKTHTCPPCP APELLGGPSVFLFPK PKDITLMISRTPEVTCV VVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQY GSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALP APIEKTISKAKG QPREPQVYTLPPSREEMTK NQVSLTCLVKGFY PSDIAVEWESNGQPENNYK TTPVLDSDGSFFL YSKLTVDKSRWQQGNV FSCVMHEALHNHYT QKSLSLSPGKGGGGSGGGG SGGGGSGGGGSG GGGSGGGGSDKTHTCPPCP APELLGGPSVFLFP PKPKDITLMISRTPEVTCV VVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVL VLHQDWLNGKEYKCKVSNKALP APIEKTISKA KGQPREPQVYTLPPSREEMTK NQVSLTCLVKG FYPSDIAVEWESNGQPENNYK TTPVLDSDGSF FLYSKLTVDKSRWQQGNV FSCVMHEALHNH YTQKSLSLSPGKGGGGSGGGG SGGGGSQVQL VQSGAEVKKPGASVKV SCKASGYTFTSYWMH WVRQAPGQCLEWIGYITPSTGYTE YNQKFKGR VTMTRDKSTSTVYME LSSLTSED TAVYYCARV HDYDRAMEYWGQGT TVTVSSGGGGSGGGG SGGGGSDIQMTQSPSSLSASV GDRVTITCKASQD INKYIAWYQQKPGKGP KLLIYYTSTLQPGVPSR

			<p>FSGSGSGTDFTFTISSLQPEDIATYYCLQYASYP FTFGCGTRLEIKSGGGGSEVQLVESGGGLVQPG GSLKLSAASGFTFNKYAMNWVRQAPGKGLE WVARIRSKYNNYATYYADSVKDRFTISRDDSK NTAYLQMNNLKTEDTAVYYCVRHGNFGNSYI SYWAYWGQGLTVTVSSGGGGSGGGGSGGGG SQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSG NYPNWVQKPGQAPRGLIGGTKFLAPGTPARF SGSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGGGTKLTVL</p>
<p>374.</p>	<p>CD22_28- B7_N655S_CC_x _I2E_x_(G4Q)3_x _scFc_x_(G4Q)3x _CD20_99- E5_CC_x_I2E_EI mod - Full Sequence</p>	<p>artificial</p>	<p>Aa</p> <p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTSY GISWVRQAPGQCLEWMGWISAYSGNAIYAQK LQGRVTMTRDTSTSTAYMELRSLRSDDTAVYY CARDPDYYGSGSYSDYWGQGLTVTVSSGGGG QGGGGQGGGGQEIQLTQSPATLSVSPGERATL SCRASQSVSSNLAWYQKPKGQAPRLLIYGASS RATGIPARFSGSGSGTEFTLTISSLQSEDFAVYY CQQYHSWPLLTFGCGTKVEIKSGGGGQEVQLV ESGGGLVQPGGSLKLSAASGFTFNKYAINWV RQAPGKGLEWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTVYLQMNNLKTEDTAVYYCAR AGNFGSSYISYWAYWGQGLTVTVSSGGGGQG GGGQGGGGQQTVVTQEPSLTVSPGGTVTITCG SSTGAVTSGNYPNWVQKPKGQAPRGLIGGTKF LAPGTPARFSGSLSGGKAALTLSGVQPEDEAEY YCVLWYSNRWVFGSGTKLTVLGGGGQGGGG QGGGGQCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEEPEVKFNWYVDGVEV HNAKTKPCEEQYGYSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTPPVLDSDGSFFLYSKLTV DKSRWQQGNVFCFSVMHEALHNHYTQKSLSL SPGKGGGGQGGGGQGGGGQGGGGQGGGGQGG GGGQCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEEPEVKFNWYVDGVEVH NAKTKPCEEQYGYSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFCFSVMHEALHNHYTQKSLSLSPG KGGGGQGGGGQGGGGQGVQLVQSGAEVKKP GASVKVSCKASGYTFTSYWMHWVRQAPGQC LEWIGYITPSTGYTEYNQKFKGRVTMTRDKST STVYMELSSLTSEDVAVYYCARVHDYDRAME YWGQGLTVTVSSGGGGQGGGGQGGGGQGGGGQ MTQSPSSLSASVGDRTITCKASQDINKYIAWY QKPKGKPKLLIYYTSTLQPGVPSRFSGSGSGT DFTFTISSLQPEDIATYYCLQYASYPFTFGCGTR LEIKSGGGGQEVQLVESGGGLVQPGGSLKLSA AASGFTFNKYAINWVRQAPGKGLEWVARIRSK YNNYATYYADAVKDRFTISRDDSKNTVYLQM NNLKTEDTAVYYCARAGNFGSSYISYWAYWG QGLTVTVSSGGGGQGGGGQGGGGQQTVVTQE PSLTVSPGGTVTITCGSSTGAVTSGNYPNWVQ KKPGQAPRGLIGGTKFLAPGTPARFSGSLSGGK</p>

				AALTLSGVQPEDEAEYYCVLWYSNRWVFGSG TKLTVL
375.	CD22_28- B7_N655S_CC_x _I2E_x_G4_x_sc Fc_x_G4_x_CD2 0_99- E5_CC_x_I2E_G Q_EImod - Full Sequence	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTSY GISWVRQAPGQCLEWMGWISAYSIGNAIYAQK LQGRVTMTRDTSTSTAYMELRSLRSDDTAVYY CARDPDYYSYSGSYSDYWGQGLVTVSSGGGG QGGGGQGGGGQEIIVLTQSPATLSVSPGERATL SCRASQSVSSNLAWYQQKPGQAPRLLIYGASS RATGIPARFSGSGSGTEFTLTISSLQSEDAVYY CQQYHSWPLLTFGCGTKVEIKSGGGGQEVQLV ESGGGLVQPGGSLKLSAASGFTFNKYAINWV RQAPGKGLEWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTVYLQMNNLKTEDTAVYYCAR AGNFGSSYISYWAYWGQGLVTVSSGGGGQG GGGQGGGGQQTVVVTQEPSLTVSPGGTVTITCG SSTGAVTSGNYPNWVQKKPGQAPRGLIGGTFK LAPGTPARFSGSLSGGKAALTLSGVQPEDEAEY YCVLWYSNRWVFGSGTKLTVLGGGGCPCPA PELLGGPSVFLFPPKPKDTLMISRTPEVTCVVV DVSHEEPEVKFNWYVDGVEVHNAKTKPCEEQ YGSTYRCVSVLTVLHQDWLNGKEYKCKVSNK ALPAPIEKTISKAKGQPREPQVYTLPPSREEMT KNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTPPVLDSDGSFFLYSKLTVDKSRWQQGNVF SCSVMHEALHNHYTQKLSLSLSPGKGGGGQGG GGQGGGGQGGGGQGGGGQGGGGQCPPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKLSLSLSPGKGGGGQVQL VQSGAEVKKPGASVKVSCKASGYTFTSYWMH WVRQAPGQCLEWIGYITPSTGYTEYNQKFKGR VTMTRDKSTSTVYMESSLTSEDTAVYYCARV HDYDRAMEYWGQGTITVTVSSGGGGQGGGGQ GGGGQEIQMTQSPSSLSASVGDRVTITCKASQD INKYIAWYQQKPGKPKLLIYTTSTLQPGVPSR FSGSGSDFTFTISSLQPEDIAATYYCLQYASYP FTFGCTRLEIKSGGGGQEVQLVESGGGLVQP GGSLKLSAASGFTFNKYAINWVRQAPGKGLE WVARIRSKYNNYATYYADAVKDRFTISRDDSK NTVYLQMNNLKTEDTAVYYCARAGNFGSSYIS YWAYWGQGLVTVSSGGGGQGGGGQGGGGQ QTVVTQEPSLTVSPGGTVTITCGSSTGAVTSGN YPNWVQKKPGQAPRGLIGGTFKFLAPGTPARFS GSLSGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGSGTKLTVL
376.	CL1 9-G4 CC - HCDR1	artificial	Aa	DYYMH
377.	CL1 9-G4 CC - HCDR2	artificial	Aa	WINPNSGGPNYAQKFQG
378.	CL1 9-G4 CC - HCDR3	artificial	Aa	EKHAVAGIGFDY
379.	CL1 9-G4 CC -	artificial	Aa	QASQDISNYLN

	LCDR1			
380.	CL1 9-G4 CC - LCDR2	artificial	Aa	AASSLES
381.	CL1 9-G4 CC - LCDR3	artificial	aa	QQANSFPLT
382.	CL1 9-G4 CC - VH	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWRQAPGQCLEWMGWNPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGLTIVTSS
383.	CL1 9-G4 CC - VL	artificial	aa	DIQMTQSPSSVSASVGDRVTITCQASQDISNYL NWXQQKPGKAPKLLIYAASSLESGVPSRFSGS GSGTDFLTITSSLPEDFATYYCQQANSFPLTFG CGTKVDIK
384.	CL1 9-G4 CC EI - VL	artificial	aa	EIQMTQSPSSVSASVGDRVTITCQASQDISNYL NWXQQKPGKAPKLLIYAASSLESGVPSRFSGS GSGTDFLTITSSLPEDFATYYCQQANSFPLTFG CGTKVDIK
385.	FL 4-E9 CC - HCDR1	artificial	aa	NARMGVS
386.	FL 4-E9 CC - HCDR2	artificial	aa	HIFSNDEKSYSTSLKS
387.	FL 4-E9 CC - HCDR3	artificial	aa	VPEYSSGWYRFDY
388.	FL 4-E9 CC - LCDR1	artificial	aa	RASQSIRSYLN
389.	FL 4-E9 CC - LCDR2	artificial	Aa	ATSSLQG
390.	FL 4-E9 CC - LCDR3	artificial	Aa	QQSYSTPFT
391.	FL 4-E9 CC - VH	artificial	Aa	QVTLKESGPTLVKPTETLTLTCTVSGFSFRNAR MGVSWIRQPPGKCLEWLAHIFSNDEKSYSTSL KSRLTISKDTSKSQVVLMTNMDPVDATATYFC ARVPEYSSGWYRFDYWGQGLTIVTSS
392.	FL 4-E9 CC - VL	artificial	Aa	DIQMTQSPSSLSASVGDRVTISCRASQSIRSYLN WYQQKPGKAPKLLIYATSSLQGGVPSRFSGSG SGTDFLTITSSLPEDFATYYCQQSYSTPFTFGC GTKVEIK
393.	FL 4-E9 CC EI - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVTISCRASQSIRSYLN WYQQKPGKAPKLLIYATSSLQGGVPSRFSGSG SGTDFLTITSSLPEDFATYYCQQSYSTPFTFGC GTKVEIK
394.	CL1 9-G4 CC x4F10.03 scFc xFL 4-E9 CC x4F10.03 mut - Full Sequence	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWRQAPGQCLEWMGWNPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGLTIVTSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWXQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFLTITSSLPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWV RQAPGKGMWVARIRSKYNNYATYYADAVK DRFTISRDDSKNTLYLQMNKLTEDTAVYYCV RAGNFGKSYISYWAYWGQGLTIVTSSGGGG GGGGSGGGGQTIVTQEPSLTVSPGGTITC GSSTGAVTSGNYPNWVQKKPGQAPRGLIGGTK

			<p>FLAPGTPARFSGSLSGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGSGTKLTVLGGGGSSGGG GSGGGGSDKTHTCPPCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSGDSFFL YSKLTVDKSRWQQGNVFCSCVMHEALHNYT QKSLSLSPGKGGGGSSGGGGSSGGGGSSGGGGSSG GGGSSGGGSDKTHTCPPCPAPPELLGGPSVFLFPP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPVLDSGDSF FLYSKLTVDKSRWQQGNVFCSCVMHEALHNH YTQKSLSLSPGKGGGGSSGGGGSSGGGGSSQVTLK ESGPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKSQVVLMTNMDPVDATYFCARVPE YSSGWYRFDYWGQGLTVTVSSGGGGSSGGGGSS GGGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGGTDFTLTISSLQPEDFATYYCQQSYSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADA VKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGK SYISYWAYWGQGLTVTVSSGGGGSSGGGGSSG GGSQTVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWVQKKPGQAPRGLIGGTFKFLAPGTPA RFSGSLSGGKAALTLSGVQPEDEAEYYCVLWY SNRWVFGSGTKLTVL</p>
395.	<p>CL1 9-G4 CC x4G10.04x scFc xFL 4-E9 CC x4G10.04 - Full Sequence</p>	artificial	<p>Aa QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSSGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSCAASGFTFSKYAMNWV REAPGKGLEWVARIRSKYNNYATYYAEAVKD RFTISRDDSKNTVYLQMNNLKTEDTAVYYCVR AENIGKSYISYWAYWGQGLTVTVSSGGGGSSG GGGSSGGGSSQTVVTQEPSLTVSPGGTVTMTCG SSTGAVTSGNYPNWVQKKPGQAPRGLIGGTFK LAPGTPARFSGSLEGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGSSGGG GSGGGGSDKTHTCPPCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSGDSFFL</p>

				<p>YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPVLDSGDSF FLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSGVTLK ESGPTLVKPTETLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKTSQVVLMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGLVTVSSGGGGSGGGGS GGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGSGTDFTLTISSLQPEDFATYYCQQSYSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFSKYAMNWWREAPGKGL EWVARIRSKYNNYATYYAEAVKDRFTISRDDS KNTVYLQMNNLKTEDTAVYYCVRAENIGKSYI SYWAYWGQGLVTVSSGGGGSGGGGSGGGG SQT VVTQEPSLTVSPGGTVTMTCGSSTGAVTS GNYPNWVQKPGQAPRGLIGGTKFLAPGTPAR FSGSLEGGKAALTLGSGVQPEDEAEYYCVLWYS NRWVFGGGTKLTVL</p>
396.	CL1 9-G4 CC x5B1.05 x scFc xFL 4-E9 CC x5B1.05 - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSCAASGFTFSKYAMNWW RQAPGKGMWVARIRSKYNNYATYYAEAVK GRFTISRDDSKNTVYLQMNNLKTEDTAVYYCV RAGNFGSSYISYWAYWGQGLVTVSSGGGGS GGGSGGGGSQT VVTQEPSLTVSPGGTVTLTC GSSTGAVTSGNYPNWVQKPGQAPRGLIGGTK FLAPGTPARFSGSLGGKAALTLGSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGSGGG GSGGGGSDKTHTCPPCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSGDSFFL YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPVLDSGDSF</p>

				<p>FLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSQVTLK ESGPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKSQVVLMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGLTVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGSGTDFTLTISSLQPEDFATYYCQQSYSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFSKYAMNWRQAPGKGM EWVARIRSKYNNYATYYAEAVKGRFTISRDDS KNTVYLQMNNLKTEDTAVYYCVRAGNFGSSY ISYWAYWGQGLTVTVSSGGGGSGGGGSGGGG SQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSG NYPNWVQKKPGQAPRGLIGGTKFLAPGTPARF SGLSGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGGGTKLTVL</p>
<p>397.</p>	<p>CL1 9-G4 CC x5B1.09 x scFc xFL 4-E9 CC xH5B1.09 - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSCAASGFTFSKYAMNWR RQAPGKGMWVARIRSKYNNYATYYADAVK GRFTISRDDSKNTVYLQMNNLKTEDTAVYYCV RAGNFGKSYISYFAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVTQEPSLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQKKPGQAPRGLIGGTKF LAPGTPARFSGLSGGKAALTLSGVQPEDEAEY YCVLYYSNRWVFGGGTKLTVLGGGGSGGGGS GGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPK DTLMISRTPPEVTCVVVDVSHEDPEVKFNWYVD GVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQ DWLNGKEYKCKVSNKALPAPIEKTISKAKGQP REPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSDGSFFLYS KLTVDKSRWQQGNVFSCSVMHEALHNHYTQK SLSLSPGKGGGGSGGGGSGGGGSGGGGSGGGG GSGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK PKDTLMISRTPPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPPVLDSDGSFFL YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSGGGG GPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTISK DTSKSQVVLMTNMDPVDTATYFCARVPEYSS GWYRFDYWGQGLTVTVSSGGGGSGGGGSGGGG GGSDIQMTQSPSSLSASVGDRVTISCRASQSIRS YLNWYQQKPGKAPKLLIYATSSLQGGVPSRFS</p>

				<p>GSGSGTDFTLTISLQPEDFATYYCQQSYSTPFT FGCGTKVEIKSGGGGSEVQLVESGGGLVQPGG SLKLSAASGFTFSKYAMNWRQAPGKGM EW VARIRSKYNNYATYYADAVKGRFTISRDDSKN TVYLQMN NLKTEDTAVYYCVRAGNFGKSYIS YFAYWGQGLVTVSSGGGGSGGGGSGGGGSQ TVVTQEPLTVSPGGTVTLTCSSTGAVTSGNY PNWVQKPGQAPRGLIGGTKFLAPGTPARFSG SLSGGKAALTLSGVQPEDEAEYYCVLYYSNR WVFGGGTKLTVL</p>
<p>398.</p>	<p>CL1 9-G4 CC x6H10.03x scFc xFL 4-E9 CC x 6H10.03 - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLVQSGAEVKKPGASVKVSKKASGYTFTD YMHWRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISLQPEDFATYY CQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWR RQAPGKGM EWVARIRSKYNNYATYYAEAVK DRFTISRDDSKNTLYLQMN NLKTEDTAVYYCV RAGNFGKSYISYWAYWGQGLVTVSSGGGGS GGGGSGGGGSQTVVTQEPLTVSPGGTVTITC GSSTGAVTSGNYPNWIQKPGQAPRGLIGGTK FLAPGTPARFSGSLEGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGSGTKLTVLGGGGSGGG GSGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTPPVLDSDGSFLL YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPELLGGPSVFLF PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTPPVLDSDGSF FLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSGVTLK ESGPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKSQVVLMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGLVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGTDFTLTISLQPEDFATYYCQQSYSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSAASGFTFNKYAMNWRQAPGKGM EWVARIRSKYNNYATYYAEAVKDRFTISR DSKNTLYLQMN NLKTEDTAVYYCVRAGNFGK SYISYWAYWGQGLVTVSSGGGGSGGGGSGG GGSQTVVTQEPLTVSPGGTVTITCSSTGAVT SGNYPNWIQKPGQAPRGLIGGTKFLAPGTPA</p>

				RFIGSLEGGKAALTLSGVQPEDEAEYYCVLWY SNRWVFGSGTKLTVL
399.	CL1 9-G4 CC x6H10.09 x scFc xFL 4-E9 CC x6H10.09 - Full Sequence	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFLTITSSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGSLKLSAASGFTFNKYAMNWV RQAPGKGMWVARIRSKYNNYATYYADAVK DRFTISRDDSKNTLYLQMNNLKTEDTAVYYCV RAGNFGSSYISYFAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVTQEPSTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGTKFL APGTPARFSGSLEGGKAALTLSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGSGGGGS GGGGSDKTHTCPPCPAPELLGGPSVFLFPPKPK DTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD GVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQ DWLNGKEYKCKVSNKALPAPIEKTISKAKGQP REPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSDGSFFLYS KLTVDKSRWQQGNVFSCSVMHEALHNHYTQK SLSLSPGKGGGGSGGGGSGGGGSGGGGSGGG GSGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPPVLDSDGSFFL YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKLSLSLSPGKGGGGSGGGGSGGGGSGVTLKES GPTLVKPTETLTLCTVSGFSFRNARMGVSWIR QPPGKCLEWLAHIFSNDEKSYSTSLKSRLTISK DTSKSQVVLMTNMDPVDATYFCARVPEYSS GWYRFDYWGQGLTVTVSSGGGGSGGGGSGG GGSDIQMTQSPSSLSASVGDRVTISCRASQSIRS YLNWYQQKPGKAPKLLIYATSSSLQGGVPSRFS GSGSGTDFLTITSSLQPEDFATYYCQSYSTPFT FGCGTKVEIKSGGGGSEVQLVESGGGLVQPGG SLKLSAASGFTFNKYAMNWVRQAPGKGMW VARIRSKYNNYATYYADAVKDRFTISRDDSK NTLYLQMNNLKTEDTAVYYCVRAGNFGSSYIS YFAYWGQGLTVTVSSGGGGSGGGGSGGGGSQ TVVTQEPSTVSPGGTVTITCGSSTGAVTSGNY PNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGS LEGGKAALTLSGVQPEDEAEYYCVLYYSNRW VFGSGTKLTVL
400.	CL1 9-G4 CC x I2C x scFc x I2C xFL 4-E9 CC - Full Sequence	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS

				<p>LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWW RQAPGKGLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVTQEPLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKF LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGSGGG GSGGGGSDKTHTCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPPVLDSDGSFLL YSKLTVDKSRWQQGNVSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPAPPELLGGPSVFLFPP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPPVLDSDGSF FLYSKLTVDKSRWQQGNVSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSEVQLV ESGGGLVQPGGSLKLSAASGFTFNKYAMNWW VRQAPGKGLEWVARIRSKYNNYATYYADSVK DRFTISRDDSKNTAYLQMNNLKTEDTAVYYCV RHGNFGNSYISYWAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVTQEPLTVSPGGTVTLTCG GSSTGAVTSGNYPNWVQQKPGQAPRGLIGGTK FLAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGSQVTL KESGPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKSVVLTMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGLTVTVSSGGGGSGGGGS GGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGSGTDFTLTISSLQPEDFATYYCQSYSTP FTFGCGTKVEIK</p>
401.	CL1 9-G4 CC x I2C x(G4S)3xscFc x(G4S)3 xFL 4-E9 CC x I2C - Full Sequence	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWW RQAPGKGLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVTQEPLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKF</p>

			<p>LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGSGGG GSGGGGSDKTHTCPPCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSDGSFFL YSKLTVDKSRWQQGNVFCSCVMHEALHNYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLF PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPVLDSDGSF FLYSKLTVDKSRWQQGNVFCSCVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSGVTLK ESGPTLVKPTETLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKSQVVLMTNMDPVDATYFCARVPE YSSGWYRFDYWGQGLTVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGGTDFTLTISSLQPEDFATYYCQQSSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWVRQAPGKGL EWVARIRSKYNNYATYYADSVKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRHGNFGNSY ISYWAYWGQGLTVTVSSGGGGSGGGGSGGGG SQT VVTQEPLTVSPGGTVTLTCGSSTGAVTSG NYPNWVQQKPGQAPRGLIGGTKFLAPGTPARF SGSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGGGTKLTVL</p>
402.	CL1 9-G4 CC x I2Ccc x scFc xFL 4-E9 CC x I2Ccc - Full Sequence	artificial	<p>aa QVQLVQSGAEVKKPGASVKVSKKASGYTFD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSCAASGFTFNKYAMNWV RQAPGKCLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLTVTVSSGGGGSG GGGSGGGGSQT VVTQEPLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKF LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGSGGG GSGGGGSDKTHTCPPCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSDGSFFL</p>

				<p>YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPVLDSGDSF FLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGSGGGGSGVTLK ESGPTLVKPTETLTLCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKTSQVVLMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGLVTVSSGGGGSGGGGS GGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGSGTDFTLTISSLQPEDFATYYCQQSYSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWRQAPGKCL EWVARIRSKYNNYATYYADS VKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRHGNFGNSY ISYWAYWGQGLVTVSSGGGGSGGGGSGGGG SQT VVTQEPSLTVSPGGTVTLTCGSSTGAVTSG NYPNWVQQKPGQAPRGLIGGTKFLAPGTPARF SGSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWFVFCGTKLTVL</p>
403.	CL1 9-G4 CC x I2Ccc x scFc x I2Ccc xFL 4-E9 CC - Full Sequence	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRLSDDTAV YYCAREKHAVAGIGFDYWGQGLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSCAASGFTFNKYAMNWR RQAPGKCLEWVARIRSKYNNYATYYADS VKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLVTVSSGGGGSG GGGSGGGGSQT VVTQEPSLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKF LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWFVFCGTKLTVLGGGGSGGG GSGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTPVLDSGDSFFL YSKLTVDKSRWQQGNVFSCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTPVLDSGDSF</p>

				<p>FLYSKLTVDKSRWQQGNVFSVCSVMHEALHNH YTQKSLSLSPGKGGGGSGGGGGSGGGGSEVQLV ESGGGLVQPGGSLKLSAASGFTFNKYAMNW VRQAPGKCLEWVARIRSKYNNYATYYADSVK DRFTISRDDSKNTAYLQMNNLKTEDTAVYYCV RHGNFGNSYISYWAYWGQGLTVTVSSGGGGG GGGGSGGGGSQTVVTQEPLTVSPGGTVTLTC GSSTGAVTSGNYPNWVQQKPGQAPRGLIGGTK FLAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGGSQVTL KESGPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSKQVVLMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGLTVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGTDFTLTISSLQPEDFATYYCQQSYSTP FTFGCGTKVEIK</p>
404.	CL1 9-G4 CC x FL 4-E9 CC xscFc x I2Ccc x I2Ccc - Full Sequence	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSQVTLKE SGPTLVKPTETLTLTCTVSGFSFRNARMGVS RQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTISK DTSKQVVLMTNMDPVDTATYFCARVPEYSS GWYRFDYWGQGLTVTVSSGGGGSGGGGSGG GGSDIQMTQSPSSLSASVGDRVTISCRASQSIRS YLNWYQQKPGKAPKLLIYATSSLQGGVPSRFS GSGSGTDFTLTISSLQPEDFATYYCQQSYSTPFT FGCGTKVEIKSGGGGDKTHTCPPCPAPPELLGGP SVFLFPPPKDTLMISRTPEVTCVVDVSHEDP EVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC VSVLTVLHQDWLNGKEYKCKVSNKALPAPIEK TISKAKGQPREPQVYTLPPSREEMTKNQVSLT LVKGFYPSDIAVEWESNGQPENNYKTTTPVLD SDGSFFLYSKLTVDKSRWQQGNVFSVCSVMHEA LHNHYTQKSLSLSPGKGGGGSGGGGSGGGGS GGGGSGGGGSGGGGSDKTHTCPPCPAPPELLGG PSVFLFPPPKDTLMISRTPEVTCVVDVSHED PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTTPVLD DSDGSFFLYSKLTVDKSRWQQGNVFSVCSVMHE ALHNHYTQKSLSLSPGKGGGGGEVQLVESGGGL VQPGGSLKLSAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGLTVTVSSGGGGSGGGGSG GGGSQTVVTQEPLTVSPGGTVTLTCGSSTGA VTSGNYPNWVQQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL</p>

				<p>WYSNRWVFGCGTKLTVLGGGGSGGGGSGGG GSEVQLVESGGGLVQPGGSLKLSCAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGLVTV SSGGGSGGGGSGGGGSGTQVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQKPGQAP RGLIGGTKFLAPGTPARFSGSLLGGKAALTLG VQPEDEAEYYCVLWYSNRWVFGCGTKLTVL</p>
<p>405.</p>	<p>CL1 9-G4 CC x FL 4-E9 CCx I2Ccc x scFc x I2Ccc - Full Sequence</p>	<p>artificial</p>	<p>aa</p>	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWVRQAPGQCLEWMGWNPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPKGAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSQVTLKE SGPTLVKPTETLTLTCTVSGFSFRNARMGVSWI RQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTISK DTSKSQVVL TMTNMDPVDATYFCARVPEYSS GWYRFDYWGQGLVTVSSGGGSGGGGSGG GGSDIQMTQSPSSLSASVGDRVTISCRASQSIRS YLNWYQQKPKGAPKLLIYATSSLQGGVPSRFS GSGSGTDFTLTISLQPEDFATYYCQSYSTPFT FGCGTKVEIKSGGGGSEVQLVESGGGLVQPGG SLKLSCAASGFTFNKYAMNWVRQAPGKCLEW VARIRSKYNNYATYYADSVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRHGNFGNSYIS YWAYWGQGLVTVSSGGGSGGGGSGGGGSG QTVVTQEPSLTVSPGGTTLTCGSSTGAVTSGN YPNWVQKPGQAPRGLIGGTKFLAPGTPARFS GSLGKAALTLGSGVQPEDEAEYYCVLWYSN RWVFGCGTKLTVLGGGGDKHTCPCPAPPELL GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH EDPEVKFNWYVDGVEVHNAKTKPCEEQYGST YRCVSVLTVLHQDWLNGKEYKCKVSNKALPA PIEKTISKAKGQPREPQVYTLPPSREEMTKNQV SLTCLVKGFYPSDIAVEWESNGQPENNYKTTPP VLDSGDGSFFLYSKLTVDKSRWQQGNVFCSSVM HEALHNHYTQKSLSLSPGKGGGGSGGGGSGG GGSGGGGSGGGGSGGGGSDKHTCPCPAPPELL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTTP PPVLDSGDGSFFLYSKLTVDKSRWQQGNVFCSS VMHEALHNHYTQKSLSLSPGKGGGGGEVQLVE SGGGLVQPGGSLKLSCAASGFTFNKYAMNWV RQAPGKCLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLVTVSSGGGSGG GGGSGGGGSGTQVVTQEPSLTVSPGGTTLTCG SSTGAVTSGNYPNWVQKPGQAPRGLIGGTKF LAPGTPARFSGSLLGGKAALTLGSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVL</p>

406.	CL1 9-G4 CC x FL 4-E9 CCx I2Ccc x I2Ccc xscFc - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGTLLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSQVTLKE SGPTLVKPTETLTLCTVSGFSFRNARMGVSWI RQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTISK DTSKSQVVL TMTNMDPVDATYFCARVPEYSS GWYRFDYWGQGTLLVTVSSGGGGSGGGGSGG GGSDIQMTQSPSSLSASVGDRVTISCRASQSIRS YLNWYQQKPGKAPKLLIYATSSSLQGGVPSRFS GSGSGTDFTLTISSLQPEDFATYYCQQSYSTPFT FGCGTKVEIKSGGGGSEVQLVESGGGLVQPGG SLKLSAASGFTFNKYAMNWRQAPGKCLEW VARIRSKYNNYATYYADSVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRHGNFGNSYIS YWAYWGQGTLLVTVSSGGGGSGGGGSGGGGS QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWWVQKPGQAPRGLIGGTKFLAPGTPARFS GSLGGAALTLSGVQPEDEAEYYCVLWYSN RWVFGCGTKLTVLGGGGSGGGGSGGGGSEVQ LVESGGGLVQPGGSLKLSAASGFTFNKYAMN WVRQAPGKCLEWVARIRSKYNNYATYYADSV KDRFTISRDDSKNTAYLQMNNLKTEDTAVYYC VRHGNFGNSYISYWAYWGQGTLLVTVSSGGGG SGGGGSGGGGSQTVVTQEPSLTVSPGGTVTLT CGSSTGAVTSGNYPNWWVQKPGQAPRGLIGGT KFLAPGTPARFSGSLGGAALTLSGVQPEDEA EYYCVLWYSNRWVFGCGTKLTVLGGGGDKT HTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTP EVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K</p>
407.	CL1 9-G4 CC x I2Ccc xG4 xscFc xG4 xFL 4-E9 CC x I2Ccc - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGTLLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE</p>

				<p>SGGGLVQPGGSLKLSCAASGFTFNKYAMNWW RQAPGKCLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVTQEPSLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQKPGQAPRGLIGGTFK LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGDKTH TCPPCPAPELLGGPSVFLFPPKPKDTLMISRTP VTCVVVDVSHEDPEVKFNWYVDGVEVHNAK TKPCEEQYGSTYRCVSVLTVLHQDWLNGKEY KCKVSNKALPAPIEKTISKAKGQPREPQVYTL PSREEMTKNQVSLTCLVKGFYPSDIAVEWESN GQPENNYKTTPPVLDSDGSFFLYSKLTVDKSR WQQGNVFSCSVMHEALHNHYTQKSLSLSPGK GGGSGGGGSGGGGSGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGQVTLKESGPTLVKPTETLTLCTVSGFS FRNARMGVSWIRQPPGKCLEWLAHIFSNDEKS YSTSLKSRLTISKDTSKSQVLTMTNMDPVDT ATYFCARVPEYSSGWYRFDYWGQGLTVTVSS GGGSGGGGSGGGGSDIQMTQSPSSLSASVGD RVTISCRASQSIRSYLNWYQKPGKAPKLLIYA TSSLQGGVPSRFSGSGSGTDFTLTISSLQPEDFA TYQCQSYSTPFTFGCGTKVEIKSGGGGSEVQL VESGGGLVQPGGSLKLSCAASGFTFNKYAMN WVRQAPGKCLEWVARIRSKYNNYATYYADSV KDRFTISRDDSKNTAYLQMNNLKTEDTAVYYC VRHGNFGNSYISYWAYWGQGLTVTVSSGGGG SGGGGSGGGGSQTVVTQEPSLTVSPGGTVTLT CGSSTGAVTSGNYPNWVQKPGQAPRGLIGGT KFLAPGTPARFSGSLLGGKAALTLSGVQPEDEA EYYCVLWYSNRWVFGCGTKLTVL</p>
408.	CL1 9-G4 CC xscFc x FL 4-E9 CC x I2Ccc x I2Ccc - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSKKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGD RVTI TCQASQDISNYLNWYQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKGGGGDKTHTCPP CPAPELLGGPSVFLFPPKPKDTLMISRTP EVTCTVVVDVSHEDPEVKFNWYVDGVEVHNAK TKPCEEQYGSTYRCVSVLTVLHQDWLNGKEY KCKVSNKALPAPIEKTISKAKGQPREPQVYTL PSPREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGSG GGGSGGGGSGGGGSGGGGSGGGGSDKTHTC PCPAPELLGGPSVFLFPPKPKDTLMISRTP EVTCT</p>

				VVVDVSHEDPEVKFNWYVDGVEVHNAKTKPC EEQYGSTYRCVSVLTVLHQDWLNGKEYKCKV SNKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSDGSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGKGGGGQ VTLKESGPTLVKPTETLTCTVSGFSFRNARM GVSWIRQPPGKCLEWLAHIFSNDEKSYSTSLKS RLTISKDTSKSQVVLMTNMDPVDTATYFCAR VPEYSSGWYRFDYWGQGTLLTVSSGGGGSSGG GGSGGGGSDIQMTQSPSSLSASVGDRVITISCR SQSIRSYLNWYQQKPGKAPKLLIYATSSLQGG VPSRFSGSGSGTDFTLTISSLQPEDFATYYCQQS YSTPFTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSCAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGTLLTVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGTVTLTCGSSTGA VTSGNYPNWVQKPGQAPRGLIGGTKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVLGGGGSGGGGSGGG GSEVQLVESGGGLVQPGGSLKLSCAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGTLLTV SSGGGGSGGGGSGGGGSQTVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQKPGQAP RGLIGGTKFLAPGTPARFSGSLLGGKAALTLSG VQPEDEAEYYCVLWYSNRWVFGCGTKLTVL
409.	CL1_9- G4_CC_x(EAAA K)10_x I2Ccc_xscFc_xFL _4- E9_CC_x(EAAA K)10_x I2Ccc - Full Sequence	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRSDDTAV YYCAREKHAVAGIGFDYWGQGTLLTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVIT TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGEAAAKEAAAK EAAAKEAAAKEAAAKEAAAKEAAAKEAAAK EAAAKEAAAKEVQLVESGGGLVQPGGSLKLS CAASGFTFNKYAMNWVRQAPGKCLEWVARIR SKYNNYATYYADSVKDRFTISRDDSKNTAYLQ MNNLKTEDTAVYYCVRHGNFGNSYISYWAY WGQGTLLTVSSGGGGSGGGGSGGGGSQTVVT QEPSLTVSPGGTVTLTCGSSTGAVTSGNYPNW VQKPGQAPRGLIGGTKFLAPGTPARFSGSLLG GKAALTLSGVQPEDEAEYYCVLWYSNRWVFG CGTKLTVLGGGGDKTHTCPPCPAPELLGGPSV FLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEV KFNWYVDGVEVHNAKTKPCEEQYGSTYRCVS VLTVLHQDWLNGKEYKCKVSNKALPAPIEKTI SKAKGQPREPQVYTLPPSREEMTKNQVSLTCL VKGFYPSDIAVEWESNGQPENNYKTTTPVLDSD DGSFFLYSKLTVDKSRWQQGNVFSCSVMHEA LHNHYTQKSLSLSPGKGGGGSGGGGSGGGGS GGGGSGGGGSGGGGSDKHTCPPCPAPELLGG

				<p>PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVVFSCVMHEALHNHYTQKSLSLSPGKGGGGQVTLKESGPTLVKPTETLTCTVSGFSFRNARMGVSWIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTISKDTSKSQVVLTMTNMDPVDTATYFCARVPEYSSGWYRFDYWGQGTLLTVSSGGGGSGGGGSGGGGSDIQMTQSPSSLSASVGDRTVISCRAQSIRSYLNWYQKPKGKAPKLLIYATSSLQGGVPSRFSGSGGTDFTLTISSLQPEDFATYYCQQSYSTPFTFGCGTKVEIKSGEAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEAAAKEVQLVESGGGLVQPGGSLKLSCAASGFTFNKYAMNWVRQAPGKCLEWVARIRSKYNNYATYYADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAVYYCVRHGNFGNSYISYWAYWGQGTLLTVSSGGGGSGGGGSGGGGSQTVVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIGTKFLAPGTPARFSGSLLGGKAALTLVSGVQPEDEAEYYCVLWYSNRWVFGCGTKLTVL</p>
410.	<p>CL1_9-G4_CC_x(G4S)3_x I2Ccc_xscFc_xFL_4- E9_CC_x(G4S)3_x I2Ccc - Full Sequence</p>	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFDYYMHWVRQAPGQCLEWMGWNPNSGGPNYAQKFQGRVTMTRDTSISTAHMELSLRSDDTAVYYCAREKHAVAGIGFDYWGQGTLLTVSSGGGGSGGGGSGGGGSDIQMTQSPSSVSASVGDRTVITCQASQDISNYLNWYQKPKGKAPKLLIYAASSLESGVPSRFSGSGSGTDFTLTISSLQPEDFATYYCQQANSFPLTFGCGTKVDIKSGGGGSGGGGSGGGSEVQLVESGGGLVQPGGSLKLSCAASGFTFNKYAMNWVRQAPGKCLEWVARIRSKYNNYATYYADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAVYYCVRHGNFGNSYISYWAYWGQGTLLTVSSGGGGSGGGGSGGGGSQTVVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIGTKFLAPGTPARFSGSLLGGKAALTLVSGVQPEDEAEYYCVLWYSNRWVFGCGTKLTVLGGGGDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVVFSCVMHEALHNHYTQKSLSLSPGKGGGGQVTLKESGPTLVKPTETLT</p>

				<p>LTCTVSGFSFRNARMGVSWIRQPPGKCLEWLA HIFSNDEKSYSTSLKSRLTISKDTSKSQVVLMT NMDPVDATATYFCARVPEYSSGWYRFDYWGQ GTLVTVSSGGGGSGGGSGGGGSDIQMTQSPS SLSASVGDRVTISCRASQSIRS YLNWYQQKPGK APKLLIYATSSLQGGVPSRFSGSGSGTDFTLTIS SLQPEDFATYYCQQSYSTPFTFGCGTKVEIKSG GGGSGGGGSGGGSEVQLVESGGGLVQPGGS LKLSCAASGFTFNKYAMNWVRQAPGKCLEWV ARIRSKYNNYATYYADSVKDRFTISRDDSKNT AYLQMNNLKTEDTAVYYCVRHGNFGNSYISY WAYWGQGTLLTVSSGGGGSGGGGSGGGGSQ TVVTQEPLTVSPGGTVTLTCGSSTGAVTSGNY PNWVQQKPGQAPRGLIGGTKFLAPGTPARFSG SLLGGKAALTLSGVQPEDEAEYYCVLWYSNR WVFGCGTKLTVL</p>
411.	CH3-G8A_6- B12x I2Cx scFc_(G4S)6 - Full Sequence	artificial	Aa	<p>EVQLLESGLLVQPGGSLRLSCAASGFSFSSYPI NWVRQAPGKGLEWVGVIVTGGGTNYASSVK GRFTISRDNKNTVYLQMNSLRAEDTAVYYCA KSRGVYDFDGRGAMDYWGQGTLLTVSSGGG GSGGGGSGGGSDIVMTQSPDSLAVSLGERATI NCKSSQSLLYSSNQKNYFAWYQQKPGQPPKLL IYWASTRESGVPDRFSGSGSGTDFLTISLQAE DVAVYYCQQYYSYPYTFGQGTKLEIKSGGGGS EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGLEWVARIRSKYNNYATYY ADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAV VYYCVRHGNFGNSYISYWAYWGQGTLLTVSS GGGGSGGGGSGGGGSGTQVVTQEPLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQQKPGQAPRG LIGGTKFLAPGTPARFSGSLLGGKAALTLSGVQ PEDEAEYYCVLWYSNRWVFGGKTKLTVLGGG GDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHGDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSDGSFFLYSKLTV DKSRWQQGNVFCFSVMHEALHNHYTQKSLSL SPGKGGGGSGGGGSGGGGSGGGGSGGGGSGG GGSDKHTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHGD WLNKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSDGSFFLYSK LTVDKSRWQQGNVFCFSVMHEALHNHYTQKS LSLSPGK</p>
412.	CL1 9-G4 CC x I2C x scFc - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGTLLTVSSGGG GSGGGGSGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFLTISLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE</p>

				<p>SGGGLVQPGGSLKLSCAASGFTFNKYAMNWW RQAPGKGLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGTLLTVSSGGGGSG GGGSGGGGSQTVVTQEPSTLVSPGGTVTLTCG SSTGAVTSGNYPNWVQKPGQAPRGLIGGTFK LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGGGTKLTVLGGGGDKTH TCPPCPAPELLGGPSVFLFPPKPKDTLMISRTP VTCVVVDVSHEDPEVKFNWYVDGVEVHNAK TKPCEEQYGSTYRCVSVLTVLHQDWLNGKEY KCKVSNKALPAPIEKTISKAKGQPREPQVYTL PSREEMTKNQVSLTCLVKGFYPSDIAVEWESN GQPENNYKTTPPVLDSDGSFFLYSKLTVDKSR WQQGNVFSCSVMHEALHNHYTQKSLSLSPGK GGGSGGGGSGGGGSGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K</p>
413.	CL1 9-G4 CC x PSMA 76-B10 x I2C x scFc - Full Sequence	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKVSKASGYTFTD YYMHWVRQAPGQCLEWMGWNPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGTLLTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSQVQLVE SGGGLVKPGESLRLSCAASGFTFSDYYMYWVR QAPGKGLEWVAIISDGGYYTYYSDIKGRFTISR DNAKNSLYLQMNSLKAEDTAVYYCARGFPLL RHGAMDYWGQGTLLTVSSGGGGSGGGGSGG GGSDIQMTQSPSSLSASVGDRVTITCKASQNV TNVAWYQQKPGQAPKSLIYSASYRYSVPSRF SGSASGTDFTLTISSVQSEDFATYYCQQYDSYP YTFGGGTKLEIKSGGGGSEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWWVRQAPGKGL EWVARIRSKYNNYATYYADSVKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRHGNFGNSY ISYWAYWGQGTLLTVSSGGGGSGGGGSGGGG SQTVVTQEPSTLVSPGGTVTLTCGSSTGAVTSG NYPNWVQKPGQAPRGLIGGTFKFLAPGTPARF SGSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGGGTKLTVLGGGGDKTHTCPPCPAPELL GGPSVFLFPPKPKDTLMISRTPVTCVVVDVSH EDPEVKFNWYVDGVEVHNAKTKPCEEQYGST YRCVSVLTVLHQDWLNGKEYKCKVSNKALPA PIEKTISKAKGQPREPQVYTLPSREEMTKNQV SLTCLVKGFYPSDIAVEWESNGQPENNYKTTP VLDSGSDGSFFLYSKLTVDKSRWQQGNVFSCSVM HEALHNHYTQKSLSLSPGKGGGGSGGGGSGG GGSGGGGSGGGGSGGGGSDKTHTCPPCPAPEL</p>

				LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTTTPVLDSGDGSFFLYSKLTVDKSRWQQGNVFCSS VMHEALHNHYTQKSLSLSPGK
414.	EGFRvIII_CC_x_ I2C x scFc - Full Sequence	artificial	Aa	QVQLVESGGGVVQSGRSLRLSCAASGFTFRNY GMHWVRQAPGKCLEWVAWIWYDGSDDKYAD SVRGRFTISRDNKNTLYLQMNSLRAEDTAVY YCARDGYDILTGNPRDFDYWGQGLVTVSSG GGGSGGGGSGGGGSDTVMQTPLSSHVTLGQP ASISCRSSQSLVHSDGNTYLSWLQQRPGQPPRL LIYRISRRFSGVPDRFSGSGAGTDFTLEISRVEA EDVGVVYCMQSTHVPRTFGCGTKVEIKSGGG GSEVQLVESGGGLVQPGGSLKLSCAASGFTFN KYAMNWVRQAPGKGLEWVARIRSKYNNYAT YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGLVTV SSGGGSGGGGSGGGGSGTQVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQQKPGQAP RGLIGGTFKFLAPGTPARFSGSLLGGKAALTLG VQPEDEAEYCYLWYSNRWVFGGGTKLTVLG GGGDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSYRCVSVLTVLHQD WLNGKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSD IAVEWESNGQPENNYKTTTPVLDSGDGSFFLYS KLTVDKSRWQQGNVFCSSVMHEALHNHYTQKS LSLSPGKGGGGSGGGGSGGGGSGGGGSGGGG SGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK KDTLMISRTPEVTCVVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSYRCVSVLTVLH QDWLNGKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTTPVLDSGDGSFFLYS KLTVDKSRWQQGNVFCSSVMHEALHNHYTQK LSLSPGK
415.	EpCAM 5-10 x I2C x scFc - Full Sequence	artificial	aa	ELVMTQSPSSLTVTAGEKVTMSCKSSQSLLS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGG EVQLLEQSGAELVRPGTSVKISCKASGYAFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGQGTITVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSCAASGFTFNKYAM NWVRQAPGKGLEWVARIRSKYNNYATYYADS VKDRFTISRDDSKNTAYLQMNNLKTEDTAVY YCVRHGNFGNSYISYWAYWGQGLVTVSSGG GGSGGGGSGGGGSGTQVVTQEPSLTVSPGGT LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTFKFLAPGTPARFSGSLLGGKAALTLGSGVQPE DEAEYCYLWYSNRWVFGGGTKLTVLGGGGD KHTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR

				TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG K
416.	FL 4-E9 CC x I2C x scFc - Full Sequence	artificial	aa	QVTLKESGPTLVKPTETLTLTCTVSGFSFRNAR MGVSWIRQPPGKCLEWLAHIFSNDEKSYSTSL KSRLTISKDTSKSQVVLMTNMDPVDATYFC ARVPEYSSGWYRFDYWGQGLTVTVSSGGGG GGGGSGGGSDIQMTQSPSSLSASVGDRVTISC RASQSIRSYLNWYQQKPGKAPKLLIYATSSLQG GVPSRFSGSGTDFTLTISSLQPEDFATYYCQQ SYSTPFTFGCGTKVEIKSGGGGSEVQLVESGGG LVQPGGSLKLSAASGFTFNKYAMNWVRQAP GKGLEWVARIRSKYNNYATYYADSVKDRFTIS RDDSKNTAYLQMNNLKTEDTAVYYCVRHGNF GNSYISYWAYWGQGLTVTVSSGGGGSGGGG GGGGSTVVTQEPSLTVSPGGTVTLTCGSSTG AVTSGNYPNWVQQKPGQAPRGLIGGKFLAPG TPARFSGSLLGGKAALTLSGVQPEDEAEYYCV LWYSNRWVFGGGTKLTVLGGGGDKTHTCPPC PAPELLGGPSVFLFPPKPKDTLMISRTPEVTCV VDVSHEDPEVKFNWYVDGVEVHNNAKTKPCEE QYGSTYRCVSVLTVLHQDWLNGKEYKCKVSN KALPAPIEKTISKAKGQPREPQVYTLPPSREEM TKNQVSLTCLVKGFYPSDIAVEWESNGQPEN YKTTTPVLDSGDSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGK
417.	MS 5-F11 x I2C x scFc - Full Sequence	artificial	aa	QVQLVESGGGLVKPGGSLRLSAAASGFTFSDY YMTWIRQAPGKGLEWLSYISSSGSTIYYADSV KGRFTISRDNKNSLFLQMNSLRAEDTAVYYC ARDRNSHFYWGQGLTVTVSSGGGGSGGGG GGGGSDIQMTQSPSSVSASVGDRVTITCRASQG INTWLAWYQQKPGKAPKLLIYGASGLQSGVPS RFSGSGSGTDFTLTISSLQPEDFATYYCQAKSF PRTFGQGTKVEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSAASGFTFNKYAMNWVRQAPGKG LEWVARIRSKYNNYATYYADSVKDRFTISRDD

				SKNTAYLQMNNLKTEDTAVYYCVRHGNFGNS YISYWAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPLTVSPGGTVLTCGSSTGAVTS GNYPNWVQQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGGGTKLTVLGGGGDKTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGSGGGGS GGGGSGGGSGGGSGGGGSDKTHTCPPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGK
418.	MSLN H2 x I2C xscFc - Full Sequence	artificial	Aa	EVQLLESGGGVVQPGRSLRLSCAASGFTFSSYG MGWVRQAPGKGLEWVAVISYDGSNKYYADS VKGRFTISRDNKNTLYLQMNSLRAEDTAVYY CAREGAHFGSGSYPLYYYYAMDVWGQGT TVTVSSGGGGSGGGSGGGGSELTLTQSPGTL LSPGERATLSCRASQSVSSSYLAWYQQKPGQA PRLLIYGASIRATGIPDRFSGSGGTDFLTISRL EPEDFAVYYCQQYGSPIFTFGPGTKVEIKSGG GGSEVQLVESGGGLVQPGGSLKLSAASGFTF NKYAMNWVRQAPGKGLEWVARIRSKYNNYA TYYADSVKDRFTISRDDSKNTAYLQMNNLKTE DTAVYYCVRHGNFGNSYISYWAYWGQGLTV VSSGGGGSGGGSGGGGSGQTVVTQEPLTVSP GGTVLTCGSSTGAVTSGNYPNWVQQKPGQA PRGLIGGTKFLAPGTPARFSGSLLGGKAALTL GVQPEDEAEYYCVLWYSNRWVFGGGTKLTVL GGGGDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSD IAVEWESNGQPENNYKTTPPVLDSDGSFFLYSK LTVDKSRWQQGNVFSVMHEALHNHYTQKS LSLSPGKGGGGSGGGSGGGGSGGGGSGGGG SGGGGSDKTHTCPPCPAPELLGGPSVFLFPPK KDTLMISRTPEVTCVVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSTYRCVSVLTVLH QDWLNGKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTPPVLDSDGSFFLYS KLTVDKSRWQQGNVFSVMHEALHNHYTQK LSLSPGK
419.	PSMA 76-B10 x FL 4-E9 CC x I2C xscFc - Full	artificial	Aa	QVQLVESGGGLVKPGESLRLSCAASGFTFSDY YMYWVRQAPGKGLEWVAIISDGGYYTYYSII KGRFTISRDNKNSLYLQMNSLKAEDTAVYYC

	Sequence			<p>ARGFPLLRHGAMDYWGQGTLLTVSSGGGGSG GGGSGGGGSDIQMTQSPSSLSASVGDRVITICK ASQNVDTNVAWYQQKPGQAPKSLIYSASYRY SDVPSRFGSASGTDFTLTISVQSEDFATYYCQ QYDSYPYTFGGGKLEIKSGGGGSQVTLKESG PTLVKPTETLTLTCTVSGFSFRNARMGVSWIRQ PPGKCLEWLAHIFSNDEKSYSTSLKSRLTISKDT SKSQVVLMTNMDPVDATYFCARVPEYSSG WYRFDYWGQGTLLTVSSGGGGSGGGGSGGG GSDIQMTQSPSSLSASVGDRVITISCRASQSIRSY LNWYQQKPGKAPKLLIYATSSLQGGVPSRFSG SGSGTDFTLTISLQPEDFATYYCQQSYSTPFTF GCGTKVEIKSGGGGSEVQLVESGGGLVQPGGS LKLSCAASGFTFNKYAMNWVRQAPGKGLEW VARIRSKYNNYATYYADSVKDRFTISRDDSKN TAYLQMNNLKTEDTAVYYCVRHGNFGNSYIS YWAYWGQGTLLTVSSGGGGSGGGGSGGGGS QTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGN YPNWWQQKPGQAPRGLIGGTKFLAPGTPARFS GSLGGAALTLGSGVQPEDEAEYYCVLWYSN RWVFGGGTKLTVLGGGGDKTHTCPPCPAPEL GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH EDPEVKFNWYVDGVEVHNAKTKPCEEQYGST YRCVSVLTVLHQDWLNGKEYKCKVSNKALPA PIEKTISKAKGQPREPQVYTLPPSREEMTKNQV SLTCLVKGFYPSDIAVEWESNGQPENNYKTTPP VLDSGDSFFLYSKLTVDKSRWQQGNVFCFSVM HEALHNHYTQKSLSLSPGKGGGGSGGGGSGG GGSGGGGSGGGGSGGGGSDKHTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTTP PPVLDSDGDSFFLYSKLTVDKSRWQQGNVFCFS VMHEALHNHYTQKSLSLSPGK</p>
420.	VH CDR1 CD3 B2	artificial	Aa	GFTFNKYAIN
421.	VH CDR2 CD3 B2	artificial	Aa	RIRSKYNNYATYYADQVK
422.	VH CDR3 CD3 B2	artificial	Aa	HANFGNSYISYWAY
423.	VL CDR1 CD3 B2	artificial	Aa	ASSTGAVTSGNYPN
424.	VL CDR2 CD3 B2	artificial	Aa	GTKFLVP
425.	VL CDR3 CD3 B2	artificial	Aa	TLWYSNRWV
426.	H VL CD3 B2 binder	artificial	Aa	<p>EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AINWVRQAPGKGLEWVARIRSKYNNYATYYA DQVKDRFTISRDDSKNTAYLQMNNLKTEDTAV VYYCVR HANFGNSYISYWAYWGQGTLLTVSSGGGGSG GGGSGGGGSQTVVTQEPSLTVSPGGTVTLTCA SSTGAVTSGNYPNWWQQKPGQAPRGLIGGTKF LVP</p>

				GTPARFSGSLLGGKAALTLSGVQPEDEAEYYC TLWYSNRWVFGGGTKLTVL
427.	CL1 9-G4 CC xI2Ccc xHSA xFL 4-E9 CC xI2Ccc	artificial	Aa	<p> QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YMHWRQAPGQCLEWMGWNPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSLRSDDTAV YYCAREKHAVAGIGFDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWV RQAPGKCLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLTVTVSSGGGGSG GGGSGGGGSQTVVVTQEPSLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTFK LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGSGGG GSGGGGSDAHKSEVAHRFKDLGEENFKALVLI AFAQYLQQCPFEDHVKL VNEVTEFAKTCVADE SAENCDKSLHTLFGDKLCTVATLRETYGEMAD CCAKQEPERNECFLQHKDDNPPLRVRPEVD VMCTAFHDNEETFLLKLYEYIARRHPYFYAPE LLFFAKRYKAAFTECCQAADKAACLLPKLDEL RDEGKASSAKQRLKCASLQKGERAFKAWAV ARLSQRFPKAEFAEVSKLVTDLTKVHTECCHG DLLECADDRADLAKYICENQDSISSKLKECCEK PLEKSHCIAEVENDEMPADLPSLAADFVESKD VCKNYAEAKDVFLGMFLYEYARRHPDYSVVL LLRLAKTYETTLEKCCAAADPHECYAKVFDEF KPLVEEPQNLKQNCLEFELGGEYKFNALLVR YTKKVPQVSTPTLVEVSRNLGKVGSKCKHPE AKRMPCAEDYLSVVLNQLCVLHEKTPVSDRV TKCTESLVNRRPCFSALEVDETYVPKEFNAET FTFHADICTLSEKERQIKKQTALVELVKHKPKA TKEQLKAVMDDFAAFVEKCKKADDKETCFAE EGKKLVAASQAALGLGGGGSGGGGSGGGGSQ VTLKESGPTLVKPTETLTLTCTVSGFSFRNARM GVSWIRQPPGKCLEWLAHIFSNDEKSYSTSLKS RLTISKDTSKSQVVLMTNMDPVDTATYFCAR VPEYSSGWYRFDYWGQGLTVTVSSGGGGSGG GSGGGGSDIQMTQSPSSLSASVGDRVITISCR QSIRSILNYLNWYQQKPGKAPKLLIYATSSLQGG VPSRFSGSGSGTDFTLTISSLQPEDFATYYCQQS YSTPFTFGCGTKVEIKSGGGGSEVQLVESGGGL VQPGGSLKLSAASGFTFNKYAMNWVRQAPG KCLEWVARIRSKYNNYATYYADSVKDRFTISR DDSKNTAYLQMNNLKTEDTAVYYCVRHGNFG NSYISYWAYWGQGLTVTVSSGGGGSGGGGGSG GGGSQTVVTQEPSLTVSPGGTVTLTCGSSTGA VTSGNYPNWVQQKPGQAPRGLIGGTFKFLAPGT PARFSGSLLGGKAALTLSGVQPEDEAEYYCVL WYSNRWVFGCGTKLTVL </p>
428.	CL1 9-G4 CC xI2Ccc xFL 4-E9 CC xI2Ccc	artificial	Aa	<p> QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YMHWRQAPGQCLEWMGWNPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSLRSDDTAV </p>

				<p>YYCAREKHAVAGIGFDYWGQGTTLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWV RQAPGKCLEWVARIRSKYNNYATYYADS VKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGTTLVTVSSGGGGSG GGGSGGGGSQT VVTQEPLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTFK LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGSQVTL KESGPTLVKPTETLTLTCTVSGFSFRNARMGVS WIRQPPGKCLEWLAHIFSNDEKSYSTSLKSRLTI SKDTSK SQVVTMTNMDPVDTATYFCARVPE YSSGWYRFDYWGQGTTLVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSLSASVGDRVTISCRASQSI RSYLNWYQQKPGKAPKLLIYATSSLQGGVPSR FSGSGSGTDFTLTISLQPEDFATYYCQSYSTP FTFGCGTKVEIKSGGGGSEVQLVESGGGLVQP GGSLKLSAASGFTFNKYAMNWVRQAPGKCL EVARIRSKYNNYATYYADS VKDRFTISRDDS KNTAYLQMNNLKTEDTAVYYCVRHGNFGNSY ISYWAYWGQGTTLVTVSSGGGGSGGGGGGGGG SQT VVTQEPLTVSPGGTVTLTCGSSTGAVTSG NYPNWVQQKPGQAPRGLIGGTFKFLAPGTPARF SGSLLGGKAALTLSGVQPEDEAEYYCVLWYSN RWVFGCGTKLTVL</p>
429.	<p>CL1 9-G4 CC xI2Ccc x(EAAAK)10xFL 4-E9 CC xI2Ccc</p>	artificial	Aa	<p>QVQLVQSGAEVKKPGASVKV SCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSR LRSDDTAV YYCAREKHAVAGIGFDYWGQGTTLVTVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFTLTISLQPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWV RQAPGKCLEWVARIRSKYNNYATYYADS VKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGTTLVTVSSGGGGSG GGGSGGGGSQT VVTQEPLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTFK LAPGTPARFSGSLLGGKAALTLSGVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLEAAAKEAA AKEAAAKEAAAKEAAAKEAAAKEAAAKEAA AKEAAAKEAAAKQVTLKESGPTLVKPTETLTL TCTVSGFSFRNARMGVS WIRQPPGKCLEWLAH IFSNDEKSYSTSLKSRLTISKDTSK SQVVTMTN MDPVDTATYFCARVPEYSSGWYRFDYWGQGT LVTVSSGGGGSGGGGSGGGGSDIQMTQSPSSL SASVGDRVTISCRASQSIRSYLNWYQQKPGKA PKLLIYATSSLQGGVPSRFSGSGSGTDFTLTISL QPEDFATYYCQSYSTPFTFGCGTKVEIKSGGG GSEVQLVESGGGLVQPGGSLKLSAASGFTFN KYAMNWVRQAPGKCLEWVARIRSKYNNYAT</p>

				YYADSVKDRFTISRDDSKNTAYLQMNNLKTED TAVYYCVRHGNFGNSYISYWAYWGQGLVTV SSGGGSGGGGSGGGGSQTVVVTQEPSLTVSPG GTVTLTCGSSTGAVTSGNYPNWVQQKPGQAP RGLIGGTKFLAPGTPARFSGSLLGGKAALTLG VQPEDEAEYYCVLWYSNRWVFGCGTKLTVL
430.	CL1 9-G4 CC xI2Ccc -scFc - scFc2 xFL 4-E9 CC xI2Ccc	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YYMHWVRQAPGQCLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHEMELSRLSDDTAV YYCAREKHAVAGIGFDYWGQGLVTVVSSGGG GSGGGGSGGGGSDIQMTQSPSSVSASVGDRVTI TCQASQDISNYLNWYQQKPGKAPKLLIYAASS LESGVPSRFSGSGSGTDFLTITSSLPEDFATYY CQQANSFPLTFGCGTKVDIKSGGGGSEVQLVE SGGGLVQPGGSLKLSAASGFTFNKYAMNWV RQAPGKCLEWVARIRSKYNNYATYYADSVKD RFTISRDDSKNTAYLQMNNLKTEDTAVYYCVR HGNFGNSYISYWAYWGQGLVTVVSSGGGSG GGGSGGGGSQTVVVTQEPSLTVSPGGTVTLTCG SSTGAVTSGNYPNWVQQKPGQAPRGLIGGTKF LAPGTPARFSGSLLGGKAALTLGTVQPEDEAE YYCVLWYSNRWVFGCGTKLTVLGGGGSGGG GSGGGGSDKTHTCPPCPAPPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HLDWLNQKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYKTTTPVLDSDGSFFL YSKLTVDKSRWQQGNVFSVCSVMHEALHNHYT QKSLSLSPGKGGGGSGGGGSGGGGSGGGGSG GGGSGGGGSDKTHTCPPCPAPPELLGGPSVFLF PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVL VLHLDWLNQKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTTTPVLDSDGSF FLYSKLTVDKSRWQQGNVFSVCSVMHEALHNH YTQKSLSLSPGKDKTHTCPPCPAPPELLGGPSV FLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEV FNWYVDGVEVHNAKTKPCEEQYGSTYRCVSV LTVLHLDWLNQKEYKCKVSNKALPAPIEKTIS KAKGQPREPQVYTLPPSREEMTKNQVSLTCLV KGFYPSDIAVEWESNGQPENNYKTTTPVLDSD GSFFLYSKLTVDKSRWQQGNVFSVCSVMHEAL HNHYTQKSLSLSPGKGGGGSGGGGSGGGGSG GGGSGGGGSGGGGSDKTHTCPPCPAPPELLGGP SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDP EVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC VSVLTVLHLDWLNQKEYKCKVSNKALPAPIEK TISKAKGQPREPQVYTLPPSREEMTKNQVSLT LVKGFYPSDIAVEWESNGQPENNYKTTTPVLD SDGSFFLYSKLTVDKSRWQQGNVFSVCSVMHEA LHNHYTQKSLSLSPGKGGGGSGGGGSGGGGSG QVTLKESGPTLVKPTETLTCTVSGFSFRNAR MGVSWIRQPPGKCLEWLAHIFSNDEKSYSTSL KSRLTISKDTSKSKVVLTMNTMMDPVDATATYFC

				<p>ARVPEYSSGWYRFDYWGGQGLTVTVSSGGGGG GGGGSGGGSDIQMTQSPSSLSASVGDRVTISC RASQSIRS YLNWYQQKPGKAPKLLIYATSSLQG GVPSRFSGSGSGTDFTLTISLQPEDFATYYCQQ SYSTPFTFGCGTKVEIKSGGGGSEVQLVESGGG LVQPGGSLKLSAASGFTFNKYAMNWVRQAP GKCLEWVARIRSKYNNYATYYADSVKDRFTIS RDDSNTAYLQMNNLKTEDTAVYYCVRHGNF GNSYISYWAYWGGQGLTVTVSSGGGGSGGGG GGGGSQTVVTQEPSLTVSPGGTTLTLCGSSTG AVTSGNYPNWVQQKPGQAPRGLIGTKFLAPG TPARFSGSLLGGKAALTLGSGVQPEDEAEYYCV LWYSNRWVFGCGTKLTVL</p>
431.	EpCAM 5-10 xI2Ccc xHSA xH2 xI2Ccc	artificial	Aa	<p>ELVMTQSPSSLTVTAGEKVTMSCKSSQSLLNS GNQKNYLTWYQQKPGQPPKLLIYWASTRESG VPDRFTGSGSGTDFTLTISVQAEDLAVYYCQN DYSYPLTFGAGTKLEIKGGGGSGGGGSGGGG EVQLLEQSGAELVRPGTSVKISCKASGYFTNY WLGWVKQRPGHGLEWIGDIFPGSGNIHYNEKF KGKATLTADKSSSTAYMQLSSLTFEDSAVYFC ARLRNWDEPMDYWGGQTTVTVSSGGGGSEV QLVESGGGLVQPGGSLKLSAASGFTFNKYAM NWVRQAPGKCLEWVARIRSKYNNYATYYADS VKDRFTISRDDSNTAYLQMNNLKTEDTAVY YCVRHGNFNGNSYISYWAYWGGQGLTVTVSSGG GGSGGGSGGGGSQTVVTQEPSLTVSPGGTTL LTCGSSTGAVTSGNYPNWVQQKPGQAPRGLIG GTKFLAPGTPARFSGSLLGGKAALTLGSGVQPE EAEYYCVLWYSNRWVFGCGTKLTVLGGGGSG GGSGGGGSDAHKSEVAHRFKDLGEENFKAL VLIAFAQYLQCCPFEDHVKL VNEVTEFAKTCV ADESAENCDKSLHTLFGDKLCTVATLRETYGE MADCCAQEPERNECFLQHKDDNPNLPRVLP EVDVMCTAFHDNEETFLLKLYEYIARRHPYFY APELFFAKRYKAAFTECCQAADKAAACLLPKL DELRDEGKASSAKQRLKASLQKFGERAFAKA WAVARLSQRFPKAEFAEVSKLVTDLTKVHTEC CHGDLLCADRADLAKYICENQDSISSKLKE CCEKPLLEKSHCIAEVENDEMPADLPSLAADFV ESKDVCKNYAEAKDVFLGMFLYEYARRHPDY SVVLLRLAKTYETTLEKCCAAADPHECYAKV FDEFKPLVEEPQNLKQNCLEFEQLGEYKFQNA LLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCC KHPEAKRMPCAEDYLSVVLNQLCVLHEKTPVS DRVTKCCTESLVNRRPCFSALEVDETYVPKEF NAETFTFHADICTLSEKERQIKKQATALVELVKH KPKATKEQLKAVMDDFAAFVEKCKADDKET CFAEEGKKLVAASQAALGLGGGGSGGGGSGG GGSEVQLLES GGGVQGRSLRLSCAASGFTFS SYGMGWVRQAPGKGLEWVAVISYDGSNKYY ADSVKGRFTISRDNKNTLYLQMNSLRAEDTA VYYCAREGAHFGSGSYYPYLYYYAMDVWGQ GTTVTVSSGGGGSGGGGSGGGGSELTLTQSPG TSLSPGERATLSCRASQSVSSSYLAWYQQKPG QAPRLLIYGASIRATGIPDRFSGSGSGTDFTLTIS RLEPEDFAVYYCQQYGGSSPIFTFGPGTKVEIKS</p>

				GGGGSEVQLVESGGGLVQPGGSLKLSCAASGFTFNKYAMNWVRQAPGKCLEWVARIRSKYNNYATYYADSVKDRFTISRDDSKNTAYLQMNNLKTEDTAVYYCVRHGNFGNSYISYWAYWGQGLVTVSSGGGGSGGGGSGGGGSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTSGNYPNWVQKPGQAPRGLIGGTKFLAPGTPARFSGSLLGGKAALTLSGVQPEDEAEYYCVLWYSNRWVFGCGTKLTVL
432.	CH3 15-E11_1_VAG_CC - HCDR2	artificial	Aa	NIAYGVAGTNYNQKFQG
433.	CH3 15-E11_1_VAG_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKVSKKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSS
434.	CH3 15-E11_1_VAG_CC x I2L x G4 x scFc x G4 x MS 15-B12 CC x I2L clipopt_DI	artificial	Aa	QVQLVQSGAEVKKPGASVKVSKKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGGQ GGGGQGGGGQDIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQKPGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFLTISLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGQEVQLVES GGGGLVQPGGSLKLSCAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGLTVTVSSGGGGGQGG GGGGGGQQT VVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWVQKPGQAPRGLIGGTKFL APGTPARFSGSLEGGKAALTLSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGCPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQQDWLNGKEYKCKVSNKAL PAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQGGG GQGGGGQGGGGQGGGGQGGGGQCPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEEPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PVLDSGDSFFLYSKLTVDKSRWQQGNVFS CS VMHEALHNHYTQKSLSLSPGKGGGGQVQLQE SGPGLVKPSETLSLTCTVSGGSISSSYFWGWIR QPPGKCLEWIGNIYYSGSSNYNPSLKSRVTISV DTSKNQFSKLSSVTAADTAVYYCARLPRGDR DAFDIWGQGMVTVSSGGGGQGGGGQGGGG QDIVMTQSPSSLSASVGDRVTITCRASQGISNY LAWYQKPGKVPKLLIYAASLTQSGVPSRFSG SSGTDFLTISLQPEDFATYYCQQS YSTPFTF GCGTKVEIKSGGGQEVQLVESGGGLVQPGGS LKLSCAASGFTFNKYAMNWVRQAPGKGMW

				VARIRSKYNNYATYYADAVKDRFTISRDDSKN TLYLQMNNLKTEDTAVYYCVRAGNFGSSYISY FAYWGQGTLLTVSSGGGGQGGGGQGGGGQ TVVTQEPLSLTVSPGGTVTITCGSSTGAVTSGNY PNWIQKKPGQAPRGLIGGTKFLAPGTPARFSGS LEGGKAALTLSGVQPEDEAEYYCVLYYSNRW VFGSGTKLTVL
435.	CH3 15- E11 ₁ VAG _{CC} x I2L x G4 x scFc x G4 x MS 15- B12 CC x I2L clipopt_EI	artificial	Aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRLSDDTA VYYCATRYFYVMDYWGQGTLLTVSSGGGGQ GGGGQGGGGQEIQMTQSPSSLSASVGDRVITIT CRASQDISNYLNWYQQKPGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFLTISSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAMNWVR QAPGKGMWVVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGTLLTVSSGGGGQGG GGQGGGGQQT VVTQEPLSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGTKFL APGTPARFSGSLEGGKAALTLSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGCPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQGGG GQGGGGQGGGGQGGGGQGGGGQCPCPAP LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEEPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGQVQLQE SGPGLVKPSETLSLTCTVSGGSISSSSYFWGWIR QPPGKCLEWIGNIYYSGSSNYPNPSLKSRTISV DTSKNQFSKLSSVTAADTAVYYCARLPRGDR DAFDIWGQGTMTVTVSSGGGGQGGGGQGGGG QEIVMTQSPSSLSASVGDRVITICRASQGISNYL AWYQQKPGKVPKLLIYAASLQSGVPSRFSGS GSGTDFLTISSLQPEDFATYYCQQSYSTPFTFG CGTKVEIKSGGGGQEVQLVESGGGLVQPGGSL KLSAASGFTFNKYAMNWVRQAPGKGMWV ARIRSKYNNYATYYADAVKDRFTISRDDSKNT LYLQMNNLKTEDTAVYYCVRAGNFGSSYISYF AYWGQGTLLTVSSGGGGQGGGGQGGGGQQT VVTQEPLSLTVSPGGTVTITCGSSTGAVTSGNYP NWIQKKPGQAPRGLIGGTKFLAPGTPARFSGSL EGGKAALTLSGVQPEDEAEYYCVLYYSNRWV FSGTKLTVL
436.	I2M - HCDR1	artificial	Aa	KYAMN
437.	I2M - HCDR2	artificial	Aa	RIRSKYNNYATYYADAVKD

438.	I2M - HCDR3	artificial	Aa	AGNFGTSYISYWAY
439.	I2M - LCDR1	artificial	Aa	GSSTGAVTSGNYPN
440.	I2M - LCDR2	artificial	Aa	GTKFLAP
441.	I2M - LCDR3	artificial	Aa	VLWYSNRWV
442.	I2M - VH	artificial	Aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGTSYISYWAYWGQGLTVTVSS
443.	I2M - VL	artificial	Aa	QTVVTQEPSLTVSPGGTVTITCGSSTGAVTSGN YPNWVQKKPGQAPRGLIGGTKFLAPGTPARFS GSLSGKAALTLGVPPEDEAEYYCVLWYSN RWVFGSGTKLTVL
444.	IgG4 subtype hinge	artificial	Aa	ESKYGPPCPCP
445.	IgG1 subtype hinge	artificial	Aa	EPKSCDKTHTCPPCP
446.	EpCAM_19124- A6_CC - HCDR1	artificial	Aa	RYDMH
447.	EpCAM_19124- A6_CC - HCDR2	artificial	Aa	IISYDGSNKYYGDAVKG
448.	EpCAM_19124- A6_CC - HCDR3	artificial	Aa	RAGFQDF
449.	EpCAM_19124- A6_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
450.	EpCAM_19124- A6_CC - LCDR2	artificial	Aa	DVSSRPS
451.	EpCAM_19124- A6_CC - LCDR3	artificial	Aa	SSYTSSTWV
452.	EpCAM_19124- A6_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMHWVRQAPGQCLEWMAIISYDGSNKYYGD AVKGRFTISRDNRSRNTLYLQMNSLRAEDTAVY HCVKRAFQDFWGQGLTVTVSS
453.	EpCAM_19124- A6_CC - VL	artificial	Aa	QSALTQPPSVSGSPGQSITISCTGTSSDVGGYNY VSWYQQHPGKAPKLMYDSSRPSGVSNRFSG SKSGNTASLTISGLQAEDEADYYCSSYTSSTW VFGCGTKLTVL
454.	EpCAM_19124- B5_CC - HCDR1	artificial	Aa	DYGMH
455.	EpCAM_19124- B5_CC - HCDR2	artificial	Aa	GISWNSGNIGYADSVKG
456.	EpCAM_19124- B5_CC - HCDR3	artificial	Aa	PDCSSTSCYRGYYFDY
457.	EpCAM_19124- B5_CC - LCDR1	artificial	Aa	GGNNIGSKSVH
458.	EpCAM_19124- B5_CC - LCDR2	artificial	Aa	DVSDRPS
459.	EpCAM_19124- B5_CC - LCDR3	artificial	Aa	QVWDSNTDHVV
460.	EpCAM_19124- B5_CC - VH	artificial	Aa	EVQLVESGGGLVQPGRSLRLSCAASGFTFDDY GMHWVRQAPGKCLEWVSGISWNSGNIGYADS VKGRFTISRDNKNSLYLQMNSLRAEDTALYY CAKPCDSSTSCYRGYYFDYWGQGLTVTVSS
461.	EpCAM_19124- B5_CC - VL	artificial	Aa	SYVLTQPASVSVAPGQTARITCGNNIGSKSVH WYQQKPGQAPILVYDVS DRPSGIPERFSGSNS GNTATLTISRVEAGDEADYYCQVWDSNTDHV

				VFGCGTKLTVL
462.	EpCAM_19124-C5_CC - HCDR1	artificial	Aa	SYAII
463.	EpCAM_19124-C5_CC - HCDR2	artificial	Aa	GIIPMFGTANYAQKFQG
464.	EpCAM_19124-C5_CC - HCDR3	artificial	Aa	VSGTYHWGY
465.	EpCAM_19124-C5_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
466.	EpCAM_19124-C5_CC - LCDR2	artificial	Aa	DVSARPS
467.	EpCAM_19124-C5_CC - LCDR3	artificial	Aa	SSYISSTSLV
468.	EpCAM_19124-C5_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGSSVKVSCASGGTFSSY AIIWVRQAPGQCLEWMGGIIPMFGTANYAQKF QGRVTITADESTSTAYMELSSLRSED TAVYYC ARVSGTYHWGYWGQGLTVTVSS
469.	EpCAM_19124-C5_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSSDVGGYN YVSWYQQHPGKAPKLMYDVSARPSGVS NRFS GSKSGNTASLTISGLQAEDEADYYCSSYISSTSL VFGCGTKLTVL
470.	EpCAM_19124-C7_N67Q_CC - HCDR1	artificial	Aa	NYDMN
471.	EpCAM_19124-C7_N67Q_CC - HCDR2	artificial	Aa	VISYDGSQKSYSDSVKG
472.	EpCAM_19124-C7_N67Q_CC - HCDR3	artificial	Aa	RGATPFDY
473.	EpCAM_19124-C7_N67Q_CC - LCDR1	artificial	Aa	TGTSNDVGGYNYVS
474.	EpCAM_19124-C7_N67Q_CC - LCDR2	artificial	Aa	DVSSRPS
475.	EpCAM_19124-C7_N67Q_CC - LCDR3	artificial	Aa	SSYARSRTFVA
476.	EpCAM_19124-C7_N67Q_CC - VH	artificial	Aa	QVQLVESGGGVLPGRSLRLSCAASGFTFRNY DMNWVRQVPGKCLEWVAVISYDGSQKSYSDS VKGRFTISRDN SKNTLSLQMN SLRNED TAVYY CAKRGATPFDYWGQGLTVTVSS
477.	EpCAM_19124-C7_N67Q_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSDVGGYN YVSWYQQHPGKAPKLMYDVSRRPSGISNRFS GSKSGNTASLTISGLQAEDEADYYCSSYARSRT FVAFGCGTKLTVL
478.	EpCAM_19124-C7_S69Y_CC - HCDR1	artificial	Aa	NYDMN
479.	EpCAM_19124-C7_S69Y_CC - HCDR2	artificial	Aa	VISYDGSNKYSDSVKG
480.	EpCAM_19124-C7_S69Y_CC -	artificial	Aa	RGATPFDY

	HCDR3			
481.	EpCAM_19124-C7_S69Y_CC - LCDR1	artificial	Aa	TGTSNDVGGYNYVS
482.	EpCAM_19124-C7_S69Y_CC - LCDR2	artificial	Aa	DVSSRPS
483.	EpCAM_19124-C7_S69Y_CC - LCDR3	artificial	Aa	SSYARSRTFVA
484.	EpCAM_19124-C7_S69Y_CC - VH	artificial	Aa	QVQLVESGGGVLPGRSLRLSCAASGFTFRNY DMNWVRQVPGKCLEWVAVISYDGSNKYYSDS VKGRFTISRDNKNTLSLQMNSLRNEDTAVYY CAKRGATPFDYWGQGLVTVSS
485.	EpCAM_19124-C7_S69Y_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSNDVGGYN YVSWYQQHPGKAPKLMYDVSSRPSGISNRFS GSKSGNTASLTISGLQAEDEADYYCSSYARSRT FVAFGCGTKLTVL
486.	EpCAM_19124-D3_CC - HCDR1	artificial	Aa	NYDMN
487.	EpCAM_19124-D3_CC - HCDR2	artificial	Aa	VISYDGS DKHYTDSVKG
488.	EpCAM_19124-D3_CC - HCDR3	artificial	Aa	RGATPVDY
489.	EpCAM_19124-D3_CC - LCDR1	artificial	Aa	KSSQSLLHSNGYNYLG
490.	EpCAM_19124-D3_CC - LCDR2	artificial	Aa	FGSSRAS
491.	EpCAM_19124-D3_CC - LCDR3	artificial	Aa	MQALQTPFT
492.	EpCAM_19124-D3_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGS DKHYTD SVKGRFTISRDNKNTLYLQMNSLRTEDTAVY YCAKRGATPVDYWGQGLVTVSS
493.	EpCAM_19124-D3_CC - VL	artificial	Aa	EIVMTQSPLSLPVTGEPASISCKSSQSLLHSNG YNYLGWYLQKPGQSPQLLIYFGSSRASGVPDR FSGSGGTDFTLKISGVEAEDVGVYYCMQALQ TPFTFGCGTKVDIK
494.	EpCAM_19124-F5_CC - HCDR1	artificial	Aa	SYAII
495.	EpCAM_19124-F5_CC - HCDR2	artificial	Aa	GIPIFGTANYAQKFQG
496.	EpCAM_19124-F5_CC - HCDR3	artificial	Aa	VSGTYHWGY
497.	EpCAM_19124-F5_CC - LCDR1	artificial	Aa	TGTSSDIGSFNLVS
498.	EpCAM_19124-F5_CC - LCDR2	artificial	Aa	EGYKRPS
499.	EpCAM_19124-F5_CC - LCDR3	artificial	Aa	SSYISSSTLV
500.	EpCAM_19124-F5_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGSSVKVSCKVSGGTFSSY AIWVRQAPGQCLEWMGGIPIFGTANYAQKFQ GRVTITADESTSTAYMELSSLRSDDTAVYYCA RVSGTYHWGYWGQGLVTVSS
501.	EpCAM_19124-	artificial	Aa	QSALTQPPSASGSPGQSITISCTGTSSDIGSFNLV

	F5_CC - VL			SWYQQHPGKAPKLMYEGYKRPSGVSDRFSGS KSGNTASLTISGLQAEDEADYYCSSYISSSTLVF GCGTKLTVL
502.	EpCAM_19124- G7_CC - HCDR1	artificial	Aa	RYWMS
503.	EpCAM_19124- G7_CC - HCDR2	artificial	Aa	EINPDSSTINYTPSLKD
504.	EpCAM_19124- G7_CC - HCDR3	artificial	Aa	YPWFTY
505.	EpCAM_19124- G7_CC - LCDR1	artificial	Aa	RSSQSLVHSNGNTYLH
506.	EpCAM_19124- G7_CC - LCDR2	artificial	Aa	KVSNRFS
507.	EpCAM_19124- G7_CC - LCDR3	artificial	Aa	SQSTHVPFT
508.	EpCAM_19124- G7_CC - VH	artificial	Aa	EVQLVESGGGLVQPGGSLKLSAASGFDFSR Y WMSWVRQAPGKCLEWIGEINPDSSTINYTPSL KDKFIVSRDNAKNTLYLQMSKVRSEDTALYYC ARYPWFTYWGQGLTVTVSS
509.	EpCAM_19124- G7_CC - VL	artificial	Aa	EIVMTQTPLSLPVSLGDQASISCRSSQSLVHSNG NTYLHWYLQKPGQSPKLLIYKVSNRFSGVPDR FSGSGGTDFTLKISRVEAEDLGVYFCSQSTHV PFTFGCGTKLEIK
510.	EpCAM_19124- H1T69Y_CC - HCDR1	artificial	Aa	NYDMN
511.	EpCAM_19124- H1T69Y_CC - HCDR2	artificial	Aa	VISYDGSNKYYTDSVKG
512.	EpCAM_19124- H1T69Y_CC - HCDR3	artificial	Aa	RGATPVDY
513.	EpCAM_19124- H1T69Y_CC - LCDR1	artificial	Aa	RSSQSLVHSNGYNYLG
514.	EpCAM_19124- H1T69Y_CC - LCDR2	artificial	Aa	LGSSRAS
515.	EpCAM_19124- H1T69Y_CC - LCDR3	artificial	Aa	MQALQTPFT
516.	EpCAM_19124- H1T69Y_CC - VH	artificial	Aa	EVQLLESVGGGLVQGRSLRLSCAASGFTFRNY DMNWVRQVPGKCLEWVAVISYDGSNKYYTD SVKGRFTISRDNKNTLYLQMNSLRTEDTAVY YCAKRGATPVDYWGQGLTVTVSS
517.	EpCAM_19124- H1T69Y_CC - VL	artificial	Aa	EIVMTQSPLSLPVTPGEPASISCRSSQSLVHSNG YNYLGWYLQKPGQSPQLLIYLGSSRASGVPDR FSGSGGTDFTLKISRVEAEDVGVYVCMQALQ TPFTFGCGTKLEIK
518.	EpCAM_19124- H1_N67Q_CC - HCDR1	artificial	Aa	NYDMN
519.	EpCAM_19124- H1_N67Q_CC - HCDR2	artificial	Aa	VISYDGSQKTYTDSVKG

520.	EpCAM_19124-H1_N67Q_CC - HCDR3	artificial	Aa	RGATPVDY
521.	EpCAM_19124-H1_N67Q_CC - LCDR1	artificial	Aa	RSSQSLLSNGYNYLG
522.	EpCAM_19124-H1_N67Q_CC - LCDR2	artificial	Aa	LGSSRAS
523.	EpCAM_19124-H1_N67Q_CC - LCDR3	artificial	Aa	MQALQTPFT
524.	EpCAM_19124-H1_N67Q_CC - VH	artificial	Aa	EVQLLESGGGLVQPGRSLRLSCAASGFTFRNY DMNWVRQVPGKCLEWVAVISYDGSQKTYTDS VKGRFTISRDN SKNTLYLQMNSLR TEDTAVYY CAKRGATPVDYWGQGLTVTVSS
525.	EpCAM_19124-H1_N67Q_CC - VL	artificial	Aa	EIVMTQSPLSLPVTPGEPASISCRSSQSLLSNG YNYLGWYLQKPGQSPQLLIYLGSSRASGVPDR FSGSGGTDFTLKISRVEAEDVGVYYCMQALQ TPFTFGCGTKLEIK
526.	EpCAM_19125-A6_CC - HCDR1	artificial	Aa	RYDMH
527.	EpCAM_19125-A6_CC - HCDR2	artificial	Aa	IISYDGSNKYYGDAVKG
528.	EpCAM_19125-A6_CC - HCDR3	artificial	Aa	RAGFQDF
529.	EpCAM_19125-A6_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
530.	EpCAM_19125-A6_CC - LCDR2	artificial	Aa	EVSKRPA
531.	EpCAM_19125-A6_CC - LCDR3	artificial	Aa	SSYAGSNNWV
532.	EpCAM_19125-A6_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMHWVRQAPGQCLEWMAIISYDGSNKYYGD AVKGRFTISRDN SRNTLYLQMNSLRAEDTAVY HCVKRAGFQDFWQGLTVTVSS
533.	EpCAM_19125-A6_CC - VL	artificial	Aa	QSALTQPPSASGSPGQSVTISCTGTSSDVGGYN YVSWYRQHPGKAPKLMIEVSKRPAGVPDRFS GSKSGNTASLTVSGLQAEDEADYYCSSYAGSN NWFVFCGKLTVL
534.	EpCAM_19125-G6_C107A_CC - HCDR1	artificial	Aa	RYDMN
535.	EpCAM_19125-G6_C107A_CC - HCDR2	artificial	Aa	FISYDGSNEDYPAVKG
536.	EpCAM_19125-G6_C107A_CC - HCDR3	artificial	Aa	VGASPFDY
537.	EpCAM_19125-G6_C107A_CC - LCDR1	artificial	Aa	TGTSNDVGGYNYVS
538.	EpCAM_19125-G6_C107A_CC - LCDR2	artificial	Aa	EVSKRPS

539.	EpCAM_19125-G6_C107A_CC - LCDR3	artificial	Aa	ASYTGGRTYVG
540.	EpCAM_19125-G6_C107A_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMNWVRQAPGKCLEWVAFISYDGSNEDYPDA VKGRFTISRDN SKNTLYLQMNSLRAEDTAVYY CAKVGASPFDYWGQGLTVTVSS
541.	EpCAM_19125-G6_C107A_CC - VL	artificial	Aa	QSALTQPPSVSGSPGQSITISCTGTSNDVGGYN YVSWYQQHPGKAPKLM IYEVSKRPSGVPDRFS GSKSGNTASLTISGLQAEDEADYYCASYTGGR TYVGF GCGTKLTVL
542.	EpCAM_19125-G6_C107L_CC - HCDR1	artificial	Aa	RYDMN
543.	EpCAM_19125-G6_C107L_CC - HCDR2	artificial	Aa	FISYDGSNEDY PDAVKG
544.	EpCAM_19125-G6_C107L_CC - HCDR3	artificial	Aa	VGASPFDY
545.	EpCAM_19125-G6_C107L_CC - LCDR1	artificial	Aa	TGTSNDVGGYNYVS
546.	EpCAM_19125-G6_C107L_CC - LCDR2	artificial	Aa	EVSKRPS
547.	EpCAM_19125-G6_C107L_CC - LCDR3	artificial	Aa	LSYTGGRTYVG
548.	EpCAM_19125-G6_C107L_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMNWVRQAPGKCLEWVAFISYDGSNEDYPDA VKGRFTISRDN SKNTLYLQMNSLRAEDTAVYY CAKVGASPFDYWGQGLTVTVSS
549.	EpCAM_19125-G6_C107L_CC - VL	artificial	Aa	QSALTQPPSVSGSPGQSITISCTGTSNDVGGYN YVSWYQQHPGKAPKLM IYEVSKRPSGVPDRFS GSKSGNTASLTISGLQAEDEADYYCLSYTGGR TYVGF GCGTKLTVL
550.	EpCAM_19126-D5_CC - HCDR1	artificial	Aa	TYTIS
551.	EpCAM_19126-D5_CC - HCDR2	artificial	Aa	GIIPILGAPNYAQKFQG
552.	EpCAM_19126-D5_CC - HCDR3	artificial	Aa	DPFSRY
553.	EpCAM_19126-D5_CC - LCDR1	artificial	Aa	RSSQSLLSNGYNYLD
554.	EpCAM_19126-D5_CC - LCDR2	artificial	Aa	LGSNRAS
555.	EpCAM_19126-D5_CC - LCDR3	artificial	Aa	MQALQTPRT
556.	EpCAM_19126-D5_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGSSVKV SCKVSGGTFSTY TISWVRQAPGQCLEWMGGIIPILGAPNYAQKF QGRVSITADESTSTSYMELTSLRSED TAVYYCA RDPFSRYWGQGLTVTVSS
557.	EpCAM_19126-D5_CC - VL	artificial	Aa	EIVMTQSPLSLPVTPGEPASISCRSSQSLLSNG YNYLDWYLQKPGQSPQLLIYLG SNRASGVPDR

				FSGSGSGTDFTLKISRVEAEDVGVVYCMQALQ TPRTFGCGTKVEIK
558.	EpCAM_19127- B6_CC - HCDR1	artificial	Aa	SYAII
559.	EpCAM_19127- B6_CC - HCDR2	artificial	Aa	GIIPMFGTANYAQKFQG
560.	EpCAM_19127- B6_CC - HCDR3	artificial	Aa	VSGTYHWGY
561.	EpCAM_19127- B6_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
562.	EpCAM_19127- B6_CC - LCDR2	artificial	Aa	DVSARPS
563.	EpCAM_19127- B6_CC - LCDR3	artificial	Aa	SSYISITTLV
564.	EpCAM_19127- B6_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGSSVKVSCASGGTFRSY AIIWVVRQAPGQCLEWMGGIIPMFGTANYAQKF QGRVTITADESTSTAYMELSRLESDTAVYYC ARVSGTYHWGYWGQGLVTVSS
565.	EpCAM_19127- B6_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSSDVGGYN YVSWYQQRPGRAPKLMYDVSARPSGVSNRFS GSKSGNTASLTISGLQAEDEADYYCSSYISITTL VFGCGTKLTVL
566.	EpCAM_19127- G11_CC - HCDR1	artificial	Aa	RYDMH
567.	EpCAM_19127- G11_CC - HCDR2	artificial	Aa	IISYDGSIRYYADSVKG
568.	EpCAM_19127- G11_CC - HCDR3	artificial	Aa	RAGFQFDS
569.	EpCAM_19127- G11_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
570.	EpCAM_19127- G11_CC - LCDR2	artificial	Aa	EVSKRPA
571.	EpCAM_19127- G11_CC - LCDR3	artificial	Aa	SSYAGGNNFVV
572.	EpCAM_19127- G11_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRSY DMHWVVRQAPGQCLEWMAIISYDGSIRYYADS VKGRFTISRDNRSNTLYLQMNLSRAEDTAVYY CVKRAFQFDSWGQGLVTVSS
573.	EpCAM_19127- G11_CC - VL	artificial	Aa	QSALTQPPSASGSPGQSVTISCTGTSSDVGGYN YVSWYQQHPGKAPKLMYEVSKRPAGVPDRFS GSKSGNTASLTVSGLQAEDEADYYCSSYAGGN NFVVFGCGTKLTVL
574.	EpCAM_19128- H8_CC - HCDR1	artificial	Aa	EYWMS
575.	EpCAM_19128- H8_CC - HCDR2	artificial	Aa	EIIPDSSKINYTPSLKD
576.	EpCAM_19128- H8_CC - HCDR3	artificial	Aa	PLYGGYDEGFAY
577.	EpCAM_19128- H8_CC - LCDR1	artificial	Aa	RSSQSLVHSNGNTYLE
578.	EpCAM_19128- H8_CC - LCDR2	artificial	Aa	KVSNRFS

579.	EpCAM_19128-H8_CC - Lcdr3	artificial	Aa	FQGSHVPTYT
580.	EpCAM_19128-H8_CC - VH	artificial	Aa	EVQLVESGGGLVQPGRSLKLSAASGDFSEY WMSWVRQAPGKCLEWIGEIPDSSKINYTPSLK DKFIISRDNKNTLYLQMSKVRSEDTALYYCA RPLYYGDEGFAYWGQTTVTVSS
581.	EpCAM_19128-H8_CC - VL	artificial	Aa	EIVMTQTPLSLPVSLGDQASISCRSSQSLVHSNG NTYLEWYLQKPGQSPKLLIYKVSNRFSGVPDR FSGSGGTDFTLKISRVEAEDLGVYYCFQGS HV PYTFGCGTRLEIK
582.	EpCAM_19129-A4_CC - HCDR1	artificial	Aa	SYAMH
583.	EpCAM_19129-A4_CC - HCDR2	artificial	Aa	RVRKSDNYATYYADSVKD
584.	EpCAM_19129-A4_CC - HCDR3	artificial	Aa	PLFTTVEVTNALDY
585.	EpCAM_19129-A4_CC - Lcdr1	artificial	Aa	SASSISSNYLH
586.	EpCAM_19129-A4_CC - Lcdr2	artificial	Aa	RTSVLSS
587.	EpCAM_19129-A4_CC - Lcdr3	artificial	Aa	QQGSSMPFT
588.	EpCAM_19129-A4_CC - VH	artificial	Aa	EVQLVESGGGLVQPKGSLKLSAASGFTFNSY AMHWVRQAPGRRCMEWVGRVRSKSDNYATYY ADSVKDRFTISRDDSQSMYLYQMNNLKTEDTA IYYCVRPLFTTVEVTNALDYWGQGTTLTVSS
589.	EpCAM_19129-A4_CC - VL	artificial	Aa	EIVLTQSPTTMAASPGEKITITCSASSISSNYLH WYQQKPGFSPKLLIYRTSVLSSGVPARFSGSGS GTSYSLTIDTMEAEDVATYFCQQGSSMPFTFGC GTRLEIK
590.	EpCAM_19129-E3_CC - HCDR1	artificial	Aa	NYWMQ
591.	EpCAM_19129-E3_CC - HCDR2	artificial	Aa	AIYPGEGETRYTQKFKG
592.	EpCAM_19129-E3_CC - HCDR3	artificial	Aa	PYAGYYLYAMDQ
593.	EpCAM_19129-E3_CC - Lcdr1	artificial	Aa	RSSQSIVHSNGNTYLE
594.	EpCAM_19129-E3_CC - Lcdr2	artificial	Aa	KVSNRFS
595.	EpCAM_19129-E3_CC - Lcdr3	artificial	Aa	SQSTHVPTYT
596.	EpCAM_19129-E3_CC - VH	artificial	Aa	QVQLVQSGAELARPGASVKLSCKASGYIFSNY WMQWVKQRPGQCLEWIGAIYPGEGETRYTQK FKGKATLTADTSSSTAYMQLSSLASEDSAVYY CARPYAGYYLYAMDQWGQGTTLTVSS
597.	EpCAM_19129-E3_CC - VL	artificial	Aa	EIVMTQTPLSLPVSLGDQASISCRSSQSI VHSNG NTYLEWYLQKPGQSPKLLIYKVSNRFSGVPDR FSGSGGTDFTLKISRVEAEDLGVYFCSQSTHV PYTFGCGTRLEIK
598.	EpCAM_19130-C11_CC - HCDR1	artificial	Aa	NYDMN
599.	EpCAM_19130-C11_CC -	artificial	Aa	VISYDGSNKYYTDSVKG

	HCDR2			
600.	EpCAM_19130-C11_CC - HCDR3	artificial	Aa	RGATPVDY
601.	EpCAM_19130-C11_CC - LCDR1	artificial	Aa	RSSQSLLSNGYNYLG
602.	EpCAM_19130-C11_CC - LCDR2	artificial	Aa	FGSSRAS
603.	EpCAM_19130-C11_CC - LCDR3	artificial	Aa	MQALQTPFT
604.	EpCAM_19130-C11_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSNY DMNWVRQAPGKCLEWVAVISYDGSNKYYTD SVKGRFTISRDNKNTLYLQMNSLRTEDEVY YCAKRGATPVDYWGQGTLVTVSS
605.	EpCAM_19130-C11_CC - VL	artificial	Aa	EIVMTQSPLSLPVTPGEPASISCRSSQSLLSNG YNYLGWYLVKPGQSPQLLIYFGSSRASGVPDR FSGSGGTDFTLKISGVEAEDVGVYYCMQALQ TPFTFGCGTKVDIK
606.	EpCAM_19131-B6_CC - HCDR1	artificial	Aa	RYDMH
607.	EpCAM_19131-B6_CC - HCDR2	artificial	Aa	FISYDGSNEDYPAVKG
608.	EpCAM_19131-B6_CC - HCDR3	artificial	Aa	VGASPFDY
609.	EpCAM_19131-B6_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
610.	EpCAM_19131-B6_CC - LCDR2	artificial	Aa	EVSKRPS
611.	EpCAM_19131-B6_CC - LCDR3	artificial	Aa	TSYAGSNNLV
612.	EpCAM_19131-B6_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMHWVRQAPGKCLEWVAFISYDGSNEDYPA VKGRFTISRDNKNTLYLQMNSLRAEDTAVYY CAKVGASPFDYWGQGTLVTVSS
613.	EpCAM_19131-B6_CC - VL	artificial	Aa	QSALTQPASVSGSPGRSVTISCTGTSSDVGGYN YVSWYQQHPGKAPKLMIIYVSKRPSGVPVRF GSKSDNTASLTVSGLQAEDAEDYYCTSYAGSN NLVFGCGTKLTVL
614.	EpCAM_19131-H3_hu_N67Q_CC - HCDR1	artificial	Aa	NYDMN
615.	EpCAM_19131-H3_hu_N67Q_CC - HCDR2	artificial	Aa	VISYDGSQKSYSDSVKG
616.	EpCAM_19131-H3_hu_N67Q_CC - HCDR3	artificial	Aa	RGATPFDY
617.	EpCAM_19131-H3_hu_N67Q_CC - LCDR1	artificial	Aa	SGDKLGDKYAS
618.	EpCAM_19131-H3_hu_N67Q_CC - LCDR2	artificial	Aa	QDSRRPS
619.	EpCAM_19131-H3_hu_N67Q_CC	artificial	Aa	QVWDYSSDHWV

	- LCDR3			
620.	EpCAM_19131- H3_hu_N67Q_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSQKSYSDS VKGRFTISRDNKNTLSLQMNLSLRNEDSAVYY CAKRGATPFDYWGQGLVTVSS
621.	EpCAM_19131- H3_hu_N67Q_CC - VL	artificial	Aa	SYELTQPPSVSVSPGQTASITCSGDKLGDKYAS WYQQKPGQSPVLVIYQDSRRPSGIPERFSGSNS GNTATLTISGTQAMDEADYYCQVWDYSSDHW VFGCGTKLTVL
622.	EpCAM_19132- E12_hu_CC - HCDR1	artificial	Aa	NYDMN
623.	EpCAM_19132- E12_hu_CC - HCDR2	artificial	Aa	VISYDGS DKHYTDSVKG
624.	EpCAM_19132- E12_hu_CC - HCDR3	artificial	Aa	RGATPVDY
625.	EpCAM_19132- E12_hu_CC - LCDR1	artificial	Aa	SASSSISNSLH
626.	EpCAM_19132- E12_hu_CC - LCDR2	artificial	Aa	RTSNLAS
627.	EpCAM_19132- E12_hu_CC - LCDR3	artificial	Aa	QQGSSIPRT
628.	EpCAM_19132- E12_hu_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGS DKHYTD SVKGRFTISRDNKNTLFLQMNLSLRTEDTAVY YCAKRGATPVDYWGQGLVTVSS
629.	EpCAM_19132- E12_hu_CC - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVTITCSASSSISNSLH WYQQKPGKAPKLLIYRTSNLASGVPSRFSGSGS GTDFTLTISLQPEDFATYYCQQGSSIPRTFGCG TKLEIK
630.	EpCAM_19143- C11_CC - HCDR1	artificial	Aa	RYDMN
631.	EpCAM_19143- C11_CC - HCDR2	artificial	Aa	FISYDGSNEDY PDAVKG
632.	EpCAM_19143- C11_CC - HCDR3	artificial	Aa	VGASPFDY
633.	EpCAM_19143- C11_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
634.	EpCAM_19143- C11_CC - LCDR2	artificial	Aa	GASSRAT
635.	EpCAM_19143- C11_CC - LCDR3	artificial	Aa	QQYGSSPRT
636.	EpCAM_19143- C11_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMNWVRQAPGKCLEWVAFISYDGSNEDY PDA VKGRFTISRDNKNTLYLQLNSLRAEDTAVYY CAKVGASPFDYWGQGLVTVSS
637.	EpCAM_19143-	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL

	C11_CC - VL			AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPRTFG CGTKVEIK
638.	EpCAM_19143- E11_CC - HCDR1	artificial	Aa	NYDMN
639.	EpCAM_19143- E11_CC - HCDR2	artificial	Aa	VISYDGSNKYYTDSVKG
640.	EpCAM_19143- E11_CC - HCDR3	artificial	Aa	RGATPFDY
641.	EpCAM_19143- E11_CC - LCDR1	artificial	Aa	RASQSVNSNLA
642.	EpCAM_19143- E11_CC - LCDR2	artificial	Aa	GASTRAT
643.	EpCAM_19143- E11_CC - LCDR3	artificial	Aa	QQYNNWPYT
644.	EpCAM_19143- E11_CC - VH	artificial	Aa	QVQLVESGGGVVLPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSNKYYTD SVKGRFTISRDNRSRNTLYLQMNSLRTEDTAVY SCTKRGATPFDYWGQGTLVTVSS
645.	EpCAM_19143- E11_CC - VL	artificial	Aa	EIVLTQSPATLSVSPGERATLSCRASQSVNSNL AWYQQKPGQAPRLLIYGASTRATGIPARFSGS GSGTEFTLTISLQSEDFAVYYCQQYNNWPYTF GCGTKLEIK
646.	EpCAM_19145- C4_CC - HCDR1	artificial	Aa	NYDMN
647.	EpCAM_19145- C4_CC - HCDR2	artificial	Aa	VISYDGSNDKHYTDSVKG
648.	EpCAM_19145- C4_CC - HCDR3	artificial	Aa	RGATPVDY
649.	EpCAM_19145- C4_CC - LCDR1	artificial	Aa	RSSQSLLSHNGYNYLD
650.	EpCAM_19145- C4_CC - LCDR2	artificial	Aa	LGSNRAS
651.	EpCAM_19145- C4_CC - LCDR3	artificial	Aa	MQALQAPLT
652.	EpCAM_19145- C4_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSNDKHYTD SVKGRFTISRDNKNTLYLQMNSLRTEDTAVY YCAKRGATPVDYWGQGTLVTVSS
653.	EpCAM_19145- C4_CC - VL	artificial	Aa	EIVMTQTPLSLPVTTPGEPASISCRSSQSLLSHNG YNYLDWYLQKPGQSPQLLIYLGSNRASGVPDR FSGSGGTDFTLKISRVEAEDVGVYYCMQALQ APLTFGCGTKVDIK
654.	EpCAM_19145- F12_CC - HCDR1	artificial	Aa	RYDMN
655.	EpCAM_19145- F12_CC - HCDR2	artificial	Aa	FISYDGSNEDYPPDAVKG
656.	EpCAM_19145- F12_CC - HCDR3	artificial	Aa	VGASPFDY
657.	EpCAM_19145- F12_CC - LCDR1	artificial	Aa	RSSQSLLSHNGYNYLG
658.	EpCAM_19145- F12_CC - LCDR2	artificial	Aa	SGSSRAS
659.	EpCAM_19145- F12_CC - LCDR3	artificial	Aa	MQALQTPFT

660.	EpCAM_19145-F12_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY DMNWVVRQAPGKCLEWVAFISYDGSNEDYPPDA VKGRFTISRDNKNTLYLQMNSLRAEDTAVYY CAKVGASPFDYWGQGTLVTVSS
661.	EpCAM_19145-F12_CC - VL	artificial	Aa	EIVMTQSPLSLPVTGPGEASISCRSSQSLLSNG YNYLGWYLQKPGQSPQLLIYSGSSRASGVPDR FSGSGSGTDFTLKISRVEAEDVGVYYCMQALQ TPFTFGCGTKVEIK
662.	EpCAM_19168-H9_CC - HCDR1	artificial	Aa	RYYMH
663.	EpCAM_19168-H9_CC - HCDR2	artificial	Aa	VIWHDGSNKYYADSVKG
664.	EpCAM_19168-H9_CC - HCDR3	artificial	Aa	EAPSLAY
665.	EpCAM_19168-H9_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
666.	EpCAM_19168-H9_CC - LCDR2	artificial	Aa	GASSRAT
667.	EpCAM_19168-H9_CC - LCDR3	artificial	Aa	QQYGSSPLT
668.	EpCAM_19168-H9_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY YMHVVRQAPGKCPEWVAVIWHDGSNKYYAD SVKGRFTISRDNKNTLYLQMNSLRAEDTAVY YCAREAPSLAYWGQGTLVTVSS
669.	EpCAM_19168-H9_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPLTFG CGTKVEIK
670.	EpCAM_19171-A5_CC - HCDR1	artificial	Aa	RYYMH
671.	EpCAM_19171-A5_CC - HCDR2	artificial	Aa	VIWHDGSNKYYADSVKG
672.	EpCAM_19171-A5_CC - HCDR3	artificial	Aa	EAPSLAY
673.	EpCAM_19171-A5_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
674.	EpCAM_19171-A5_CC - LCDR2	artificial	Aa	GASSRAT
675.	EpCAM_19171-A5_CC - LCDR3	artificial	Aa	QQYGSSIT
676.	EpCAM_19171-A5_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY YMHVVRQAPGKCPEWVAVIWHDGSNKYYAD SVKGRFTISRDNKNTLYLQMNSLRAEDTAVY YCAREAPSLAYWGQGTLVTVSS
677.	EpCAM_19171-A5_CC - VL	artificial	Aa	EIVMTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSITFGCG TRLEIK
678.	EpCAM_19171-D3_CC - HCDR1	artificial	Aa	RYYMH
679.	EpCAM_19171-D3_CC - HCDR2	artificial	Aa	VIWHDGSNKYYADSVKG
680.	EpCAM_19171-D3_CC - HCDR3	artificial	Aa	EAPSLAY
681.	EpCAM_19171-	artificial	Aa	RASQSVSSSYLA

	D3_CC - LCDR1			
682.	EpCAM_19171-D3_CC - LCDR2	artificial	Aa	GASSRAT
683.	EpCAM_19171-D3_CC - LCDR3	artificial	Aa	QQYGSSPWT
684.	EpCAM_19171-D3_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSRY YMHWVRQAPGKCPPEWVAVIWHDGSNKYYAD SVKGRFTISRDN SKNTLYLQMNSLRAEDTAVY YCAREAPSLAYWGQGTLLTVSS
685.	EpCAM_19171-D3_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPWTFG CGTKVEIK
686.	EpCAM_19171-E11_CC - HCDR1	artificial	Aa	SYYS
687.	EpCAM_19171-E11_CC - HCDR2	artificial	Aa	RVYTSGSTDYNPSLKS
688.	EpCAM_19171-E11_CC - HCDR3	artificial	Aa	DSGNFWGFLDH
689.	EpCAM_19171-E11_CC - LCDR1	artificial	Aa	RSSQSLLSHNGYNYLD
690.	EpCAM_19171-E11_CC - LCDR2	artificial	Aa	LGSNRAS
691.	EpCAM_19171-E11_CC - LCDR3	artificial	Aa	MQALQTPWT
692.	EpCAM_19171-E11_CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSYY WSWIRQPAGKCLEWIGRVYTSGSTDYNPSLKS RVTMSLDTSKSKQFSLKLRVTAADTAVYYCAR DSGNFWGFLDHWGQGTLLTVSS
693.	EpCAM_19171-E11_CC - VL	artificial	Aa	EIVLTQSPLSLPVTPEPASISCRSSQSLLSHNGY NYLDWYLQKPGQSPQLLIYLGSNRASGVDRF SGSGSGTDFTLKISRVEAEDVGIYYCMQALQTP WTFGCGTKVEIK
694.	EpCAM_19180-B12_CC - HCDR1	artificial	Aa	NYDMN
695.	EpCAM_19180-B12_CC - HCDR2	artificial	Aa	VISYDGSNKYYTDSVKG
696.	EpCAM_19180-B12_CC - HCDR3	artificial	Aa	RGATPFDY
697.	EpCAM_19180-B12_CC - LCDR1	artificial	Aa	TGTNSDVGSYNLVS
698.	EpCAM_19180-B12_CC - LCDR2	artificial	Aa	DVSHRPS
699.	EpCAM_19180-B12_CC - LCDR3	artificial	Aa	SSYISSSLV
700.	EpCAM_19180-B12_CC - VH	artificial	Aa	EVQLLESGGGLVQPGGSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSNKYYTD SVKGRFTISRDN SRNTLYLQMNSLRTEAVY SCTKRGATPFDYWGQGTLLTVSS
701.	EpCAM_19180-B12_CC - VL	artificial	Aa	QSALTQPPSVSGSPQSITISCTGTNSDVGSYNL VSWYQQHPGKTPKLMYDVS HRPSGVS NRFSG SKSGNTASLTISGLQAEDEADYICSSYISSSLV

				FGCGTKLTVL
702.	EpCAM_19180-B6_N67Q_CC - HCDR1	artificial	Aa	NYDMN
703.	EpCAM_19180-B6_N67Q_CC - HCDR2	artificial	Aa	VISYDGSQKSYSDSVKG
704.	EpCAM_19180-B6_N67Q_CC - HCDR3	artificial	Aa	RGATPFDY
705.	EpCAM_19180-B6_N67Q_CC - LCDR1	artificial	Aa	GGNNIGSKNVH
706.	EpCAM_19180-B6_N67Q_CC - LCDR2	artificial	Aa	RDSKRPS
707.	EpCAM_19180-B6_N67Q_CC - LCDR3	artificial	Aa	QAWDRSTAV
708.	EpCAM_19180-B6_N67Q_CC - VH	artificial	Aa	EVQLLESGGGSAQPGGSLRLSCVASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSQKSYSDS VKGRFTISRDN SKNTLSLQMNSLRNEDTAVYY CAKRGATPFDYWGQGLVTVSS
709.	EpCAM_19180-B6_N67Q_CC - VL	artificial	Aa	SYELTQPPSVSVAPGQTARITCGGNNIGSKNVH WYQQKPGQAPVLVIYRDSKRPSGIPERFSGSNS GNTATLTISGTQAMDEADYYCQAWDRSTAVF GCGTKLTVL
710.	EpCAM_19180-D10S69Y_CC - HCDR1	artificial	Aa	NYDMN
711.	EpCAM_19180-D10S69Y_CC - HCDR2	artificial	Aa	VISYDGSNKYYSDSVKG
712.	EpCAM_19180-D10S69Y_CC - HCDR3	artificial	Aa	RGATPFDY
713.	EpCAM_19180-D10S69Y_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
714.	EpCAM_19180-D10S69Y_CC - LCDR2	artificial	Aa	DVSVRPS
715.	EpCAM_19180-D10S69Y_CC - LCDR3	artificial	Aa	SSYISSTTLV
716.	EpCAM_19180-D10S69Y_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSNKYYSDS VKGRFTISRDN SKNTLSLQMNSLRNEDTAVYY CAKRGATPFDYWGQGLVTVSS
717.	EpCAM_19180-D10S69Y_CC - VL	artificial	Aa	QSALTQPPSASGSPGQSITISCTGTSSDVGGYNY VSWYQQHPGKAPKLMYDVSVRPSGVSNRFSG SKSGNTASLTISGLQAEDEADYYCSSYISSTTLV FGCGTKLTVL
718.	EpCAM_19180-D10_N67Q_CC -	artificial	Aa	NYDMN

	HCDR1			
719.	EpCAM_19180-D10_N67Q_CC - HCDR2	artificial	Aa	VISYDGSQKSYSDSVKG
720.	EpCAM_19180-D10_N67Q_CC - HCDR3	artificial	Aa	RGATPFDY
721.	EpCAM_19180-D10_N67Q_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
722.	EpCAM_19180-D10_N67Q_CC - LCDR2	artificial	Aa	DVSVRPS
723.	EpCAM_19180-D10_N67Q_CC - LCDR3	artificial	Aa	SSYISSTTLV
724.	EpCAM_19180-D10_N67Q_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSQKSYSDS VKGRFTISRDNKNTLSLQMNSLRNEDTAVYY CAKRGATPFDYWGQGLVTVSS
725.	EpCAM_19180-D10_N67Q_CC - VL	artificial	Aa	QSALTQPPSASGSPGQSITISCTGTSSDVGGYNY VSWYQQHPGKAPKLMYDVSVRPSGVSNRFSG SKSGNTASLTISGLQAEDEADYCYSSYISSTTLV FGCGTKLTVL
726.	EpCAM_19180-G7_CC - HCDR1	artificial	Aa	GYVMH
727.	EpCAM_19180-G7_CC - HCDR2	artificial	Aa	WINPNSGGTNYAQKFQG
728.	EpCAM_19180-G7_CC - HCDR3	artificial	Aa	TGALAGALKH
729.	EpCAM_19180-G7_CC - LCDR1	artificial	Aa	RSSQSLLSHNGYNYLD
730.	EpCAM_19180-G7_CC - LCDR2	artificial	Aa	LGSNRAS
731.	EpCAM_19180-G7_CC - LCDR3	artificial	Aa	MQALQTPFT
732.	EpCAM_19180-G7_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKVCKASGYTFTG YYMHVWRQAPGQCLEWMGWINPNSGGTNYA QKFQGRITMTRDTSISTAYMELSLRSDDTAVY YCARTEGALAGALKHWGQGLVTVSS
733.	EpCAM_19180-G7_CC - VL	artificial	Aa	EIVMTQSPLSLPVTPEPAPASISCRSSQSLLSHNG YNYLDWYLQKPGQSPQLLIYLGSNRASGVPDR FSGSGGTDFTLKISRVEAEDVGVYYCMQALQ TPFTFGCGTKVEIK
734.	EpCAM_19182-H8_CC - HCDR1	artificial	Aa	NYDMN
735.	EpCAM_19182-H8_CC - HCDR2	artificial	Aa	VISYDGSQKHYTDSVKG
736.	EpCAM_19182-H8_CC - HCDR3	artificial	Aa	RGATPVDY
737.	EpCAM_19182-H8_CC - LCDR1	artificial	Aa	TGTNSDVGGYNYVS
738.	EpCAM_19182-H8_CC - LCDR2	artificial	Aa	DVSKRPS
739.	EpCAM_19182-	artificial	Aa	SSYISSSSLV

	H8_CC - LCDR3			
740.	EpCAM_19182-H8_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGS DKHYTD SVKGRFTISRDN SKNTLYLQMNSLRTE DTAVY YCAKRGATPVDYWGQGLTVTVSS
741.	EpCAM_19182-H8_CC - VL	artificial	Aa	QSALTQPASVSGSPGRSVTISCTGTNSDVGGYN YVSWYQQHPGKAPKLM IYDVSKRPSGVS NRFS GSKSGNTASLTISGLQAEDEADYYC SSYISSSSL VFGCGTKLTVL
742.	EpCAM_19187-B6_N67Q_CC - HCDR1	artificial	Aa	NYDMN
743.	EpCAM_19187-B6_N67Q_CC - HCDR2	artificial	Aa	VISYDGSQKSYSDSVKG
744.	EpCAM_19187-B6_N67Q_CC - HCDR3	artificial	Aa	RGATPFDY
745.	EpCAM_19187-B6_N67Q_CC - LCDR1	artificial	Aa	GGNNIGSKSVH
746.	EpCAM_19187-B6_N67Q_CC - LCDR2	artificial	Aa	QDSKRPS
747.	EpCAM_19187-B6_N67Q_CC - LCDR3	artificial	Aa	QAWDSSTAV
748.	EpCAM_19187-B6_N67Q_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFRNY DMNWVRQAPGKCLEWVAVISYDGSQKSYSDS VKGRFTISRDN SKNTLSLQMNSLRNEDTAVYY CAKRGATPFDYWGQGLTVTVSS
749.	EpCAM_19187-B6_N67Q_CC - VL	artificial	Aa	SYELTQPPSVSVAPGQTARITCGNNIGSKSVH WYQQKPGQSPVLVIYQDSKRPSGIPDRFSGSNS GNTATLTISGTQAMDEADYYCQAWDSSTAVF GCGTKLTVL
750.	MSLN_13203-C2_CC - HCDR1	artificial	Aa	SNSAAWN
751.	MSLN_13203-C2_CC - HCDR2	artificial	Aa	RTYYRSKWyNDYAVSVKS
752.	MSLN_13203-C2_CC - HCDR3	artificial	Aa	AIFVVPAAAMRFDY
753.	MSLN_13203-C2_CC - LCDR1	artificial	Aa	RSSQSLLSHNGYNYLD
754.	MSLN_13203-C2_CC - LCDR2	artificial	Aa	LGSNRAS
755.	MSLN_13203-C2_CC - LCDR3	artificial	Aa	MQALQTPT
756.	MSLN_13203-C2_CC - VH	artificial	Aa	QVQLQQSGPGLVKPSQTLSTCAISGDSVSSNS AAWNWIRQSPSRCLEWLGRTYYRSKWyNDY AVSVKSRITINPDISKNQFSLQLNSVTPEDTAVY YCARAIFVVPAAAMRFDYWGQGLTVTVSS
757.	MSLN_13203-C2_CC - VL	artificial	Aa	EIVMTQSPLSLPVTGPGEPAISCRSSQSLLSHNG YNYLDWY LQKPGQSPQLLIYLG SNRASGVPDR FSGSGGTDFTLKISRVEAEDVGVYYCMQALQ TPTFGCGTKVDIK

758.	MSLN_13203-F11_CC - HCDR1	artificial	Aa	SNYMS
759.	MSLN_13203-F11_CC - HCDR2	artificial	Aa	VIYSSGNTYYADSVKG
760.	MSLN_13203-F11_CC - HCDR3	artificial	Aa	GSYYAFDI
761.	MSLN_13203-F11_CC - LCDR1	artificial	Aa	GLSSGSVSTTYPS
762.	MSLN_13203-F11_CC - LCDR2	artificial	Aa	STNTRSS
763.	MSLN_13203-F11_CC - LCDR3	artificial	Aa	VLYMGSGIWV
764.	MSLN_13203-F11_CC - VH	artificial	Aa	EVQLVESGGGLIQPGGSLRLSCAVSGFTVSSNY MSWVRQAPGKCLEWVSVIYSSGNTYYADSVK GRFTISRDNKNTLYLQMNSLRAEDTAVYYCA SGSYYAFDIWGQGTMTVSS
765.	MSLN_13203-F11_CC - VL	artificial	Aa	QTVVTQEPSLTVSPGGTVTLTCGLSSGSVSTTY YPSWYQQTPGQAPRTLIYSTNTRSSGVPDRFSG SILGNKAALTITGAQADDESYYCVLYMGSGI WVFGCGTKLTVL
766.	MSLN_13204-A9_CC - HCDR1	artificial	Aa	NAWMS
767.	MSLN_13204-A9_CC - HCDR2	artificial	Aa	RIKTKTDGGTTDYAAPVKG
768.	MSLN_13204-A9_CC - HCDR3	artificial	Aa	DFRIMGATWFDP
769.	MSLN_13204-A9_CC - LCDR1	artificial	Aa	SGDKLGDKYAS
770.	MSLN_13204-A9_CC - LCDR2	artificial	Aa	QHSRRPS
771.	MSLN_13204-A9_CC - LCDR3	artificial	Aa	QAWDSSTVV
772.	MSLN_13204-A9_CC - VH	artificial	Aa	EVQLVESGGGLVKPGGSLRLSCAASGFTFSNA WMSWVRQAPGKCLEWVGRIKTKTDGGTTDY AAPVKGRFTISRDDSKNTLYLQMNSLKTEDTA VYYCTTDFRIMGATWFDPWGQGLTVTVSS
773.	MSLN_13204-A9_CC - VL	artificial	Aa	SYELTQPPSVSVSPGQTASITCSGDKLGDKYAS WYQQKPGQSPVLYIQHSRRPSGIPERFSGSNS GNTATLTISGTQAMDEADYYCQAWDSSTVVF GCGTKLTVL
774.	MSLN_13204-D11_CC - HCDR1	artificial	Aa	SYSTEMN
775.	MSLN_13204-D11_CC - HCDR2	artificial	Aa	SISSRSSYIHYADSVKG
776.	MSLN_13204-D11_CC - HCDR3	artificial	Aa	VQRAGLDY
777.	MSLN_13204-D11_CC - LCDR1	artificial	Aa	TGSSSDVGNYNLVS
778.	MSLN_13204-D11_CC - LCDR2	artificial	Aa	EVSNRPS
779.	MSLN_13204-D11_CC - LCDR3	artificial	Aa	SSYTSSSTWV

780.	MSLN_13204-D11_CC - VH	artificial	Aa	EVQLVESGGGLVKPGGSLRLSCAASGFTFSSYS MNWVRQAPGKCLEWVSSISSRSSYIHADSVK GRFTISRDNKNSLNLQMNSLRAEDTAVYYCA RVQRAGLDYWGQGTLVTVSS
781.	MSLN_13204-D11_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGSSSDVGNYNL VSWYQQHPGKAPKLMISEVSNRPSGVSDRFSG SKSGNTASLTISGLQAEDAEDYCYSSYTSSSTW VFGCGTKLTVL
782.	MSLN_13204-F11_CC - HCDR1	artificial	Aa	SSSYYWG
783.	MSLN_13204-F11_CC - HCDR2	artificial	Aa	SIYYSGSTNYNPSLKS
784.	MSLN_13204-F11_CC - HCDR3	artificial	Aa	PSNYDAFDI
785.	MSLN_13204-F11_CC - LCDR1	artificial	Aa	TGSSSNIGAGYDVH
786.	MSLN_13204-F11_CC - LCDR2	artificial	Aa	GNSNRPS
787.	MSLN_13204-F11_CC - LCDR3	artificial	Aa	QSYDSSLGGWV
788.	MSLN_13204-F11_CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSLSSSS YYWGWIRQPPGKCLEWIGSIYYSGSTNYNPSL KSRVTISADTSKNQFSLKLSSVTAADTAVYYC ARPSNYDAFDIWGQGMVTVSS
789.	MSLN_13204-F11_CC - VL	artificial	Aa	QSVLTQPPSVSGAPGQRVTISCTGSSSNIGAGY DVHWHYQQLPGTAPKLLIYGNSNRPSGVDPDRFS GSKSGTSASLAITGLQAEDAEDYCYQSYDSSLG GWVFGCGTKLTVL
790.	MSLN_13204-H6_CC - HCDR1	artificial	Aa	SGGFFWS
791.	MSLN_13204-H6_CC - HCDR2	artificial	Aa	YIYYSGSTYYNPSLRS
792.	MSLN_13204-H6_CC - HCDR3	artificial	Aa	DPGSYRVWFDP
793.	MSLN_13204-H6_CC - LCDR1	artificial	Aa	RASQNIKNYLN
794.	MSLN_13204-H6_CC - LCDR2	artificial	Aa	DASSLQS
795.	MSLN_13204-H6_CC - LCDR3	artificial	Aa	QQSYSTPFT
796.	MSLN_13204-H6_CC - VH	artificial	Aa	QVQLQESGPGLVKPSQTLSTCTVSGGSISSGG FFWSWIRQHPGKCLEWIGYIYYSGSTYYNPSLR SRVTISVDTSKNQFSLKLSSVTAADTAVYYCAR DPGSYRVWFDPWGQGTTLVTVSS
797.	MSLN_13204-H6_CC - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVTITCRASQNIKNYL NWXQQKPGRAPKLLIYDASSLQSGDPSRFSGS GSGTDFTLTISLQPEDFATYYCQQSYSTPFTFG CGTKVEIK
798.	MSLN_13213-A9_CC - HCDR1	artificial	Aa	DHYMS
799.	MSLN_13213-A9_CC - HCDR2	artificial	Aa	YISNSGSIYYVDSVKG
800.	MSLN_13213-A9_CC - HCDR3	artificial	Aa	DVRTAFDY
801.	MSLN_13213-	artificial	Aa	RASQSIGSWLA

	A9_CC - LCDR1			
802.	MSLN_13213-A9_CC - LCDR2	artificial	Aa	AASSLQS
803.	MSLN_13213-A9_CC - LCDR3	artificial	Aa	QQANSFPPT
804.	MSLN_13213-A9_CC - VH	artificial	Aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDH YMSWIRQAPGKCLEWISYISNSGSIYYVDSVK GRFTISRDNKNSLYLQMNSLRAEDTAVYYCA RDVRTAFDYWGQGTLLVTVSS
805.	MSLN_13213-A9_CC - VL	artificial	Aa	EIQMTQSPSSVSASVGDRVTITCRASQSIGSWL AWYQQKPGKAPNLLIYAASSLQSGVPSRFGSG GSGTDFTLTISSLQPEDFATYYCQQANSFPPTFG CGTKVEIK
806.	MSLN_13215-B12_CC - HCDR1	artificial	Aa	SSSYFWG
807.	MSLN_13215-B12_CC - HCDR2	artificial	Aa	NIYYSGSSNYNPSLKS
808.	MSLN_13215-B12_CC - HCDR3	artificial	Aa	LPRGDRDAFDI
809.	MSLN_13215-B12_CC - LCDR1	artificial	Aa	RASQGISNYLA
810.	MSLN_13215-B12_CC - LCDR2	artificial	Aa	AASTLQS
811.	MSLN_13215-B12_CC - LCDR3	artificial	Aa	QQSYSTPFT
812.	MSLN_13215-B12_CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSS
813.	MSLN_13215-B12_CC - VL	artificial	Aa	EIVMTQSPSSLSASVGDRVTITCRASQGISNYLA WYQQKPGKVPKLLIYAASTLQSGVPSRFGSGG SGTDFTLTISSLQPEDFATYYCQQSYSTPFTFGC GTKVEIK
814.	MSLN_13215-B12_LC_V3Q_C - HCDR1	artificial	Aa	SSSYFWG
815.	MSLN_13215-B12_LC_V3Q_C - HCDR2	artificial	Aa	NIYYSGSSNYNPSLKS
816.	MSLN_13215-B12_LC_V3Q_C - HCDR3	artificial	Aa	LPRGDRDAFDI
817.	MSLN_13215-B12_LC_V3Q_C - LCDR1	artificial	Aa	RASQGISNYLA
818.	MSLN_13215-B12_LC_V3Q_C - LCDR2	artificial	Aa	AASTLQS
819.	MSLN_13215-B12_LC_V3Q_C - LCDR3	artificial	Aa	QQSYSTPFT
820.	MSLN_13215-	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY

	B12_LC_V3Q_C C - VH			FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSS
821.	MSLN_13215- B12_LC_V3Q_C C - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVTITCRASQGISNYLA WYQQKPGKVPKLLIYAASLTQSGVPSRFSGSG SGTDFLTITSSLQPEDFATYYCQSYSTPFTFGC GTKVEIK
822.	MSLN_13216- B12_CC - HCDR1	artificial	Aa	SGGHFWS
823.	MSLN_13216- B12_CC - HCDR2	artificial	Aa	YIYYSGSTYSTPSLTS
824.	MSLN_13216- B12_CC - HCDR3	artificial	Aa	EGQSGSFDI
825.	MSLN_13216- B12_CC - LCDR1	artificial	Aa	TGTSSDVGGSDYVS
826.	MSLN_13216- B12_CC - LCDR2	artificial	Aa	EVSNRPS
827.	MSLN_13216- B12_CC - LCDR3	artificial	Aa	SSYTTTGTLV
828.	MSLN_13216- B12_CC - VH	artificial	Aa	QVQLQESGPGLVKPSQTLSTCTVSGGSISSGG HFWSWIRQHPPGKCLEWIGYIYYSGSTYSTPSLT SRVTMSRDTSKNQFSLKLSSVTAADTAVYYCA REGQSGSFDIWGQGTMTVTVSS
829.	MSLN_13216- B12_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSSDVGGSDY VSWYRQHPPGKAPKLIIEVSNRPSGVSNRFSGS KSGNTASLTISGLQAEDEADYYCSSYTTTGTLV FGCGTKLTVL
830.	MSLN_13216- B4_CC - HCDR1	artificial	Aa	SYGMH
831.	MSLN_13216- B4_CC - HCDR2	artificial	Aa	VIWKDGNNKYYADSVKG
832.	MSLN_13216- B4_CC - HCDR3	artificial	Aa	GLNYYYYGMDV
833.	MSLN_13216- B4_CC - LCDR1	artificial	Aa	TRSNNGGIANNYVQ
834.	MSLN_13216- B4_CC - LCDR2	artificial	Aa	ENNQRPS
835.	MSLN_13216- B4_CC - LCDR3	artificial	Aa	QSYDGS HHVV
836.	MSLN_13216- B4_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMHWVRQAPGKCLEWVAIVWKDGNNKYYA DSVKGRFTISRDN SKNTLYLQMNSLRAEDTAV YYCARGLNYYYYGMDVWGQGTITVTVSS
837.	MSLN_13216- B4_CC - VL	artificial	Aa	NFMLTQPHSVSESPGKTATISCTRSNNGIANNY VQWYQQRPGSSPTIVYIYENNQRPSGVPDRFSGS IDSSNSASLTISGLKTEDEADYYCQSYDGS HH VVFSGTKLTVL
838.	MSLN_13216- C1_CC - HCDR1	artificial	Aa	GYYIH
839.	MSLN_13216- C1_CC - HCDR2	artificial	Aa	WINPKSGGTHYAQKFQG
840.	MSLN_13216-	artificial	Aa	AEARLAARQEYYYYFYGMDV

	C1_CC - HCDR3			
841.	MSLN_13216-C1_CC - LCDR1	artificial	Aa	SGDKLGDKYAS
842.	MSLN_13216-C1_CC - LCDR2	artificial	Aa	QDSKRPS
843.	MSLN_13216-C1_CC - LCDR3	artificial	Aa	QAWDSSTVV
844.	MSLN_13216-C1_CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTG YYIHWVRQAPGQCLEWMGWINPKSGGTHYA QKFQGRVTMTRDTSISTAYMELSRLLRSDDTAV YYCARAEARLARQEYYFYGMVWGQGT VTVSS
845.	MSLN_13216-C1_CC - VL	artificial	Aa	SYELTQPASVSVSPGQTASITCSGDKLGDKYAS WYQQKPGQSPVLVIYQDSKRPSGIPERFSGSNS GNTATLTISGTQAMDEADYYCQAWDSSTVVF GCGTKLTVL
846.	MSLN_13229-C9_CC - HCDR1	artificial	Aa	SGAYFWS
847.	MSLN_13229-C9_CC - HCDR2	artificial	Aa	YIYYSGSTYTNPSLRD
848.	MSLN_13229-C9_CC - HCDR3	artificial	Aa	EGAGYVFDI
849.	MSLN_13229-C9_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
850.	MSLN_13229-C9_CC - LCDR2	artificial	Aa	EVSNRPS
851.	MSLN_13229-C9_CC - LCDR3	artificial	Aa	QVWDSSSDHVV
852.	MSLN_13229-C9_CC - VH	artificial	Aa	QVQLQESGPGLVKPSQTLSTCTVSGGSISSGA YFWSWIRQHHPGKCLEWIGYIYYSGSTYTNPSLR DRLKISVDTSKNQFSLKLSVTAADTAMYYCA REGAGYVFDIHWGQGTMTVTVSS
853.	MSLN_13229-C9_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSSDVGGYN YVSWYQQHPGKAPKLMIIYVSNRPSGVSNRFS GSKSGNTASLTISGLQAGDEADYFCQVWDSSS DHVVFVCGTKLTVL
854.	MSLN_13238-G11_CC - HCDR1	artificial	Aa	SGGYWN
855.	MSLN_13238-G11_CC - HCDR2	artificial	Aa	YIFYSGITYSNPSLKS
856.	MSLN_13238-G11_CC - HCDR3	artificial	Aa	GLVRGAPDAFDI
857.	MSLN_13238-G11_CC - LCDR1	artificial	Aa	QASQDISNYLN
858.	MSLN_13238-G11_CC - LCDR2	artificial	Aa	AASSLQG
859.	MSLN_13238-G11_CC - LCDR3	artificial	Aa	QQSYSTPFT
860.	MSLN_13238-G11_CC - VH	artificial	Aa	QVQLQESGPGLVKPSQTLSTCTVSGGSISSGG YYWNWIRQHHPGQCLEWIGYIFYSGITYSNPSLK SLFTISLDTSKNQFSLKLSVTAADTAVYYCAR GLVRGAPDAFDIHWGQGTMTVTVSS

861.	MSLN_13238-G11_CC - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVITTCQASQDISNYLN WYQLKPGKAPKLLIQAASSLQGGVPSRFSGSG SGTDFTLTISSLQPEDFATYYCQSYSTPFTFGC GTKVEIK
862.	MSLN_13239-D5_CC - HCDR1	artificial	Aa	SYYS
863.	MSLN_13239-D5_CC - HCDR2	artificial	Aa	RIYYNGNTYYNPSLKS
864.	MSLN_13239-D5_CC - HCDR3	artificial	Aa	PKLGIDAFDI
865.	MSLN_13239-D5_CC - LCDR1	artificial	Aa	TGSSSNIGAGYDVH
866.	MSLN_13239-D5_CC - LCDR2	artificial	Aa	GNSNRPS
867.	MSLN_13239-D5_CC - LCDR3	artificial	Aa	QSHDSSLSGSV
868.	MSLN_13239-D5_CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSYY WSWIRQPPGKCLEWIGRIYYNGNTYYNPSLKS RVTISGDTSKNQFSLKLSSVTAADTAVYYCARP KLGIDAFDIWGQGTMTVSS
869.	MSLN_13239-D5_CC - VL	artificial	Aa	QSVLTQPPSVSGAPGQRVTISCTGSSSNIGAGY DVHWWYQKLPGTAPKLLIYGNSNRPSGVPDRFS GSKSGTSASLAITGLQAEDEADYYCQSHDSSL GSVFGCGTKLTVL
870.	MSLN_13254-B10_CC - HCDR1	artificial	Aa	SGGYFWS
871.	MSLN_13254-B10_CC - HCDR2	artificial	Aa	YIYYSGSTYTNPSLRD
872.	MSLN_13254-B10_CC - HCDR3	artificial	Aa	EGAGYAFDI
873.	MSLN_13254-B10_CC - LCDR1	artificial	Aa	TGTSSDVGGYNYVS
874.	MSLN_13254-B10_CC - LCDR2	artificial	Aa	EVSNRPS
875.	MSLN_13254-B10_CC - LCDR3	artificial	Aa	SSYTSSSTLV
876.	MSLN_13254-B10_CC - VH	artificial	Aa	QVQLQESGGGLVKPSETLSLTCTVSGGSISSGG YFWSWIRQHPPGKCLEWIGYIYYSGSTYTNPSLR DRLKISVDTSKNQFSLKLSSVTAADTAMYYCA REGAGYAFDIWGQGTMTVSS
877.	MSLN_13254-B10_CC - VL	artificial	Aa	QSALTQPASVSGSPGQSITISCTGTSSDVGGYN YVSWYQQHPGKAPKLMIIYVSNRPSGVSNRFS GSKSGNTASLTISGLQAEDEADYYCSSYSSST LVFGCGTKLTVL
878.	MSLN_13256-H4_CC - HCDR1	artificial	Aa	SYGMH
879.	MSLN_13256-H4_CC - HCDR2	artificial	Aa	VISYDGSNKYYADSVKG
880.	MSLN_13256-H4_CC - HCDR3	artificial	Aa	EGAYFGSGSYYPYLYYYAMDV
881.	MSLN_13256-H4_CC - LCDR1	artificial	Aa	RASQSVSSSYLA

882.	MSLN_13256-H4_CC - LCDR2	artificial	Aa	GASIRAT
883.	MSLN_13256-H4_CC - LCDR3	artificial	Aa	QQYGSSLFT
884.	MSLN_13256-H4_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMHWVRQAPGKCLEWVAVISYDGSNKYYAD SVKGRFTISRDN SKNTLYLQMNSLRAEDTAVY YCAREGAYFGSGSYYP LYYYYAMDVWGQGT TVT VSS
885.	MSLN_13256-H4_CC - VL	artificial	Aa	EIVMTQSPGTL SLS PGERATL SCRASQSVSSSYL AWYQQKPGQAPRL LIYGASIRATGIPDRFSGSG SGTDFLTISRLEPEDFAVYYCQQYGSSLFTFG CGTRLEIK
886.	MSLN_13266-C1_CC - HCDR1	artificial	Aa	DHYMS
887.	MSLN_13266-C1_CC - HCDR2	artificial	Aa	YISSSGSTIYYVDSVKG
888.	MSLN_13266-C1_CC - HCDR3	artificial	Aa	DVRTAFDY
889.	MSLN_13266-C1_CC - LCDR1	artificial	Aa	RASQGISSWLA
890.	MSLN_13266-C1_CC - LCDR2	artificial	Aa	AASGLQS
891.	MSLN_13266-C1_CC - LCDR3	artificial	Aa	QQANSFPPT
892.	MSLN_13266-C1_CC - VH	artificial	Aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDH YMSWIRQTPGKCLEWVSYISSSGSTIYYVDSVK GRFTISRDN AKNSLYLQMNSLRAEDTAVYYCA RDVRTAFDYWGQGT LVT VSS
893.	MSLN_13266-C1_CC - VL	artificial	Aa	EIQMTQSPSSVSASVGDRVTITCRASQGISSWL AWYQQKPGKAPKLLIYAASGLQSGVPSRFSGS GSGTDFLT LTISSLPEDFATYYCQQANSFPPTFG CGTKVEIK
894.	MSLN_13268-A4_CC - HCDR1	artificial	Aa	SYTMS
895.	MSLN_13268-A4_CC - HCDR2	artificial	Aa	AISGSGGNTYYADSVKG
896.	MSLN_13268-A4_CC - HCDR3	artificial	Aa	VGRAALDY
897.	MSLN_13268-A4_CC - LCDR1	artificial	Aa	TGTSSDVGSYNLVS
898.	MSLN_13268-A4_CC - LCDR2	artificial	Aa	EVSKRPS
899.	MSLN_13268-A4_CC - LCDR3	artificial	Aa	SSYTSSTTV
900.	MSLN_13268-A4_CC - VH	artificial	Aa	EVQLLES GGGLVQPGGSPRLSCAVSGFTFSSYT MSWVRQAPGKCLEWVSAISGSGGNTYYADSV KGRSTISRDN SRNTLYLQMNSLRAEDTAVYYC AKVGRAALDYWGQGT LVT VSS
901.	MSLN_13268-A4_CC - VL	artificial	Aa	QSALTQPPSVSGSPGQSITISCTGTSSDVGSYNL VSWYQQHPGKAPKLM IYEVSKRPSGVS NRFSG SKSGNTASLTISGLQAEDEADYYCSSYTSSTV VFGCGTKLTVL
902.	MSLN_13269-A6_CC - HCDR1	artificial	Aa	SYAMS

903.	MSLN_13269-A6_CC - HCDR2	artificial	Aa	AISGSGGSTYYADSVKG
904.	MSLN_13269-A6_CC - HCDR3	artificial	Aa	EGYYDSSGYPLYYYFGMDV
905.	MSLN_13269-A6_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
906.	MSLN_13269-A6_CC - LCDR2	artificial	Aa	GASSRAT
907.	MSLN_13269-A6_CC - LCDR3	artificial	Aa	QRYGSSPIFT
908.	MSLN_13269-A6_CC - VH	artificial	Aa	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGGSTYYADSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AREGYDSSGYPLYYYFGMDVWGQGTITVTS S
909.	MSLN_13269-A6_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGS SGTDFTLISRLEPEDFAVYYCQRYGSSPIFTFG CGTKVEIK
910.	MSLN_13270-A3_CC - HCDR1	artificial	Aa	NAWMS
911.	MSLN_13270-A3_CC - HCDR2	artificial	Aa	RIKTKTDGGTTDYAAPVKG
912.	MSLN_13270-A3_CC - HCDR3	artificial	Aa	DFRIMGATWFDP
913.	MSLN_13270-A3_CC - LCDR1	artificial	Aa	SGSSSNIGSYSVN
914.	MSLN_13270-A3_CC - LCDR2	artificial	Aa	SNNQRPS
915.	MSLN_13270-A3_CC - LCDR3	artificial	Aa	AAWDDSLSGRVA
916.	MSLN_13270-A3_CC - VH	artificial	Aa	EVQLVESGGGLVKPGGSLRLSCAASGFTFSNA WMSWVRQAPGKCLEWVGRIKTKTDGGTTDY AAPVKGRFTISRDDSKNTLYLQMNSLKTEDTA VYYCTTDFRIMGATWFDPWGQGLTVTVSS
917.	MSLN_13270-A3_CC - VL	artificial	Aa	QSVLTQPSASGTPGQRVTISCSGSSSNIGSYSV NWKYQQLPGTAPKLLIYSNNQRPSGVPDRFSGS KSGTSASLAISGLRSEDEADYYCAAWDDSLSG RGVAFGCGTKLTVL
918.	MSLN_13317-C7_CC - HCDR1	artificial	Aa	DYYMS
919.	MSLN_13317-C7_CC - HCDR2	artificial	Aa	YISSSGSMIYYIDSVKG
920.	MSLN_13317-C7_CC - HCDR3	artificial	Aa	DLGPSFDY
921.	MSLN_13317-C7_CC - LCDR1	artificial	Aa	RASQGIGSWLA
922.	MSLN_13317-C7_CC - LCDR2	artificial	Aa	GASGLQS
923.	MSLN_13317-C7_CC - LCDR3	artificial	Aa	QQANSFPRT
924.	MSLN_13317-C7_CC - VH	artificial	Aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDY YMSWIRQAPGKCLEWISYISSSGSMIYYIDSVK GRFTISRDNKNSLYLQMNSLRAEDTAVYYCA RDLGPSFDYWGQGLTVTVSS

925.	MSLN_13317- C7_CC - VL	artificial	Aa	EIQMTQSPSSVAATVGDRVTITCRASQGIGSWL AWYQQKPGKAPKLLIYGASGLQSGVPSRFSGS GSGTDFLTLISSLQPEDFATYYCQQANSFPRTF GCGTKVEIK
926.	MSLN_13317- F9_CC - HCDR1	artificial	Aa	DHYMS
927.	MSLN_13317- F9_CC - HCDR2	artificial	Aa	YISNSGSTIYYADSVKG
928.	MSLN_13317- F9_CC - HCDR3	artificial	Aa	DQRNAFDI
929.	MSLN_13317- F9_CC - LCDR1	artificial	Aa	RASQGIGSWLA
930.	MSLN_13317- F9_CC - LCDR2	artificial	Aa	AASGLQS
931.	MSLN_13317- F9_CC - LCDR3	artificial	Aa	QSYSNPLT
932.	MSLN_13317- F9_CC - VH	artificial	Aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDH YMSWIRQAPGKCLEWISYISNSGSTIYYADSVK GRFTISRDNAKNSLYLQMNSLRAEDTAVYYCA RDQRNAFDIWGQGMVTVSS
933.	MSLN_13317- F9_CC - VL	artificial	Aa	AIQMTQSPSSLSASVGDRVTITCRASQGIGSWL AWYQQKPGKAPKLLIYAASGLQSGVPSRFSGS GSGTDFLTLISSLQPEDFATYYCQQSYSNPLTFG CGTKVEIK
934.	MSLN_13318- B9_CC - HCDR1	artificial	Aa	SSSYYWG
935.	MSLN_13318- B9_CC - HCDR2	artificial	Aa	SIYSGTTRYNP SLRS
936.	MSLN_13318- B9_CC - HCDR3	artificial	Aa	PGAGHDGFDI
937.	MSLN_13318- B9_CC - LCDR1	artificial	Aa	SGSSSNIGSNYVY
938.	MSLN_13318- B9_CC - LCDR2	artificial	Aa	DNNKRPS
939.	MSLN_13318- B9_CC - LCDR3	artificial	Aa	AAWDDSLSGWV
940.	MSLN_13318- B9_CC - VH	artificial	Aa	QVQLQESGPGLLKPSSETLSLTCTVSGGSISSSSY YWGWIRQPPGKCLEWIGSIYSGTTRYNP SLRS RVTTSLDASKNRLSLQLSSVTAADTAVYYCAR PGAGHDGFDIWGQGMVTVSS
941.	MSLN_13318- B9_CC - VL	artificial	Aa	QSVLTQPPSASGTPGQRVTISCSGSSSNIGSNYV YWYQQLPGTAPKLLIYDNNKRPSGIPDRFSGSK SGTSASLAISGLRSEDEADYYCAA WDDSLSGW VFGCGTKLTVL
942.	MSLN_13319- B8_CC - HCDR1	artificial	Aa	SYYS
943.	MSLN_13319- B8_CC - HCDR2	artificial	Aa	RIYSSGSANYNP SLKS
944.	MSLN_13319- B8_CC - HCDR3	artificial	Aa	EGQWRVPAQYYYFGMDV
945.	MSLN_13319- B8_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
946.	MSLN_13319- B8_CC - LCDR2	artificial	Aa	GASSRAT
947.	MSLN_13319-	artificial	Aa	QQYGSSIT

	B8_CC - LCDR3			
948.	MSLN_13319-B8_CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSYY WSWIRQPAGKCLEWIGRIYSSGSANYNPSLKS VTMSVDTSKNQFSLKLNSVTAADTAVYYCAR EGQWRVPAQYYYFGMDVWGQGTITVTVSS
949.	MSLN_13319-B8_CC - VL	artificial	Aa	EIVMTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSITFGCG TRLEIK
950.	MSLN_18025-E3_CC - HCDR1	artificial	Aa	SSSYFWV
951.	MSLN_18025-E3_CC - HCDR2	artificial	Aa	SIYYSGSTYYNPSLKS
952.	MSLN_18025-E3_CC - HCDR3	artificial	Aa	LPRGDRMTFDI
953.	MSLN_18025-E3_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
954.	MSLN_18025-E3_CC - LCDR2	artificial	Aa	GASSRAT
955.	MSLN_18025-E3_CC - LCDR3	artificial	Aa	QQYGSSPFT
956.	MSLN_18025-E3_CC - VH	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWWIRQPPGKCLEWIGSIYYSGSTYYNPSLKS RVTISVDTSKNQFSLKLNSVTAADTAVYYCAR LPRGDRMTFDIWGQGTMTVTVSS
957.	MSLN_18025-E3_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPFTFGC GTKLEIK
958.	MSLN_18026-C1_CC - HCDR1	artificial	Aa	SYGMH
959.	MSLN_18026-C1_CC - HCDR2	artificial	Aa	VIWNRYSNKYYADAVKG
960.	MSLN_18026-C1_CC - HCDR3	artificial	Aa	DVPYYYGMDV
961.	MSLN_18026-C1_CC - LCDR1	artificial	Aa	TRSSGSIGDNYVQ
962.	MSLN_18026-C1_CC - LCDR2	artificial	Aa	ENNQRPS
963.	MSLN_18026-C1_CC - LCDR3	artificial	Aa	QSYHGSNVV
964.	MSLN_18026-C1_CC - VH	artificial	Aa	QVQLVESGGGVVLPGRSLRLSCAASGFPFSSYG MHWVRQAPGKCLEWVAVIWNRYSNKYYADA VKGRFTISRDN SKNTLYLQMNSLRAEDTAVYY CARDVPIYYGMDVWGQGTITVTVSS
965.	MSLN_18026-C1_CC - VL	artificial	Aa	NFMLTQPHSVSESPGKTVIISCTRSSGSIGDNYV QWYQQRPGSSPTTVIYENNQRPSGVPDRFSGSI DSSSNSASLTISGLKTEDEADYYCQSYHGSNVV FGCGTKLTVL
966.	MSLN_18035-B6_CC - HCDR1	artificial	Aa	SYGMH
967.	MSLN_18035-B6_CC - HCDR2	artificial	Aa	VIWNDASNKYYADAVKG
968.	MSLN_18035-B6_CC - HCDR3	artificial	Aa	DVPYYYGMDV

969.	MSLN_18035-B6_CC - LCDR1	artificial	Aa	TRSSGSIGDNYVQ
970.	MSLN_18035-B6_CC - LCDR2	artificial	Aa	ENNQRPS
971.	MSLN_18035-B6_CC - LCDR3	artificial	Aa	QSYQQSNVV
972.	MSLN_18035-B6_CC - VH	artificial	Aa	QVQLVESGGGVVLPGRSLRLSCAASGFPFSSYG MHWVRQAPGKCLEWVAVIWNDAENKYYADA VKGRFTISRDNKNTLYLQMNSLRAEDTAVYY CARDVPYYYGMDVWGQGTITVTVSS
973.	MSLN_18035-B6_CC - VL	artificial	Aa	NFMLTQPHSVSESPGKTVIISCTRSSGSIGDNYV QWYQQRPGSSPTTVIYENNRPSGVPDRFSGSI DSSSNSASLTISGLKTEDEADYYCQSYQQSNVV FGCGTKLTVL
974.	MSLN_18036-C10_CC - HCDR1	artificial	Aa	SYAMS
975.	MSLN_18036-C10_CC - HCDR2	artificial	Aa	AISGSGEFSYAAAVKG
976.	MSLN_18036-C10_CC - HCDR3	artificial	Aa	VRNYYGSGSLDY
977.	MSLN_18036-C10_CC - LCDR1	artificial	Aa	RASQSVSSTYLA
978.	MSLN_18036-C10_CC - LCDR2	artificial	Aa	GASIRAT
979.	MSLN_18036-C10_CC - LCDR3	artificial	Aa	QQYGSSLT
980.	MSLN_18036-C10_CC - VH	artificial	Aa	EVQLLESGGGVVQPGRSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGEFSYAAAV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKVRNYYGSGSLDYWGQGTITVTVSS
981.	MSLN_18036-C10_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSTYL AWYQQKPGQAPRLLIYGASIRATGIPDRFSGSG SGTDFLTISRLEPEDFAVYYCQQYGSSLTFCG GTKVEIK
982.	MSLN_18036-C5_CC - HCDR1	artificial	Aa	SYAMS
983.	MSLN_18036-C5_CC - HCDR2	artificial	Aa	AISGSGEQWYYAPSVKG
984.	MSLN_18036-C5_CC - HCDR3	artificial	Aa	VRNYYGSGSLDY
985.	MSLN_18036-C5_CC - LCDR1	artificial	Aa	RASQSFSSAYLA
986.	MSLN_18036-C5_CC - LCDR2	artificial	Aa	GASIRAT
987.	MSLN_18036-C5_CC - LCDR3	artificial	Aa	QQYGSSLT
988.	MSLN_18036-C5_CC - VH	artificial	Aa	EVQLLESGGGVVQPGRSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGEQWYYAPSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKVRNYYGSGSLDYWGQGTITVTVSS
989.	MSLN_18036-C5_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSFSSAYL AWYQQKPGQAPRLLIYGASIRATGIPDRFSGSG

				SGTDFTLTISRLEPEDFAVYYCQQYGSSLTFGC GTKVEIK
990.	MSLN_18037- B3_CC - HCDR1	artificial	Aa	SYAMS
991.	MSLN_18037- B3_CC - HCDR2	artificial	Aa	AISGSGGSTYYAPSVKG
992.	MSLN_18037- B3_CC - HCDR3	artificial	Aa	EGYYPVSGYPLYYYYFGMDV
993.	MSLN_18037- B3_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
994.	MSLN_18037- B3_CC - LCDR2	artificial	Aa	GASSRAT
995.	MSLN_18037- B3_CC - LCDR3	artificial	Aa	QQYGSSPIFT
996.	MSLN_18037- B3_CC - VH	artificial	Aa	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGGSTYYAPSV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKEGYYPVSGYPLYYYYFGMDVWGQTTVTVS S
997.	MSLN_18037- B3_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPIFTFG CGTKVEIK
998.	MSLN_18037- G4_CC - HCDR1	artificial	Aa	SYAMS
999.	MSLN_18037- G4_CC - HCDR2	artificial	Aa	AISGSGGSTYYAGNVKG
1000.	MSLN_18037- G4_CC - HCDR3	artificial	Aa	EGYYPTSGYPLYYYYFGMDV
1001.	MSLN_18037- G4_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
1002.	MSLN_18037- G4_CC - LCDR2	artificial	Aa	GASSRAT
1003.	MSLN_18037- G4_CC - LCDR3	artificial	Aa	QQYGSSPIFT
1004.	MSLN_18037- G4_CC - VH	artificial	Aa	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGGSTYYAGNV KGRFTISRDNKNTLYLQMNSLRAEDTAVYYC AKEGYYPVSGYPLYYYYFGMDVWGQTTVTVS S
1005.	MSLN_18037- G4_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRLLIYGASSRATGIPDRFSGSG SGTDFTLTISRLEPEDFAVYYCQQYGSSPIFTFG CGTKVEIK
1006.	MSLN_18126- H2_CC - HCDR1	artificial	Aa	SYGMH
1007.	MSLN_18126- H2_CC - HCDR2	artificial	Aa	VIGSRESNKNYAESVKG
1008.	MSLN_18126- H2_CC - HCDR3	artificial	Aa	ALRIAVAASYYYYGLDV
1009.	MSLN_18126- H2_CC - LCDR1	artificial	Aa	RASQSVRSFLN
1010.	MSLN_18126- H2_CC - LCDR2	artificial	Aa	TASSLQS
1011.	MSLN_18126-	artificial	Aa	QQSYEMPIT

	H2_CC - LCDR3			
1012.	MSLN_18126- H2_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMHWVRQAPGKCLEWVAVIGSRESNKNYAES VKGRFTISRDN SKNTLYLQMNSLRAEDTAVYY CASALRIAVAASYYYYGLDVWGQGTITVTVSS
1013.	MSLN_18126- H2_CC - VL	artificial	Aa	EIQMTQSPSSLSASVGDRVTITCRASQSVRSFLN WYQQKPGKAPKLLIFTASSLQSGVPSRFSGSGS GTDFTLTISLQPEDFATYYCQQS YEMPITFGC GTRLEIK
1014.	MSLN_18183- C2_CC - HCDR1	artificial	Aa	SYGMG
1015.	MSLN_18183- C2_CC - HCDR2	artificial	Aa	VISYEASNKYYAEAVKG
1016.	MSLN_18183- C2_CC - HCDR3	artificial	Aa	EGAHFGSGSYYP LYYYYAMDV
1017.	MSLN_18183- C2_CC - LCDR1	artificial	Aa	RASQSVSSSYLA
1018.	MSLN_18183- C2_CC - LCDR2	artificial	Aa	GASIRAT
1019.	MSLN_18183- C2_CC - LCDR3	artificial	Aa	QQYGSSPIFT
1020.	MSLN_18183- C2_CC - VH	artificial	Aa	QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMGWVRQAPGKCLEWVAVISYEASNKYYAE AVKGRFTISRDN SKNTLYLQMNSLRAEDTAVY YCAREGAHFGSGSYYP LYYYYAMDVWGQGT TVT VSS
1021.	MSLN_18183- C2_CC - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRASQSVSSSYL AWYQQKPGQAPRL LIYGASIRATGIPDRFSGSG SGTDFLTISRLEPEDFAVYYCQQYGSSPIFTFG CGTKVEIK
1022.	MSLN_18201- G11_CC - HCDR1	artificial	Aa	DYYMT
1023.	MSLN_18201- G11_CC - HCDR2	artificial	Aa	YISSSGSTIYYAEAVKG
1024.	MSLN_18201- G11_CC - HCDR3	artificial	Aa	DRNSHFDY
1025.	MSLN_18201- G11_CC - LCDR1	artificial	Aa	RASQGIRTWLA
1026.	MSLN_18201- G11_CC - LCDR2	artificial	Aa	GASGLQS
1027.	MSLN_18201- G11_CC - LCDR3	artificial	Aa	QQAESFPRT
1028.	MSLN_18201- G11_CC - VH	artificial	Aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDY YMTWIRQAPGKCLEWLSYISSSGSTIYYAEAV KGRFTISRDN AKNSLFLQMNSLRAEDTAVYYC ARDRNSHFDYWGQGT LTVTVSS
1029.	MSLN_18201- G11_CC - VL	artificial	Aa	EIQMTQSPSSVSASVGDRVTITCRASQGIRTWL AWYQQKPGKAPKLLIYGASGLQSGVPSRFSGSGS GSGTDFLTISLQPEDFATYYCQQAESFPRTFG CGTKVEIK
1030.	MSLN_MS_R4L_ CC - HCDR1	artificial	Aa	GYYYH

1031.	MSLN_MS_R4L_ CC - HCDR2	artificial	Aa	WINPNSGGTNYAQKFQG
1032.	MSLN_MS_R4L_ CC - HCDR3	artificial	Aa	VEAVAGREYYYYFSGMDV
1033.	MSLN_MS_R4L_ CC - LCDR1	artificial	Aa	SGEKLGDKYVY
1034.	MSLN_MS_R4L_ CC - LCDR2	artificial	Aa	QSTKRPS
1035.	MSLN_MS_R4L_ CC - LCDR3	artificial	Aa	QAYHASTAV
1036.	MSLN_MS_R4L_ CC - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKV SCKASGYTFTG YYIHWVRQAPGQCLEWMGWINPNSGGTNYA QKFQGRVTMTRDTSISTAYMELSR LRSDDTAV YYCARVEAVAGREYYYYFSGMDV VWGQGTVT VSS
1037.	MSLN_MS_R4L_ CC - VL	artificial	Aa	SYELTQPPSVSVSPGQTASITCS GEKLGDKYVY WYQQKPGQSPVLVIYQSTKRPSG VPERFSGSNS GNTATLTISGTQAMDEADYYCQAY HASTAVF GCGTKLTVL
1038.	MSLN_R195L_C C - HCDR1	artificial	Aa	SYAMS
1039.	MSLN_R195L_C C - HCDR2	artificial	Aa	AISGSGEFSYAAAVKG
1040.	MSLN_R195L_C C - HCDR3	artificial	Aa	VRNYYGSGSLDY
1041.	MSLN_R195L_C C - LCDR1	artificial	Aa	RASQSVSSTYLA
1042.	MSLN_R195L_C C - LCDR2	artificial	Aa	GASIRAT
1043.	MSLN_R195L_C C - LCDR3	artificial	Aa	QQYQSSLT
1044.	MSLN_R195L_C C - VH	artificial	Aa	EVQLLESGGGVVQPGRSLRLS CAASGFTFSSYA MSWVRQAPGKCLEWVSAISGSGE FSYAAAV KGRFTISRDNKNTLYLQMNSLRA EDTAVYYC AKVRNYYGSGSLDYWGQGLTVSS
1045.	MSLN_R195L_C C - VL	artificial	Aa	EIVLTQSPGTLSPGERATLSCRA SQSVSTYL AWYQQKPGQAPRLLIYGASIRAT GIPDRFSGSG SGTDFTLISRLEPEDFAVYYCQQY QSSLTFCG GTKVEIK
1046.	scFv_anti_CDH19 _2G6.007 CC - HCDR1	artificial	Aa	SYGMH
1047.	scFv_anti_CDH19 _2G6.007 CC - HCDR2	artificial	Aa	FIWYEGSNKYAESVKD
1048.	scFv_anti_CDH19 _2G6.007 CC - HCDR3	artificial	Aa	RAGIIGTIGYYYGMDV
1049.	scFv_anti_CDH19 _2G6.007 CC - LCDR1	artificial	Aa	SGDRLGEKYTS
1050.	scFv_anti_CDH19 _2G6.007 CC - LCDR2	artificial	Aa	QDTKRPS
1051.	scFv_anti_CDH19	artificial	Aa	QAWESSTVV

	_2G6.007 CC- LCDR3			
1052.	CDH19_2G6.007 CC- VH	artificial	Aa	QVQLVESGGGVVQPGGSLRLSCAASGFTFSSY GMHWVRQAPGKCLEWVAFIWIYEGSNKYIAE SVKDRFTISRDNKNTLYLQMNSLRAEDTAVY YCARRAGIIGTIGYIYGMVWVGQGTITVTVSS
1053.	CDH19_2G6.007 CC- VL	artificial	Aa	SYELTQPPSVSVSPGQTASITCSGDRLEGEKYTS WYQQRPGQSPLLVIYQDTRKPSGIPERFSGSNS GNTATLTISGTQAMDEADYYCQAWESSTVVF GCGTKLTVLS
1054.	scFv_anti_FOLR1 _C145209_(2A3) - HCDR1	artificial	Aa	SNSVIWN
1055.	scFv_anti_FOLR1 _C145209_(2A3) - HCDR2	artificial	Aa	RTYYRSKWYNDYAVSVKS
1056.	scFv_anti_FOLR1 _C145209_(2A3) - HCDR3	artificial	Aa	TVYYYGMDV
1057.	scFv_anti_FOLR1 _C145209_(2A3) - LCDR1	artificial	Aa	SGDKLGNNYAA
1058.	scFv_anti_FOLR1 _C145209_(2A3) - LCDR2	artificial	Aa	QDSKRPS
1059.	scFv_anti_FOLR1 _C145209_(2A3) - LCDR3	artificial	Aa	QSWDSSTVV
1060.	FOLR1_C145209 _(2A3) - VH	artificial	Aa	QVQLQQSGPGLVKPSQTLTLTCAISGDSVSSNS VIWNWIRQSPSRGLEWLGRTYYRSKWYNDYA VSVKSRITINPDTSKNQFSLQLNSVTPEDTAVY YCAGTVYYYGMVWVGQGTITVTVSS
1061.	FOLR1_C145209 _(2A3) - VL	artificial	Aa	SYELTQPPSVSVSPGQTGSITCSGDKLGNNYAA WYQQKPGQSPVLVIYQDSKRPSGIPERFSGSNS GNTATLTISGTQAVDEADYYCQSWDSSTVVF GGTKLTVLGS
1062.	scFv_anti_MSLN _C147862_(6F12) - HCDR1	artificial	Aa	SGANYWT
1063.	scFv_anti_MSLN _C147862_(6F12) - HCDR2	artificial	Aa	YIYYSGSTYLNPSLRG
1064.	scFv_anti_MSLN _C147862_(6F12) - HCDR3	artificial	Aa	ESGSSYGFDY
1065.	scFv_anti_MSLN _C147862_(6F12) - LCDR1	artificial	Aa	RTSQSITSYLN
1066.	scFv_anti_MSLN _C147862_(6F12) - LCDR2	artificial	Aa	ASSSLQS
1067.	scFv_anti_MSLN _C147862_(6F12) - LCDR3	artificial	Aa	QQSYSGPFT
1068.	MSLN_C147862_	artificial	Aa	QVQLQESGPGLVKPSQTLTLTCTVSGGSISSGA

	(6F12) - VH			NYWTWIRQHPGKGLEWIGYIYYSGSTYLNPSL RGRVTMSVDTSKNQFSLKLSSVTAADTAVYYC ARESGSSYGFDYWQGTLVTVSS
1069.	MSLN_C147862_ (6F12) - VL	artificial	Aa	DIQMTQSPSSLSASVGDRTITCRTSQSITSYLN WYQQKPGQAPKLLIYASSSLQSGVPSRFSGSGS GTDFTLTISLQPEDFATYYCQSYSQPFTFGPG TKVDIKRS
1070.	CH3 15-E11 CC x I2Lopt x G4 x scFc SEFL2 clipopt x G4 x MS 15-B12 CC x I2L_GQ - Nucleotide Sequence	artificial	na	CAGGTTCAGTTGGTTCAGTCTGGCGCCGAAG TGAAGAAACCAGGCGCTTCTGTGAAGGTGTC CTGCAAGGCCTCTGGCTACACCTTACCAAC TACTGGATGAACTGGGTCCGACAGGCTCCTG GCCAGTGTCTGGAATGGATGGGCAATATCGC TTACGGCGTGAAGGGCACCAACTACAACCAG AAATTCCAGGGCAGAGTGACCATGACCGTGG ACACCTCTCCTCCACCGCCTACATGGAAC GTCCCGGCTGAGATCTGACGACACCGCCGTG TACTACTGCGCCACCAGATACTTCTACGTGA TGGACTATTGGGGCCAGGGCACCCCTGGTTAC AGTTTCTTCTGGCGGCGGAGGACAAGGCGGT GGTGGTCAAGGCGGAGGCGGACAGGATATC CAGATGACCCAGTCTCCTTCCAGCCTGTCTG CCTCTGTGGGCGACAGAGTGACAATCACCTG TCGGGCCTCTCAGGACATCTCCAACCTACCTG AACTGGTATCAGCAGAAACCCGGCAAGGTG CCCAAGCTGCTGATCTACTACACCTCCAGAC TGCACCTCCGGCGTGCCCTCTAGATTTTCTGGC TCTGGATCTGGCACCGACTTCACCCTGACCA TCAGTTCTCTGCAGCCTGAGGACGTGGCCAC CTACTACTGTGTGCAGTACGCCAGTTTCTCCT TGACCTTCGGCTGTGGCACCAAGGTGGAAAT CAAAGCGGTGGCGGAGGCCAAGAGGTGCA GCTTGTTGAATCTGGCGGAGGATTGGTGCAG CCTGGCGGATCTCTGAAGCTGTCTTGTGCCG CCTCCGGCTTACCTTCAACAAATACGCCAT GAATTGGGTTTCGACAAGCCCCAGGCAAAGG CATGGAATGGGTCCGCCGATCAGATCCAAG TACAACAACACTACGCTACCTACTACGCCGACG CCGTGAAGGACAGATTCACCATCTCTCGGGA CGACTCCAAGAACACCCTGTACCTGCAGATG AACAACTGAAAACCGAGGATAACCGCCGTCT ATTACTGTGTCAGAGCCGGCAACTTCGGCTC CTCCTACATCTCCTACTTTGCCTACTGGGGAC AGGGAACCCTCGTGACTGTTTCTAGCGGTGG TGCGGACAAGGTGGCGGTGGACAAGGCGG CGGAGGCCAACAAACAGTGGTCACACAAGA GCCAGCCTGACAGTGTCTCCTGGCGGAACA GTGACCATCACATGTGGATCTTCTACCGGCG CTGTGACCTCCGGCAACTACCCCAATTGGAT CCAGAAGAAGCCCGGCCAGGCTCCTAGAGG ACTGATCGGCGGAACAAAGTTTCTGGCCCCT GGCACACCAGCCAGATTCTCAGGATCTCTGG AAGGCGGCAAGGCCGCTCTGACATTGTCTGG CGTTCAGCCAGAGGATGAGGCCGAGTACTAT TGCGTGCTGTACTACTCCAACAGATGGGTGT TCGGCTCCGGCACAAAGCTGACAGTTCTCGG AGGTGGCGGATGCCCTCCTTGTCTGCTCCT

				GAATTGCTCGGCGGACCCTCCGTGTTCCCTGTT TCCTCCAAAGCCTAAGGACACCCTGATGATC TCTCGGACCCCTGAAGTGACCTGCGTGGTGG TGGATGTGTCCCACGAGGAACCAGAAGTGA AGTTCAATTGGTACGTGGACGGCGTGGAAGT GCACAACGCTAAGACCAAGCCTTGCGAGGA ACAGTACGGCAGCACCTACAGATGTGTGTCC GTGCTGACCGTGCTGCACCAGGACTGGCTGA ATGGCAAAGAGTACAAGTGCAAGGTGTCCA ACAAGGCACTGCCCGTCTCTATCGAAAAGAC CATCTCCAAGGCTAAGGGCCAGCCTCGGGAA CCTCAGGTTTACACCCTGCCTCCATCTCGGG AAGAGATGACCAAGAACCAGGTGTCCCTGA CCTGCCTGGTCAAGGGCTTCTACCCTTCCGAT ATCGCCGTGGAATGGGAGTCCAATGGCCAGC CTGAGAACAACACTACAAGACCACACCTCCTGT GCTGGACTCCGACGGCTCATTCTTCTGTACT CCAAGCTGACTGTGGACAAGTCTCGGTGGCA GCAGGGCAACGTGTTCTCCTGTTCTGTGATG CACGAGGCCCTGCACAACCACTACACCAGA AGTCCCTGTCTCTGAGCCCTGGCAAAGGTGG TGGCGGTCAAGGCGGTGGTGGCCAAGGCGG CGGAGGACAAGGTGGCGGAGGCCAAGGTGG TGGCGGACAAGGCGGAGGTGGTCAATGTCT CCTTGTCCAGCACCAAGAACTCCTCGGAGGCC CTTCTGTGTTTCTGTTCCACCTAAGCCAAAG GATACACTCATGATCAGCAGGACTCCCAGAG TGACATGTGTGTCGTGGACGTTTCCCATGA AGAACCCGAAGTCAAGTTTAATTGGTATGTC GATGGCGTTCGAGGTCCACAATGCCAAGACA AAGCCCTGTGAAGAACAATACGGGTCCACCT ATAGATGCGTCAGCGTCTGACAGTCTGCA TCAGGATTGGCTCAACGGGAAAGAATACAA ATGTAAGTCTCTAACAAGGCTCTCCCAGCA CCAATCGAGAAAACCATAGCAAGGCCAAA GGACAGCCCCGCGAGCCACAAGTGTATACCC TGCCACCTAGCCGCGAGGAAATGACAAAGA ATCAAGTCTCTCTGACCTGTCTCGTGAAGGG GTTTTACCCAGCGACATTGCCGTCGAGTGG GAGTCTAACGGACAACCCGAAAACAATTATA AGACAACCCACCTGTCCTGGACAGCGACGG CTCATTTTTTCTCTACTCTAACTCACCGTGG ATAAGTCCAGATGGCAACAGGGAAATGTGTT CAGCTGCAGCGTGATGCATGAAGCTCTCCAC AATCATTATACCAGAAAAGCCTGAGCTTGT CTCCCGCAAAGGTGGCGGAGGACAGGTTC AGTTGCAAGAGTCTGGACCTGGCCTCGTGAA GCCTTCTGAGACACTGAGCCTGACCTGTACC GTGTCTGGCGGCTCCATCTCCTCCAGCTCTTA CTTCTGGGGCTGGATCAGACAGCCTCCAGGC AAGTGCCTCGAGTGGATCGGCAACATCTACT ACTCCGGCTCCAGCAACTACAATCCTAGCCT GAAGTCCCGGTGACAATCTCTGTGGATAACC TCTAAGAACCAGTTTAGCCTCAAGCTGTCCA GCGTGACCGCCGCTGATACCGCTGTGTATTA TTGCGCTAGACTGCCAGAGGCGACCGGGAT
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				<p>GCTTTCGATATTTGGGGACAAGGCACAATGG TCACCGTTTCTAGCGGAGGCGGTGGCCAAGG TGGCGGAGGCCAAGGCGGCGGTGGTCAAGA TATTGTGATGACACAGAGCCCCTCTAGCCTG AGCGCTTCCGTGGGAGATCGCGTGACCATTA CCTGTAGAGCCAGCCAGGGCATCAGCAATTA CCTGGCCTGGTATCAACAAAAGCCTGGGAAA GTCCCTAAGCTCCTCATCTACGCCGCTTCCAC ACTGCAGAGCGGCGTGCCAAGCAGATTCACT GGATCCGGCAGCGGAACCGACTTTACTCTGA CTATCTCCAGCCTGCAGCCAGAAGATTTTCGC TACCTATTACTGCCAGCAGTCTACAGCACC CCTTTCACCTTTGGCTGCGGAACTAAGGTCG AGATCAAGAGCGGAGGTGGTGGACAAGAGG TCCAGTTGGTCGAGTCAGGTGGCGGCTTGGT CCAACCAGGTGGAAGCCTGAAACTGAGCTGC GCCGTTCTGGGTTTACTTTTAAACAAATATGC TATGAACTGGGTTCCGCCAGGCACCTGGAAAA GGCATGGAATGGGTTGCCAGAATCCGCAGCA AGTATAACAATTATGCCACCTATTATGCCGA TGCTGTCAAGGATCGGTTACCATCAGCAGG GACGATAGCAAGAATACCCTCTATCTCCAAA TGAACAATCTCAAGACAGAGGACACAGCAG TGTACTATTGTGTTTCGCGCTGGCAACTTTGGC AGCAGCTACATCAGCTACTTCGCTTACTGGG GCCAAGGGACACTTGTGACCGTTAGCAGCGG AGGCGGAGGACAAGGTGGCGGAGGACAAGG CGGAGGTGGACAGCAGACAGTTGTGACCCA AGAGCCTTCTCTGACTGTGTCACCAGGCGGC ACCGTGACAATTACATGCGGAAGTTCCACAG GCGCCGTGACCAGCGGCAATTATCCTAACTG GATTCAGAAAAACCTGGACAGGCCCAAG AGGCCTGATTGGAGGCACCAAATTTCTCGCT CCCGGCACTCCTGCTCGGTTCTCTGGTAGTCT TGAAGGCGGAAAAGCTGCCCTGACTCTCTCT GGCGTGCAACCCGAAGATGAAGCTGAATATT ACTGCGTCTCTACTATAGCAATCGCTGGGT TTTCGGAAGCGGCACCAAGCTCACTGTCCTC TGA</p>
1071.	<p>CH3 15- E11_1_VAG_CC x I2L x G4 x scFc x G4 x MS 15- B12 CC x I2L clipopt_DI, AMG305 - Nucleotide Sequence</p>	artificial	na	<p>CAGGTGCAGCTGGTTCAGTCTGGCGCCGAAG TGAAGAAACCTGGCGCCTCTGTGAAGGTGTC CTGCAAGGCTTCTGGCTACACCTTTACCAAC TACTGGATGAACTGGGTCCGACAGGCCCTG GCCAGTGTGGAGTGGATGGGCAATATCGC TTACGGCGTGGCCGGCACCAACTACAACCAG AAATTCCAGGGCAGAGTGACAATGACCGTG GACACCTCCTCCTCCACCGCCTACATGGAAC TGTCCCGGCTGAGATCTGACGACACCGCCGT GTACTACTGCGCCACCAGATACTTCTACGTG ATGGACTACTGGGGCCAGGGCACCCCTGGTTA CAGTTTCTTCTGGCGGCGGAGGACAAGGCGG AGGTGGTCAAGGTGGTGGCGGACAGGATAT CCAGATGACCCAGTCTCCTTCCAGCCTGTCT GCCTCTGTGGGCGACAGAGTGACCATCACCT GTAGAGCCAGCCAGGACATCTCCAACCTACCT GAACTGGTATCAGCAGAAACCCGGCAAGGT</p>

				GCCCAAGCTGCTGATCTACTACACCTCTCGG CTGCACTCTGGCGTGCCCTCTAGATTTTCTGG CTCCGGCTCTGGCACCGACTTTACCTTGACA ATCTCCAGCCTGCAGCCTGAGGATGTGGCCA CCTACTACTGTGTGCAGTACGCCAGTTTCCT CTGACCTTCGGCTGTGGACCAAGGTGGAAA TCAAGTCTGGAGGCGGAGGCCAAGAGGTGC AGCTGGTGGAGTCCGGCGGCGGCCTGGTGCA GCCCGGCGGCTCCCTGAAGCTGTCCTGCGCC GCCTCCGGCTTCACCTTCAACAAGTACGCCA TGAACTGGGTGAGGCAGGCCCCGGCAAGG GCATGGAGTGGGTGGCCAGGATCAGGTCCA AGTACAACAACACTACGCCACCTACTACGCCGA CGCCGTGAAGGACAGGTTACCCATCTCCAGG GACGACTCCAAGAACACCCTGTACCTGCAGA TGAACAACCTGAAGACCGAGGACACCGCCG TGTAATACTGCGTGAGGGCCGGCAACTTCGG CTCCTCCTACATCTCCTACTTCGCCTACTGGG GCCAGGGCACCCCTGGTGACCGTGTCTCCGG CGGCGGCGGCCAAGGCGGCGGCGGCCAAGG CGGCGGCGGCCAACAGACCGTGGTGACCCA GGAGCCCTCCCTGACCGTGTCCCCGGCGGC ACCGTGACCATCACCTGCGGCTCCTCCACCG GCGCCGTGACCTCCGGCAACTACCCCAACTG GATCCAGAAGAAGCCCGGCCAGGCCCCAG GGGCCTGATCGGCGGCACCAAGTTCCTGGCC CCCGGCACCCCGCCAGGTTCTCCGGTCCC TGGAGGGCGGCAAGGCCGCCCTGACCCTGTC CGGCGTGCAGCCCGAGGACGAGGCCGAGTA CTACTGCGTGCTGTACTACTCCAACAGGTGG GTGTTCCGGCTCCGGCACCAAGCTGACCGTCC TAGGCGGAGGCGGCTGCCCTCCTTGTCTGC TCCTGAATTGCTCGGCGGACCCTCCGTGTCC TGTTTCCTCCAAAGCCTAAGGACACCCTGAT GATCTCTCGTACGCCTGAAGTGACCTGCGTG GTGGTGGATGTGTCCCACGAGGAACCCGAAG TGAAGTTCAATTGGTACGTGGACGGCGTGGA AGTGACAACGCCAAGACAAGCCCTGCGA GGAACAGTACGGCTCCACCTACAGATGCGTG TCCGTGCTGACAGTGCTGCACCAGGATTGGC TGAACGGCAAAGAGTACAAGTGCAAGGTGT CCAACAAGGCCCTGCCTGCTCCTATCGAAAA GACCATCTCCAAGGCCAAGGGCCAGCCTAGA GAGCCCCAGGTTTACACCCTGCCTCCAAGCA GAGAAGAGATGACCAAGAACCAGGTGTCCC TGACCTGCCTGGTCAAGGGCTTCTACCCTTCC GATATCGCCGTGGAATGGGAGAGCAATGGA CAGCCCGAGAACAACACTACAAGACCACACCTC CTGTGCTGGACTCCGACGGCTCATTCTTCTG TACTCCAAGCTGACCGTGGACAAGTCCAGAT GGCAGCAGGGCAACGTGTTCTCCTGCTCCGT GATGCACGAGGCCCTGCACAATCACTACACC CAGAAGTCCCTGTCTGTCCCCTGGAAAAG GAGGCGGAGGACAAGGCGGAGGTGGTCAAG GTGGTGGTGGCCAAGGCGGAGGCGGACAAG GCGGCGGAGGACAAGGTGGCGGTGGACAGT
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				GTCCTCCATGTCCAGCACCTGAGCTTCTCGG AGGCCCTTCTGTGTTTCTGTTCCACCTAAGC CAAAGGATACACTCATGATCAGCCGCACACC TGAAGTCACATGTGTCGTCGTGGATGTCTCT CATGAAGAACCAGAAGTCAAGTTTAATTGGT ATGTCGATGGCGTCGAGGTCCACAATGCTAA GACCAAGCCTTGTGAAGAACAATATGGCAGC ACCTATCGCTGTGTGTCTGTCTGACCGTCT GCATCAAGACTGGCTCAATGGGAAAGAATA CAAATGCAAAGTCTCTAACAAAGCTCTGCC GCACCAATCGAGAAAACCATCAGCAAGGCT AAAGGACAGCCTCGCGAGCCTCAAGTGATA CCCTGCCACCTTCTCGCGAGGAAATGACAAA AAATCAAGTCTCCCTCACCTGTCTCGTGAAG GGATTCTATCCCAGCGACATTGCCGTGAGT GGGAGTCTAATGGCCAGCCTGAAAACAATTA TAAGACAACCCACCTGTCCTGGACAGCGAC GGCTCATTTTTTCTCTACTCTAAACTCACCGT GGATAAGAGCCGGTGGCAACAGGGAAATGT GTTCAGCTGTAGCGTGATGCATGAAGCTCTC CACAACCATTATACACAGAAGAGTCTGAGCC TGTCTCCTGGCAAAGGCGGCGGAGGACAGGT GCAACTCCAGGAATCCGGGCCAGGGTTGGTG AAACCCAGCGAGACACTGTCTCTGACTTGCA CTGTTTCTGGTGGCTCCATTTCTCTAGCTCT TACTTCTGGGGTTGGATACGGCAACCACCTG GGAAGTGTCTCGAATGGATTGGTAACATCTA CTATAGTGGATCCTCCAATAACAATCCCAGC CTGAAGAGTCGTGTGACTATCAGCGTTGACA CCTCAAAGAATCAGTTCTCCCTTAAGCTGAG TTCCGTGACAGCAGCAGATACAGCCGTCTAC TACTGTGCTCGACTTCTAGGGGAGATCGGG ATGCCTTCGACATTTGGGGTCAGGGTACGAT GGTAACAGTGTCTAGTGGAGGCGGAGGTCA AGGCGGCGGAGGCCAAGGAGGAGGCGGACA AGATATCGTGATGACCCAGAGCCCATCAAGC CTGAGTGCTAGCGTTGGGGACAGGGTCACTA TCACTTGCAGAGCCTCACAGGGGATTTCAA CTATCTGGCCTGGTATCAGCAGAAACCTGGC AAGGTCCCCAAACTCCTGATATATGCTGCAA GCACGCTGCAAAGCGGGGTACCCTCTCGCTT TTCTGGGTCTGGCTCTGGCACAGACTTTACCC TGACCATCTCCAGTTTGCAGCCTGAGGACTT TGCCACCTACTATTGCCAGCAGTCCTACTCA ACACCCTTACCTTTGGCTGTGGCACCAAGG TGAGATCAAATCCGGAGGCGGAGGACAAG AAGTCCAGCTGGTTGAAAGTGGTGGCGGATT GGTTCAGCCAGGCGGCTCTCTGAAGCTGTCT TGTGCTGCCTCCGGCTTCACCTTCAACAAAT ACGCCATGAATTGGGTTTCGACAAGCCCCAGG CAAAGGCATGGAATGGGTCGCCCGGATCAG ATCCAAGTACAACAACACTACGCTACCTACTAC GCCGACGCCGTGAAGGACCGGTTACCATCT CCAGAGATGACTCCAAGAACACCCTGTACCT GCAGATGAACAACCTCAAGACCGAGGATAC CGCCGTCTATTACTGTGTCAGAGCCGGCAAC
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				<p>TTCGGCTCCTCCTACATCTCCTACTTCGCCTA CTGGGGCCAGGGAACCCTTGTGACAGTCTCT AGTGGCGGTGGTGGTCAAGGTGGTGGCGGCC AAGGCGGTGGCGGACAACAAACAGTGGTCA CCCAAGAGCCTAGCCTGACCGTTTCTCCTGG CGGCACCGTGACCATCACATGCGGATCTTCT ACCGGCGCTGTGACCTCCGGCAACTACCCCA ATTGGATCCAGAAGAAGCCAGGCCAGGCTCC TAGAGGACTGATCGGCGGCACAAAGTTTCTG GCTCCCGGCACTCCCGCCAGATTTTCTGGAT CTCTGGAAGGCGGCAAGGCTGCTCTGACATT GTCTGGCGTCCAGCCAGAGGATGAGGCCGA GTAATAATTGCGTGCTGTAATACTCCAACAGA TGGGTGTTCCGGCTCCGGCACCAAGCTGACAG TCCTATGA</p>
1072.	<p>MS 83-C2 CC x I2L x scFc x EP 71-A5 CC x I2L (G4S)3 - COMBI#11 (A8P) - Nucleotide Sequence</p>	artificial	na	<p>CAGGTGCAGCTGGTGAATCTGGTGGCGGAG TTGTGCAGCCTGGCAGATCCCTGAGACTGTC TTGTGCCGCTCCGGCTTACCTTCTCCTCTT ATGGAATGGGCTGGGTCCGACAGGCCCTGG CAAATGTTTGAATGGGTCCCGTGATCTCC TACGAGGCCTCCAACAAGTACTACGCCGAGG CCGTGAAGGGCAGATTCACCATCTCCAGAGA CAACTCCAAGAACCCTGTACCTGCAGATG AACTCCCTGAGAGCCGAGGACACCGCCGTGT ACTACTGTGCTAGAGAGGGCGCCATTTCGG CTCCGGCTCTTACTACCTCTGTAATACTACT ACGCTATGGACGTGTGGGGCCAGGGCACCAC AGTGACAGTTTCTAGCGGAGGCGGAGGAAG TGGCGGCGGAGGATCTGGCGGTGGTGGTTCT GAAATCGTGCTGACCCAGTCTCCTGGCACAC TGTCTTTGAGCCCTGGCGAGAGAGCTACCCT GAGCTGTAGAGCCTCTCAGTCCGTGTCTCC TCTTACCTGGCCTGGTATCAGCAGAAGCCCG GCCAGGCTCCTAGACTGTTGATCTACGGCGC CTCCATCAGAGCCACAGGCATCCCTGATAGA TTCTCCGGCAGCGGCTCTGGCACCGACTTCA CCCTGACAATCTCTCGGCTGGAACCCGAGGA CTTTGCTGTGTAATAATTGCCAGCAGTACGGC AGTCCCCTATCTTACCTTTGGCTGCGGCAC CAAGGTGGAATCAAGTCCGGGGGCGGAGG CTCCGAGGTGCAGCTGGTGGAGTCCGGCGGC GGCCTGGTGCAGCCCGGCGGCTCCCTGAAGC TGTCTTGCGCCGCTCCGGCTTACCTTCAAC AAGTACGCCATGAACTGGGTGAGGCAGGCC CCCGGCAAGGGCATGGAGTGGGTGGCCAGG ATCAGGTCCAAGTACAACAATACTACGCCACCT ACTACGCCGACGCCGTGAAGGACAGGTTAC CATCTCCAGGGACGACTCCAAGAACACCCTG TACCTGCAGATGAACAACCTGAAGACCGAG GACACCGCCGTGTACTACTGCGTGAGGGCCG GCAACTTCGGCTCCTCCTACATCTCCTACTTC GCCTACTGGGGCCAGGGCACCCCTGGTGACCG TGTCTCCGGCGGCGGCGGCTCCGGCGGCGG CGGCTCCGGCGGCGGCGGCTCCAGACCGTG GTGACCCAGGAGCCCTCCCTGACCGTGTCC CCGGCGGCACCGTGACCATCACCTGCGGCTC</p>

				CTCCACCGGCGCCGTGACCTCCGGCAACTAC CCCAACTGGATCCAGAAGAAGCCCGGCCAG GCCCCAGGGGCCTGATCGGCGGCACCAAGT TCCTGGCCCCCGGCACCCCGCCAGGTTCTC CGGCTCCCTGGAGGGCGGCAAGGCCGCCCTG ACCCTGTCCGGCGTGCAGCCCGAGGACGAGG CCGAGTACTACTGCGTGCTGTACTACTCCAA CAGGTGGGTGTTTCGGCTCCGGCACCAAGCTG ACCGTGCTAGGCGGCGGAGGATCTGGCGGA GGTGAAGCGGAGGCGGTGGATCTGACAAG ACCCACACATGTCCTCCATGTCCC GCCCTG AACTGCTAGGCGGACCTAGCGTGTTCTGTT CCCCCAAAGCCCAAGGACACCCTGATGATC AGCCGTACGCCGAAGTGACCTGCGTGGTGG TGGATGTGTCCCACGAGGACCCTGAAGTGAA GTTCAATTGGTACGTGGACGGCGTGGAAGTG CACAACGCCAAGACCAAGCCCTGCGAGGAA CAGTACGGCAGCACCTACAGATGCGTGTCCG TGCTGACCGTGCTGCATCAGGACTGGCTGAA CGGCAAAGAGTACAAGTGCAAGGTGTCCAA CAAGGCCCTGCCTGCCCCCATCGAGAAAACC ATCAGCAAGGCCAAGGGCCAGCCCCGCGAG CCTCAAGTGTATACCCTGCCCCCTAGCCGGG AAGAGATGACCAAGAACCAGGTGTCCCTGA CCTGTCTCGTGAAGGGCTTCTACCCCTCCGAT ATCGCCGTGGAATGGGAGAGCAACGGCCAG CCCGAGAACA ACTACAAGACCACCCCCCTG TGCTGGACAGCGACGGCTCATTCTTCTGTA CTCCAAACTGACCGTGGACAAGAGCCGGTGG CAGCAGGGCAACGTGTT CAGCTGCAGCGTGA TGCACGAGGCCCTGCACAACCACTACACCCA GAAGTCCCTGTCCCTGTCTCCCGGAAAGGC GGCGGAGGATCTGGCGGAGGCGGATCTGGG GGCGGAGGAAGTGGGGGAGGGGGAAGCGGA GGGGGAGGCTCAGGGGGGGGAGGATCCGAT AAGACCCACACCTGTCCCCTTGCCCTGCC CTGAACTGCTGGGAGGCCCTAGCGTGTTCT GTTCCCCCAAAGCCCAAGGACACCCTGATG ATCAGCCGGACCCCGAAGTGACCTGCGTGG TGGTGGATGTGTCCCACGAGGACCCTGAAGT GAAGTTCAATTGGTACGTGGACGGCGTGGA GTGCACAACGCCAAGACCAAGCCCTGCGAG GAACAGTACGGCAGCACCTACAGATGCGTGT CCGTGCTGACCGTGCTGCACCAGGACTGGCT GAACGGCAAAGAGTACAAGTGCAAGGTGTC CAACAAGGCCCTGCCTGCCCCCATCGAGAAA ACCATCAGCAAGGCCAAGGGCCAGCCCCGC GAGCCTCAAGTGTATACCCTGCCCC CAGCC GGGAAGAGATGACCAAGAACCAGGTGTCC TGACCTGTCTCGTGAAGGGCTTCTACCCCTCC GATATCGCCGTGGAATGGGAGAGCAACGGC CAGCCCGAGAACA ACTACAAGACCACCCCC CTGTGCTGGACAGCGACGGCTCATTCTTCT GTACTCCAAGCTGACAGTGGACAAGTCTAGA TGGCAGCAGGGCAACGTGTT CAGCTGCAGCG TGATGCACGAGGCCCTGCACAACCACTACAC
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				<p>CCAGAAGTCCCTGTCCCTGAGCCCCGGCAA GGTGGAGGCGGATCTGGCGGTGGCGGGAGT GGAGGAGGAGGCAGCCAGGTGCAGCTGATG GAATCTGGTGGCGGAGTTGTGCAGCCTGGCA GATCCCTGAGACTGTCTTGTGCCGCCTCCGG CTTCACCTTCAGCCGTAATATATGCACTGG GTCCGACAGGCCCTGGCAAGTGTCTGAAT GGGTTGCCGTGATCTGGCACGACGGCTCCAA CAAGTACTACGCCGACTCCGTGAAGGGCAGA TTCACCATCTCTCGGGACAACCTCCAAGAACA CCCTGTACCTGCAGATGAACTCCCTGAGAGC CGAGGACACCGCGTGTACTACTGTGCTAGA GAGGCCCTTCTCTGGCCTATTGGGGACAGG GAACACTGGTCACAGTGTCTCTGGCGGCGG AGGATCTGGCGGAGGTGGTAGCGGAGGCGG TGGATCTGAGATCGTGATGACCCAGTCTCCT GGCACACTGTCTCTGAGCCCTGGCGAGAGAG CTACCCTGTCTTGTAGAGCCTCTCAGTCCGTG TCCTCCTCCTACCTGGCTTGGTATCAGCAGA AGCCAGGCCAGGCTCCTCGGCTGTTGATCTA CGGCGCTTCTCTAGAGCCACAGGCATCCCT GACAGATTCTCCGGCTCTGGCTCTGGCACCG ACTTCACCCTGACCATCTCCAGACTGGAACC CGAGGACTTTGCTGTGTACTATTGCCAGCAG TACGGCTCCTCCATCACCTTCGGCTGTGGCA CCAGGCTGGAAATCAAGTCTGGAGGCGGAG GATCTGAAGTCCAGCTGGTTGAAAGTGGTGG CGGATTGGTTCAGCCAGGCGGCTCTCTGAAG CTGTCTTGTGCTGCCTCCGGCTTCACCTCAA CAAATACGCCATGAATTGGGTTGACAAGCC CCAGGCAAAGGCATGGAATGGGTGCGCCCGG ATCAGATCCAAGTACAACAACCTACGCTACCT ACTACGCCGACGCCGTGAAGGACCGGTTAC CATCTCCAGAGATGACTCCAAGAACACCCTG TACCTGCAGATGAACAACCTCAAGACCGAGG ATACCGCCGTCTATTACTGTGTCAGAGCCGG CAACTTCGGCTCCTCCTACATCTCCTACTTCG CCTACTGGGGCCAGGGAACCCTTGTGACAGT CTCTAGTGGCGGTGGTGGTAGTGGTGGTGGC GGCTCAGGCGGTGGCGGATCTCAAACAGTGG TCACCCAAGAGCCTAGCCTGACCGTTTCTCC TGGCGGCACCGTGACCATCACATGCGGATCT TCTACCGGCGCTGTGACCTCCGGCAACTACC CCAATTGGATCCAGAAGAAGCCAGGCCAGG CTCCTAGAGGACTGATCGGCGGCACAAAGTT TCTGGCTCCCGGCACTCCCGCCAGATTTTCTG GATCTCTGGAAGGCGGCAAGGCTGCTCTGAC ATTGTCTGGCGTCCAGCCAGAGGATGAGGCC GAGTACTATTGCGTGCTGTACTACTCCAACA GATGGGTGTTTCGGCTCCGGCACCAAGCTGAC AGTCCTATGA</p>
1073.	MS 83-C2 CC x I2L x scFc x EP 71-A5 CC x I2L (G4S)3 - COMBI#11 (A8P)	artificial	Aa	<p>QVQLVESGGGVVQPGRSLRLSCAASGFTFSSY GMGWVRQAPGKCLEWVAVISYEASNKYAE AVKGRFTISRDNKNTLYLQMNSLRAEDTAVY YCAREGAHFGSGSYYPYYAMDVWGQGT TVTSSGGGGSGGGGSGGGGSEIVLTQSPGTL</p>

				<p>LSPGERATLSCRASQSVSSSYLAWYQQKPGQA PRLLIYGASIRATGIPDRFSGSGSGTDFTLTISR EPEDFAVYYCQQYGSSPIFTFGCGTKVEIKSGG GGSEVQLVESGGGLVQPGGSLKLSAASGFTF NKYAMNWVRQAPGKGMWVARIRSKYNNYA TYYADAVKDRFTISRDDSKNTLYLQMNNLKTE DTAVYYCVRAGNFGSSYISYFAYWGQGLVT VSSGGGGSGGGSGGGGSQT VVTQEPSLTVSP GGTVTITCGSSTGAVTSGNYPNWIQKPGQAP RGLIGGTKFLAPGTPARFSGSLEGGKAALTLG VQPEDEAEYYCVLYYSNRWVFGSGTKLTVL GGGSGGGSGGGGSDKTHTCPPCPAPPELLGGP SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDP EVKFNWYVDGVEVHNAKTKPCEEQYGSTYRC VSVLTVLHQDWLNGKEYKCKVSNKALPAPIEK TISKAKGQPREPQVYTLPPSREEMTKNQVSLT LVKGFYPSDIAVEWESNGQPENNYKTTPPVL SDGSFFLYSKLTVDKSRWQQGNVFCFSVMHEA LHNHYTQKSLSLSPGKGGGGSGGGSGGGG GGGSGGGSGGGGSDKTHTCPPCPAPPELLGG PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHED PEVKFNWYVDGVEVHNAKTKPCEEQYGSTYR CVSVLTVLHQDWLNGKEYKCKVSNKALPAPIE KTISKAKGQPREPQVYTLPPSREEMTKNQVSLT CLVKGFYPSDIAVEWESNGQPENNYKTTPPVL DSDGSFFLYSKLTVDKSRWQQGNVFCFSVMHE ALHNHYTQKSLSLSPGKGGGGSGGGSGGGG SQVQLMESGGGVVQPGRSLRLS AASGFTFSR YYMHWVRQAPGKCPWVA VIWHDGSNKYYA DSVKGRFTISRDN SKNTLYLQMN SLRAEDTAV YYCAREAPSLAYWGQGLVT VSSGGGGSGGG GSGGGGSEIVMTQSPGTL SPSGERATLSCRAS QSVSSSYLAWYQQKPGQAPRLLIYGASSRATGI PDRFSGSGSGTDFTLTISRLEPEDFAVYYCQQY GSSITFGCTRLEIKSGGGGSEVQLVESGGGLV QPGGSLKLSAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMN NLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLVT VSSGGGGSGGGSGG GGSQT VVTQEPSLTVSPGGT VITITCGSSTGAVT SGNYPNWIQKPGQAPRGLIGGTKFLAPGTPA RFGSLEGGKAALTLG VQPEDEAEYYCVLYY SNRWVFGSGTKLTVL</p>
<p>1074.</p>	<p>CL1 9-G4 CC x6H10.09 x scFc xFL 4-E9 CC x6H10.09 - Nucleotide Sequence</p>	<p>artificial</p>	<p>na</p>	<p>CAGGTGCAGCTGGTTCAGTCTGGCGCCGAAG TGAAGAAACCTGGCGCCTCTGTGAAGGTGTC CTGCAAGGCTTCTGGCTACACCTTTACCGAC TACTACATGCACTGGGTCCGACAGGCCCTG GCCAGTGT TTTGGAATGGATGGGCTGGATCAA CCCCAACTCTGGCGGCCCTAATTACGCCAG AAATTCCAGGGCAGAGTGACCATGACCAGA GACACCTCCATCTCCACCGCTCACATGGAAC TGTCCCGGCTGAGATCTGACGACACCGCCGT GTACTACTGCGCCAGAGAAAAGCACGCTGTG GCCGGCATCGGCTTCGATTATTGGGGACAGG GCACCCTGGTCACCGTTTCTAGCGGAGGCGG AGGATCTGGTGGTGGTGGATCTGGCGGCGGA</p>

				GGCTCTGATATCCAGATGACCCAGTCTCCTT CCTCCGTGTCTGCCTCTGTGGGCGACAGAGT GACAATCACCTGTCAGGCCAGCCAGGACATC TCCAACCTGAACCTGGTATCAGCAGAAGC CCGGCAAGGCCCTAAGCTGCTGATCTACGC TGCTCCTCTCTGGAATCTGGCGTGCCCTCCA GATTCTCCGGCTCTGGCTCTGGCACAGACTTT ACCCTGACAATCTCCAGCCTGCAGCCTGAGG ACTTCGCCACCTACTACTGTCAGCAGGCCAA CAGCTTCCCTCTGACCTTTGGCTGTGGCACCA AGGTGGACATCAAGTCTGGTGGCGGGCGGTC CGAAGTCCAGCTGGTTGAAAGTGGTGGCGGA TTGGTTCAGCCAGGCGGCTCTCTGAAGCTGT CTTGTGCTGCCTCCGGCTTCACTTCAACAAA TACGCCATGAATTGGGTTTCGACAAGCCCCAG GCAAAGGCATGGAATGGGTGCGCCGGATCA GATCCAAGTACAACAACCTACGCTACCTACTA CGCCGACGCCGTGAAGGACCGGTTACCATC TCCAGAGATGACTCCAAGAACACCCTGTACC TGCAGATGAACAACCTCAAGACCGAGGATA CCGCCGTCTATTACTGTGTCAGAGCCGGCAA CTTCGGCTCCTCCTACATCTCCTACTTCGCT ACTGGGGCCAGGGAACCCTTGTGACAGTCTC TAGTGGCGGTGGTGGTAGTGGTGGTGGCGGC TCAGGCGGTGGCGGATCTCAAACAGTGGTCA CCCAAGAGCCTAGCCTGACCGTTTCTCCTGG CGGCACCGTGACCATCACATGCGGATCTTCT ACCGGCGCTGTGACCTCCGGCAACTACCCCA ATTGGATCCAGAAGAAGCCAGGCCAGGCTCC TAGAGGACTGATCGGCGGCACAAAGTTTCTG GCTCCCGGCACTCCCGCCAGATTTTCTGGAT CTCTGGAAGGCGGCAAGGCTGCTCTGACATT GTCTGGCGTCCAGCCAGAGGATGAGGCCGA GTACTATTGCGTGCTGTACTACTCCAACAGA TGGGTGTTCCGGCTCCGGCACCAAGCTGACAG TCCTAGGCGGCGGAGGATCTGGCGGAGGTG GAAGCGGAGGCGGTGGATCTGACAAGACCC ACACATGTCCTCCATGTCCCGCCCCTGAACT GCTAGGCGGACCTAGCGTGTTCCCTGTTCCCC CCAAAGCCCAAGGACACCCTGATGATCAGCC GTACGCCCGAAGTGACCTGCGTGGTGGTGG TGTGTCCCACGAGGACCCTGAAGTGAAGTTC AATTGGTACGTGGACGGCGTGGAAGTGCACA ACGCCAAGACCAAGCCCTGCGAGGAACAGT ACGGCAGCACCTACAGATGCGTGTCCGTGCT GACCGTGCTGCATCAGGACTGGCTGAACGGC AAAGAGTACAAGTGCAAGGTGTCCAACAAG GCCCTGCCTGCCCCATCGAGAAAACCATCA GCAAGGCCAAGGGCCAGCCCCGCGAGCCTC AAGTGTATACCCTGCCCCCTAGCCGGGAAGA GATGACCAAGAACCAGGTGTCCCTGACCTGT CTCGTGAAGGGCTTCTACCCCTCCGATATCG CCGTGGAATGGGAGAGCAACGGCCAGCCCG AGAACAACCTACAAGACCACCCCCCTGTGCT GGACAGCGACGGCTCATTCTTCCCTGTACTCC AAACTGACCGTGGACAAGAGCCGGTGGCAG
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				CAGGGCAACGTGTTTCAGCTGCAGCGTGATGC ACGAGGCCCTGCACAACCACTACACCCAGAA GTCCCTGTCCCTGTCTCCCGGGAAAGGCGGC GGAGGATCTGGCGGAGGCGGATCTGGGGGC GGAGGAAGTGGGGGAGGGGAAGCGGAGG GGGAGGCTCAGGGGGGGGAGGATCCGATAA GACCCACACCTGTCCCCCTTGCCCTGCCCTG AACTGCTGGGAGGCCCTAGCGTGTTCTGT CCCCCAAAGCCCAAGGACACCCTGATGATC AGCCGGACCCCCGAAGTGACCTGCGTGGTGG TGGATGTGTCCCACGAGGACCCTGAAGTGAA GTTCAATTGGTACGTGGACGGCGTGGAAGTG CACAAACGCAAGACCAAGCCCTGCGAGGAA CAGTACGGCAGCACCTACAGATGCGTGTCCG TGCTGACCGTGCTGCACCAGGACTGGCTGAA CGGCAAAGAGTACAAGTGCAAGGTGTCCAA CAAGGCCCTGCCTGCCCCCATCGAGAAAACC ATCAGCAAGGCCAAGGGCCAGCCCCGCGAG CCTCAAGTGTATACCCTGCCCCCGAGCCGGG AAGAGATGACCAAGAACCAGGTGTCCCTGA CCTGTCTCGTGAAGGGCTTCTACCCCTCCGAT ATCGCCGTGGAATGGGAGAGCAACGGCCAG CCCGAGAACAACCTACAAGACCACCCCCCTG TGCTGGACAGCGACGGCTCATTCTTCTGTA CTCCAAGCTGACAGTGGACAAGTCTAGATGG CAGCAGGGCAACGTGTTTCAGCTGCAGCGTGA TGCACGAGGCCCTGCACAACCACTACACCCA GAAGTCCCTGTCCCTGAGCCCCGGCAAAGGT GGAGGCGGATCTGGCGGTGGCGGGAGTGGA GGAGGAGGCAGCCAGGTGACTCTGAAAGAA TCCGGTCCCCTCTCGTCAAGCCTACCGAAA CTCTGACCCTGACGTGTACTGTCAGTGGGTTT TCCTTCAGGAATGCACGAATGGGTGTAAGCT GGATACGCCAACCACTGGCAAATGCCTGGA ATGGCTCGCTCACATCTTCAGCAATGACGAG AAGTCCTATTCTACCTCCCTGAAATCCCGGTT GACCATTTCCAAGGATACGAGCAAGTCTCAG GTTGTGCTGACCATGACCAACATGGATCCCG TGGATACAGCCACCTACTTCTGTGCTCGTGT CCCGAGTATAGCTCTGGCTGGTATCGGTTTG ACTACTGGGGACAGGGCACATTGGTGACAGT ATCTTCAGGAGGCGGCGGGTCAGGTGGCGG AGGATCAGGCGGTGGTGGTTCTGACATTCAG ATGACTCAGAGCCCATCAAGTCTGAGTGCCA GTGTTGGAGATAGAGTGACCATCAGTTGCAG AGCCTCTCAGTCTATCAGGAGCTACCTTAAC TGGTATCAGCAGAAACCCGGCAAAGCTCCTA AGCTGCTGATCTACGCAACTAGCAGCCTTCA AGGAGGGGTGCCATCCCGCTTTAGTGGGTCA GGATCTGGCACTGACTTTACCCTCACAATCA GTCCTTGCAACCTGAGGACTTTGCCACCTA CTACTGCCAGCAGTCTATTCCACACCCTTCA CATTCCGGGTGTGGGACAAAGGTCGAGATTAA GTCCGGAGGCGGAGGATCTGAAGTGCAGCT GGTTGAATCTGGCGGCGGATTGGTTCAGCCT GGCGGATCTCTGAAGCTGTCTTGTGCCGCT
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				<p>CTGGCTTCACCTTCAACAAATACGCCATGAA CTGGGTCCGACAGGCCCTGGCAAAGGCATG GAATGGGTCCGCCGGATCAGATCCAAGTACA ACAACACTACGCTACCTACTACGCCGACGCCGT GAAGGACCGGTTACCATCTCCAGAGATGAC TCCAAGAACACCCTGTACCTGCAGATGAACA ACCTCAAGACCGAGGACACCGCCGTGTACTA CTGTGTCAGAGCCGGCAACTTCGGCTCCTCC TACATCTCCTACTTCGCCTATTGGGGCCAGG GCACCCTGGTCACAGTTAGTTCAGGTGGCGG TGGATCAGGCGGCGGAGGTTCTGGTGGCGGA GGCTCTCAAACAGTGGTCACCCAAGAGCCTA GCCTGACCGTTTCTCCTGGCGGCACCGTGAC CATCACCTGTGGATCTTCTACCGGCGCTGTG ACCTCCGGCAACTACCCAATTGGATCCAGA AGAAGCCC GGCCAGGCTCCTAGAGGACTGAT CGGAGGCACCAAGTTTCTGGCTCCCGGCACT CCTGCCAGATTCTCCGGTTCTCTGGAAGGCG GAAAGGCCGCTCTGACATTGTCTGGCGTGCA GCCTGAGGATGAGGCTGAGTACTACTGCGTG CTGTACTACTCCAACAGATGGGTGTTCCGGCT CCGGCACCAAGCTGACAGTGCTT</p>
1075.	MS 15-B12 CC x4F10.03 I2M xscFc xCH3 15- E11 CC x4F10.03 I2M - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFLTISLQPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG TSYISYWAYWGQGLVTVSSGGGGSGGGGSG GGGSQTVVTQEPSLTVSPGGTVTITCGSSTGAV TSGNYPNWVQKKPGQAPRGLIGGTKFLAPGTP ARFSGSLSGGKAALTLGSGVQPEDEAEYYCVLW YSNRWVFGSGTKLTVLGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGYSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSQSVSMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGYSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSQSVSMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFKQGRVTMTVDTSS</p>

				<p>STAYMELSRLSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQOKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKGMWVVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGTSYISYWAYWGG GTLTVTVSSGGGGSGGGGSGGGGSGTQVVTQEPS LTVSPGGTVTITCGSSTGAVTSGNYPNWVQKK PGQAPRGLIGGTKFLAPGTPARFSGSLSGGKAA LTLGSGVQPEDEAEYYCVLWYSNRWVFGSGTK LTVL</p>
1076.	MS 15-B12 CC x I2L x scFc xCH3 15-E11 CC x I2M2 - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQKPKGKVPKLLIYAASLTQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQSSY TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGGTLTVTVSSGGGGSGGGGSGG GGGSGTQVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKPKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAAALTLGSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSVMSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSVMSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFQGRVTMTVDTSS STAYMELSRLSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQOKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAINWVREAPGKGLEWVARIRSKYN NYATYYADAVKDRFTISRDDSKNTAYLQMN</p>

				LKTEDTAVYYCVRNANFGTSYISYFAYWGQG TLVTVSSGGGGSGGGGGSGGGGSQTVVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQKKP GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGSGTKL TVL
1077.	MS 15-B12 CC x I2C x scFc xCH3 15-E11 CCx I2C0 - Full Sequence	artificial	Aa	QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSY TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GLEWVARIRSKYNNYATYYADSVKDRFTISR DSKNTAYLQMNNLKTEDTAVYYCVRHGNFGN SYISYWAYWGQGTTLVTVSSGGGGSGGGGSGG GGSTVVTQEPSLTVSPGGTVTLTCGSSTGAVT SGNYPNWVQKPGQAPRGLIGGTKFLAPGTPA RFSGSLLGGKAALTLSGVQPEDEAEYYCVLWY SNRWVFGGGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHGDWLNKG EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHGDWLN KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEW SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKQGRVTMTVDTSS STAYMELSRLRSDDTAVYYCATRYFYVMDYW GQGTTLVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQQK GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKGLEWVARIRSKYN NYATYYADSVKDRFTISRDDSKNTAYLQMN LKTEDTAVYYCVRHGNFGNSYISYWAYWGQG TLVTVSSGGGGSGGGGSGGGGSQTVVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQK GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGGGTKL TVL
1078.	MS 15-B12 CC x I2L x scFc x CH3	artificial	Aa	QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYSGSSNYNPSLKS

	<p>15-E11 CC x I2L - Full Sequence</p>			<p>RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAMNWVRQAPGK GMEWVARIRSKYNNYATYYADAVKDRFTISR DDSKNTLYLQMNNLKTEDTAVYYCVRAGNFG SSYISYFAYWGQGLTVTVSSGGGGSGGGGSGG GGSTVVTQEPSLTVSPGGTVTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVLGGGGSGGGGSGGGGSD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFQGRVTMTVDTSS STAYMELSRLRSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFLT TISSLPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSCAAS GFTFNKYAMNWVRQAPGKMEWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ GTLTVTVSSGGGGSGGGGSGGGGSGTQVVTQEPS LTVSPGGTVTITCGSSTGAVTSGNYPNWIQKKP GQAPRGLIGGTKFLAPGTPARFSGSLEGGKAAL TLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL</p>
<p>1079</p>	<p>MS 15-B12 CC x I2M2 x scFc x CH3 15-E11 CC x I2M2 - Full Sequence</p>	<p>artificial</p>	<p>Aa</p>	<p>QVQLQESGPGLVKPKSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFLTITSSLPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSCAASGFTFNKYAINWVREAPGKG LEWVARIRSKYNNYATYYADAVKDRFTISRDD</p>

				<p>SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGGGGSGGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTPVLDSGDSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYYWMMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFOGRVTMTVDTSS STAYMELSRLRSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRVTITCRASQDISNYLNWYQQKP GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSKAAS GFTFNKYAINWVREAPGKGLEWVARIRSKYN NYATYYADAVKDRFTISRDDSKNTAYLQMNN LKTEDTAVYYCVRNANFGTSYISYFAYWGQ TLTVTVSSGGGGSGGGGSGGGGSGQTVVTQEPSL TVSPGGTVTLTCGSSTGAVTSGNYPNWVQKKP GQAPRGLIGGTKFLAPGTPARFSGSLLGGKAAL TLSGVQPEDEAEYYCVLWYSNRWVFGSGTKL TVL</p>
1080.	MS 15-B12 CC x I2M2 x scFc x CH3 15-E11 x I2L - Full Sequence	artificial	Aa	<p>QVQLQESGPGLVKPSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGMVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRVTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASSTLQSGVP SRFSGSGSGTDFTLTISSLQPEDFATYYCQQSYS TPFTFGCGTKVEIKSGGGGSEVQLVESGGGLV QPGGSLKLSKAASGFTFNKYAINWVREAPGK LEWVARIRSKYNNYATYYADAVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRNANFGTS YISYFAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQKKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGSGTKLTVLGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA</p>

				<p>KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYKTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGSGGGSGGGSGGGSGGGSGGGGS DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTPPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGSGGGSGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYFTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFQGRVTMTVDTSS STAYMELSRLRSDDTAVYYCATRYFYVMDYW GQGLTVTVSSGGGGSGGGSGGGSGDIQMTQS PSSLSASVGDRTITCRASQDISNYLNWYQQKQ GKVPKLLIYYTSRLHSGVPSRFSGSGSGTDFTL TISSLQPEDVATYYCVQYAQFPLTFGCGTKVEI KSGGGGSEVQLVESGGGLVQPGGSLKLSKAAS GFTFNKYAMNWVRQAPGKGMWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ GTLTVTVSSGGGGSGGGSGGGSGTQVVTQEPS LTVSPGGTVTITCGSSTGAVTSGNYPNWIQKKP GQAPRGLIGGTKFLAPGTPARFSGSLEGGKAAL TLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLT VL</p>
1081.	Ubiquitin	artificial	Aa	<p>MQIFVKTLTGKTITLEVEPSDTIENVKAKIQDKE GIPPDQQRLLIFAGKQLEDGRTLSDYNIQKESTL HLVLRLRGG</p>
1082.	HSP70-1	artificial	Aa	<p>AAAIGIDLGTTYSCVGVFQHGKVEIANDQGNR TTPSYVAFTDTERLIGDAAKNQVALNPQNTVF DAKRLIGRKFQDPVVQSDMKHWPFFQVINDGD KPKVQVSYKGETKAFYPEEISSMVLTKMKEIA EAYLGYPTNAVITVPAYFNDSQRQATKDAGV IAGLNVLRIINEPTAAAIA YGLDRTGKGERNVLI FDLGGGTFDVSILTIDDGIFEVKATAGDTHLGG EDFDNRLVNHFVEEFKRKHKKDISQNKRAVRR LRTACERAKRTLSSSTQASLEIDSLFEGIDFYTSI TRARFEELCSDLFRSTLEPVEKALRDAKLDKA QIHDLVLVGGSTRIPKVQKLLQDFFNDRDLNKS INPDEAVAYGAAVQAAILM</p>
1083.	beta 2 microglobulin	artificial	Aa	<p>MIQRTPKIQVYSRHPAENGKSNFLNCYVSGFHP SDIEVDLLKNGERIEKVEHSDLSFSKDWSFYLL YYTEFTPTEKDEYACRVNHVTLSPKIVKWD</p>
1084.	SAND domain	artificial	Aa	<p>DMEIAYPITCGESKAILLWKKFVCPGINVKCVK FNDQLISPKHFVHLAGKSTLKDWKRAIRLGGI MLRKMMDSGQIDFYQHDKVCSNTR</p>
1085.	Green fluorescent protein (GFP)	artificial	Aa	<p>MSKGEELFTGVVPIVVELDGDVNGHKFSVSGE GEGDATYGKLTCLKFICTTGKLPVPWPTLVTTFT YGVQCFSRYPDHMKRHDFFKSAMPEGYVQER TIFFKDDGNYKTRAEVKFEGDTLVNRIELKGID FKEDGNILGHKLEYNYNSHNVYIMADKQKNGI</p>

				KVNFKIRHNIEDGSVQLADHYQQNTPIGDGPV LLPDNHYLSTQSALSKDPNEKRDHMLLEFVT AAGITHGMDELK
1086.	VHH antibody lama domain	artificial	Aa	QVQLQESGGGLVQAGDSLKLSCEASGDSIGTY VIGWFRQAPGKERIYLATIGRNLVGPSDFYTRY ADSVKGRFAVSRDNAKNTVNLQMNSLKPEDT AVYYCAAKTTTWGGNDPNNWNYWGQGTQV TVSS
1087.	PSI domain from Met-receptor	artificial	Aa	GSAMGCRHFQSCSQCLSAPPFVQCGWCHDKC VRSEECLSGTWTQQICL
1088.	Fibronectin type III domain from tenascin	artificial	Aa	RLDAPSQIEVKDVTDTTALITWFKPLAEIDGIEL TYGIKDVPGDRRTTIDLTEDENQYSIGNLKPDE YEVSLISRRGDMSSNPAKETFTT
1089.	Granulocyte- macrophage colony-stimulating factor (GM-CSF)	artificial	Aa	APARSPSPSTQPWEHVNAIQEARLLNLSRDTA AEMNETVEVISEMFDLQEPTCLQTRLELYKQG LRGSLTKLKGPLTMMASHYKQHCPTPETS CA TQIITFESFKENLKDFLLVIPFDCWEPVQE
1090.	Interleukin-4	artificial	Aa	HKCDITLQEIITLNSLTEQKTLCTELTVDIFA ASKNTEKETFCRAATVLRQFYSHHEKDTRCL GATAQQFHRHKQLIRFLKRLDRNLWGLAGLNS CPVKEANQSTLENFLERLKTIMREKYSKCSS
1091.	CD137L Ectodomain	artificial	Aa	DPAGLLDLRQGMFAQLVAQNVLLIDGPLSWYS DPGLAGVSLTGGLSYKEDTKELVAKAGVYY VFFQMELELRVAGEGSGSVSLALHLMPLRSAA GAAALALTVDLPPASSEARNSAFGFQGRLLHL SAGQRLGVHLHTEARARHAWQLTQGATVGLL FRVTPEIPA
1092.	Interleukin-2	artificial	Aa	APTSSSTKKTQLQLEHLLDLQMLNGINNYKN PKLTRMLTFKPYMPKKATELKHLCLEELKP LEEVLNLAQSKNFHLRPRDLISNINVIVLELKGS ETTFMCEYADETATIVEFLNRWITFAQSIISTLT
1093.	PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1)	artificial	Aa	AFTVTVPKDLVYVEYGSNMTIECKFPVEKELD LAALIVYWEMEDKNIIQFVHGEECLKVQHSSY RQRARLLKDQLSLGNAALQITDVKLQDAGVY RCMISYGGADYKRITVKVNAPYAAALEHHHH
1094.	Tim-3 (AS 24- 130)	artificial	Aa	SEYRAEVLGQNAYLPCFYTPAAPGNLVPVCWG KGACPVFECGNVLRRTDERDVNYWTSRYWLN GDFRKGDVSLTIENVTLADSGIYCCRIQIPGIMN DEKFNKLVK
1095.	MiniSOG	artificial	Aa	MEKSFVITDPRLPDNPIIFASDGFLELTEYSREEI LGRNGRFLQGPETDQATVQKIRDAIRDQREITV QLINYTKSGKKFWNLLHLQPMRDQKQELQYFI GVQLDGEFIPNPLLG
1096.	A(EAAAK)4ALE A(EAAAK)4A	artificial	Aa	AEAAAKEAAAKEAAAKEAAAKALEAEAAAK EAAAKEAAAKEAAAKA
1097.	(PA)25P	artificial	Aa	PAPAPAPAPAPAPAPAPAPAPAPAPAPAPAP APAPAPAPAPAPAPAPAP
1098.	SpyCatcher	artificial	Aa	VTTLSGLSGEQGPSGDMTTEEDSATHIKFSKRD EDGRELAGATMELRDSSGKTISTWISDGHVKD FYLYPGKYTFVETAAPDGYEVATAITFTVNEQ GQVTVNGEATKGAHT
1099.	SpyTag	artificial	Aa	VPTIVMVDAYKRYK
1100.	DogTag	artificial	Aa	DIPATYEFTDGKHYITNEPIPPK

1101.	SnoopTagJr	artificial	Aa	KLGSIEFIKVNK
1102.	MS 15-B12 CC x G4S3x heFc(A) x G4S3 x CH3 15-E11 CC	artificial	Aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSSGGGGSGGGG SGGGGSDIVMTQSPSSLSASVGDRTITCRASQ GISNYLAWYQQKPGKVPKLLIYAASTLQSGVP SRFSGSGGTDFLTITSSLPEDFATYYCQSYS TPFTFGCGTKVEIKSGGGGSGGGGSGGGGSDK THTCPPCPAPELLGGPSVFLFPPKPKDTLMISRT PEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG KGGGGSGGGGSGGGGSGVQLVQSGAEVKKPG ASVKVSCKASGYTFTNYWMNWVRQAPGQCL EWMGNIAYGVKGTNYNQKFGQGRVTMTVDTSS STAYMELSRLRSDDTAVYYCATRYFYVMDYW GQGTLVTVSSGGGGSGGGGSGGGGSDIQMTQS PSSLSASVGDRTITCRASQDISNYLNWYQQK GKVPKLLIYYTSRLHSGVPSRFSGSGGTDFLT TISSLPEDVATYYCVQYAQFPLTFGCGTKVEI K
1103.	6H10-09 x G4S3 x heFc(B) x GS3 x 6H10.09	artificial	Aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDA VYYCVRAGNFGSSYISYFAYWGQGTLVTVSSG GGGSGGGGSGGGGSGTQVVTQEPSLTVSPGGTV TITCGSSTGAVTSGNYPNWIQKPGQAPRGLIG GTKFLAPGTPARFSGSLEGGKAALTLGVPQED EAEYYCVLYYSNRWVFGSGTKLTVLGGGGSG GGGSGGGGSDKTHTCPPCPAPELLGGPSVFLFP PKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSRKEMTKNQVSLTCLVKG FYPSDIAVEWESNGQPENNYKTPPVLKSDF FLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKLSLSPGKGGGGSGGGGSGGGGSEVQLV ESGGGLVQPGGSLKLSAASGFTFNKYAMNW VRQAPGKGMWVARIRSKYNNYATYYADAV KDRFTISRDDSKNTLYLQMNNLKTEDTAVYYC VRAGNFGSSYISYFAYWGQGTLVTVSSGGGG GGGSGGGGSGTQVVTQEPSLTVSPGGTVTITC GSSTGAVTSGNYPNWIQKPGQAPRGLIGTK FLAPGTPARFSGSLEGGKAALTLGVPQEDAE YYCVLYYSNRWVFGSGTKLTVL
1104.	CL1 9-G4 scFab8 x G4S x I2Ccc x G4 x scFc x G4 x FL 4-E9 scFab8 xG4S xI2Ccc - Full Sequence	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTD YMHWVRQAPGQGLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGTLVTVSSAST KGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEP VTVSWNSGALTSGVHTFPAVLQSSGLYSLSSV VTPVSSSLGTQTYICNVNHKPSNTKVDKKEP

	- HCDR1			
1106.	CL1 9-G4 scFab8 - HCDR2	artificial	Aa	WINPNSGGPNYAQKFQG
1107.	CL1 9-G4 scFab8 - HCDR3	artificial	Aa	EKHAVAGIGFDY
1108.	CL1 9-G4 scFab8 - LCDR1	artificial	Aa	QASQDISNYLN
1109.	CL1 9-G4 scFab8 - LCDR2	artificial	Aa	AASSLES
1110.	CL1 9-G4 scFab8 - LCDR3	artificial	Aa	QQANSFPLT
1111.	CL1 9-G4 scFab8 - VH	artificial	Aa	QVQLVQSGAEVKKPGASVKVSCASGYTFTD YYMHWVRQAPGQGLEWMGWINPNSGGPNYA QKFQGRVTMTRDTSISTAHMELSRLRSDDTAV YYCAREKHAVAGIGFDYWGQGTLTVSS
1112.	CL1 9-G4 scFab8 - VL	artificial	Aa	DIQMTQSPSSVSASVGDRVTITCQASQDISNYL NWXQQKPGKAPKLLIYAASSLESGVPSRFGSG GSGTDFTLTISSLQPEDFATYYCQQANSFPLTFG PGTKVDIK
1113.	CL1 9-G4 scFab8 - CH1	artificial	Aa	ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSL SVVTVPSSSLGTQTYICNVNHKPSNTKVDKKV EPKSC
1114.	CL1 9-G4 scFab8 - CLK	artificial	Aa	RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFY PREAKVQWKVDNALQSGNSQESVTEQDSKDS TYSLSSTLTLSKADYEKHKVYACEVTHQGLSS PVTKSFNRGEC
1115.	FL 4-E9 scFab8 - HCDR1	artificial	Aa	NARMGV
1116.	FL 4-E9 scFab8 - HCDR2	artificial	Aa	HIFSNDEKSYSTSLKS
1117.	FL 4-E9 scFab8 - HCDR3	artificial	Aa	VPEYSSGWYRFDY
1118.	FL 4-E9 scFab8 - LCDR1	artificial	Aa	RASQSIRSYLN
1119.	FL 4-E9 scFab8 - LCDR2	artificial	Aa	ATSSLQG
1120.	FL 4-E9 scFab8 - LCDR3	artificial	Aa	QQSYSTPFT
1121.	FL 4-E9 scFab8 - VH	artificial	Aa	QVTLKESGPTLVKPTETLTLTCTVSGFSFRNAR MGVSWIRQPPGKALEWLAHIFSNDEKSYSTSL KSRLTISKDTSKSQVVLTMNMDPVDATATYFC ARVPEYSSGWYRFDYWGQGTLTVSS
1122.	FL 4-E9 scFab8 - VL	artificial	Aa	DIQMTQSPSSLSASVGDRVTISCRASQSIRSYLN WYQQKPGKAPKLLIYATSSLQGGVPSRFGSG SGTDFTLTISSLQPEDFATYYCQQSYSTPFTFGP GTKVEIK
1123.	FL 4-E9 scFab8 - CH1	artificial	Aa	ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSL SVVTVPSSSLGTQTYICNVNHKPSNTKVDKKV EPKSC
1124.	FL 4-E9 scFab8 - CLK	artificial	Aa	RTVAAPSVFIFPPSDEQLKSGTASVVCLLNNFY PREAKVQWKVDNALQSGNSQESVTEQDSKDS TYSLSSTLTLSKADYEKHKVYACEVTHQGLSS PVTKSFNRGEC

1153.	CD3 FHar-LCDR3	artificial	Aa	TLWYSNRWV
1154.	CDH3 HCDR1 consensus	artificial	Aa	XXXXXX
1155.	CDH3 HCDR2 consensus	artificial	Aa	XIXXXXXXTXYXXXXXG
1156.	CDH3 HCDR3 consensus	artificial	Aa	SRGVYDXXXXXXXXYXMDX
1157.	CDH3 VH consensus	artificial	Aa	XVQLXXSGXXXXXPGXSXXXSCXASGXXFXX XXXXWVRQXPGXCLEWXXXIXXXXXTXYYX XXXGRXTXXDXSXXTYXXXXXXLXXDXA VYYCAXXRGVYDFKXXXALXXDXWGQGT XVTVSS
1158.	CDH3 LCDR1 consensus	artificial	Aa	XXSXXXLYSSNQXXYXX
1159.	CDH3 LCDR2 consensus	artificial	Aa	XXXXXXXX
1160.	CDH3 LCDR3 consensus	artificial	Aa	XXXXXXPXT
1161.	CDH3 VL consensus	artificial	Aa	XIXXTQSPXXLXXSXGXRXTXXCXXSXXXLY NQXXYXXWYQQKPGXXPXLXYYXXXXXXG VPXRFSGSGSGTFTLXISXLQXEDXXXYYCX XXXXXPXTFGCGTKXXIK
1162.	MSLN HCDR1 consensus	artificial	Aa	SSXYXXX
1163.	MSLN HCDR2 consensus	artificial	Aa	XIXXXXXXXXXYXXXXXX
1164.	MSLN HCDR3 consensus	artificial	Aa	XXXXXGXXSYXPXXXXXXXXDX
1165.	MSLN VH consensus	artificial	Aa	XVQLXXSGXXXXXPXXXXXXXXCXXSGGSXX XXYXXXWXRQXPGXXLEWXXXIXXXXXXX YXXXXXXRXTXXDXXXXXXXXXXXXXXXXX XXDTAVYYCAXXXXXXXXXXSYPXYXXXX DXWGQGTXTVTVSS
1166.	MSLN LCDR1 consensus	artificial	Aa	RXXXXXXXXXXXX
1167.	MSLN LCDR2 consensus	artificial	Aa	XXXXXXXX
1168.	MSLN LCDR3 consensus	artificial	Aa	QXXXXXXXXIXX
1169.	MSLN VL consensus	artificial	Aa	XXXXTQXPXXXSXSXG1XXXXXCRXXXXXXXX XXXXWYQQKPGXXPXLXIYXXXXXXXXGX RFSGSXSGXXTLTISXXXXDXAXYYCQXXX XXXIXXFGXGTKXXXX
1170.	Human CDH3	human	Aa	MGLPRGPLASLLLLQVCWLQCAASEPCRAVFR EAEVTLEAGGAEQEPGQALGKVMGCPGQEP ALFSTDNDFFTVRNGETVQERRSLKERNPLKIF PSKRILRRHKRDWVAPISVPENGKGFPPQRLN QLKSNKDRDTKIFYSITGPGADSPPEGVFAVEK ETGWLLLNKPLDREEIAKYELFGHAVSENGAS VEDPMNISIIVTDQNDHKPKFTQDTRGVSLEG VLPGTSVMQVTATDEDDAIYTYNGVVAYSIS QEPKDPHDLMFTHIRSTGTISVISSGLDREKVPE YTLTIQATDMDGDGSTTTAVAVVEILDANDNA PMFDPQKYEAHVPENAVGHEVQRLTVTDLDA

				PNSPAWRATYLIMGGDDGDHFTITTHPESNQGI LTTRKGLDFEAKNQHTLYVEVTNEAPFVLKLP TSTATIVVHVEDVNEAPVFPVPSKVVEVQEGIP TGEPVCVYTAEDPKENQKISYRILRDPAGWL AMDPDSGQVTA VGTLDREDEQFVRNNIYEV VLAMDNGSPPTTGTGTLTLLTLIDVNDHGPVPEP RQITICNQSPVRQVLNITDKDLSPHSPFQAQLT DDSDIYWTAEVNEEGDTVVL SLKKFLKQDTYD VHLSLSDHGNKEQLTVIRATVCDCHGHVETCP GPWKGGFILPVLGAVLALLFLLLVLVLLVRKK RKIKEPLLPEDDTRDNVFFYGGEEGGEEEDQD YDITQLHRGLEARPEVVL RNDVAPTIIPTPMYR PRPANPDEIGNFIIENLKAANTDPTAPPYDTLLV FDYEGSGSDAASLSSLTSSASDQDQDYDYLNE WGSRFKKLADMYGGGEDD
1171.	Human CDH3 epitope cluster D2B	human	Aa	VAYSIHSEQPKDPHDL MFTIHRSTGTISVISSGL DREK
1172.	Human CDH3 epitope cluster D2C	human	Aa	VPEYTLTIQATDMDGDGSTTTAVAVVEILDAN DNAPM
1173.	Human CDH3 epitope cluster D3A	human	Aa	FDPQKYEAHVPENAVGHEVQRLTVTDLDAPNS PAWR
1174.	Human CDH3 epitope cluster D4B	human	Aa	YRILRDPAGWLAMDPDSGQVTA VGTLDREDE QFVRN
1175.	Human MSLN epitope cluster E1	human	Aa	EVEKTACPSGKKAREIDESLIFYKKWELEACVD AALLATQMDRVNAIPFTY
1176.	Human MSLN epitope cluster E2	human	Aa	EQLDVLKHKLDEL YPQGYPESVIQHLGYLFLK MSPEDI
1177.	Human MSLN epitope cluster E3	human	Aa	RKWNVT SLETLKALLEVNKGHEMSPQVATLID RFVKGRGQLDKDTLDTLTA FYPGYLCSLSPEEL SSVP
1178.	Human MSLN epitope cluster E4	human	Aa	PSSIWAVRPQDLDTCDPRQLDVL YPKARLAFQ NMNGSEYFVKIQSFLG
1179.	Human MSLN epitope cluster E5	human	Aa	GAPTEDLKALSQQNVSM DLATFMKLRTDAVL PLTVAEVQKLLGP
1180.	Human MSLN v1 NM_005823	human	Aa	MALPTARPLL GSCGTPALGSL LFLSLGWVQP SRTLGETGQEAAPLDGVL ANPPNISSLSRQL LGFPCA EVSGLSTERVRELAVALAQKNVKLST EQLRCLAHRLSEPPEDLDALPLDLLLFLNPDAF SGPQACTRFFSRITKANVDLLPRGAPERQRLLP AALACWGV RGSLLSEADV RALGGLACDLPGR FVAESAEVLLPRLVSCPGPLDQDQQAARAAL QGGGPPYGGPSTWSVSTMDALRGLLPVLGQPII RSIPQGIVAAWRQRSSRDPSWRQPERTILRPRF RREVEKTACPSGKKAREIDESLIFYKKWELEAC VDAALLATQMDRVNAIPFTYEQLDVLKHKLD ELYPQGYPESVIQHLGYLFLKMSPEDIRKWNV TSLETLKALLEVNKGHEMSPQVATLIDRFVKG RGQLDKDTLDTLTA FYPGYLCSLSPEELSSVPP SSIWAVRPQDLDTCDPRQLDVL YPKARLAFQN MNGSEYFVKIQSFLGGAPTEDLKALSQQNVSM DLATFMKLRTDAVLPLTVAEVQKLLGPHVEGL

				KAEERHRPV RDWILRQRQDDLDLTLGLGLQGGI PNGYL VLDLSMQEALSGTPCLLGPVLT VLA LLLASTLA
1181.	Human MSLN v2 NM_013404	human	Aa	MALPTARPLLGSCGTPALGSLLFLLFSLGWVQP SRTLAGE TGQEAAPLDGVLANPPNISSLSRQL LGFPCA EVSGLSTERVRELAVALAQKNVKLST EQLRCLAHRLSEPPEDLDALPLDLLLFLNPDAF SGPQACTRFFSRITKANVDLLPRGAPERQRLLP AALACWGV RGSLLSEADV RALGGLACDLPGR FVAESA EVLLPRLVSCPGPLDQDQQAARAAL QGGGPPY GPPSTWSVSTMDALRGLLPVLGQPII RSIPQGIVAAWRQRSSRDPSWRQPERTILRPRF RREVEKTACPSGKKAREIDESLIFYKKWELEAC VDAALLATQMDRVNAIPFTYEQLDVLKHKLD ELYPQGY P ESVIQHLGYLFLKMSPEDIRKWNV TSLETLKALLEVNKGHEMSPQAPRRPLPQVAT LIDRFVKGRGQLDKDTLDTLTA FYPGYLCSLSP EELSSVPPSSIWAVRPQDLDTCDPRQLDVLYPK ARLAFQNMNGSEYFVKIQSFLGGAPTEDLKAL SQQNVSM DLATFMKLRTDAVLPLTVAEVQKL LGPHVEGLKAEERHRPV RDWILRQRQDDLDLTL GLGLQGGIPNGYL VLDLSMQEALSGTPCLLGP GPVLT VLA LLLASTLA
1182.	Human MSLN v6 AY743922	human	Aa	MALPTARPLLGSCGTPALGSLLFLLFSLGWVQP SRTLAGE TGQEAAPLDGVLANPPNISSLSRQL LGFPCA EVSGLSTERVRELAVALAQKNVKLST EQLRCLAHRLSEPPEDLDALPLDLLLFLNPDAF SGPQACTHFFSRITKANVDLLPRGAPERQRLLP AALACWGV RGSLLSEADV RALGGLACDLPGR FVAESA EVLLPRLVSCPGPLDQDQQAARAAL QGGGPPY GPPSTWSVSTMDALRGLLPVLGQPII RSIPQGIVAAWRQRSSRDPSWRQPERTILRPRF RREVEKTACPSGKKAREIDESLIFYKKWELEAC VDAALLATQMDRVNAIPFTYEQLDVLKHKLD ELYPQGY P ESVIQHLGYLFLKMSPEDIRKWNV TSLETLKALLEVNKGHEMSPQVATLIDRFVKG RGQLDKDTLDTLTA FYPGYLCSLSPEELSSVPP SSIWAVRPQDLDTCDPRQLDVLYPKARLAFQ MNGSEYFVKIQSFLGGAPTEDLKALSQQNVSM DLATFMKLRTDAVLPLTVAEVQKLLGPHVEGL KAEERHRPV RDWILRQRQDDLDLTLGLGLQGGI PNGYL VLDLSVQEALSGTPCLLGPVLT VLA L LLASTLA
1183.	MSLN 5F11 xI2C -scFc _HLE_bispecific molecule	artificial	aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDY YMTWIRQAPGKGLEWLSYISSSGSTIYYADSV KGRFTISRDNKNSLFLQMNSLRAEDTAVYYC ARDRNSHF DYWGQGLVTVSSGGGGSGGGGS GGGSDIQMTQSPSSVSASVGDRTITCRASQG INTWLAWYQQKPGKAPKLLIYGASGLQSGVPS RFSGSGSGTDFTLTISSLQPEDFATYYCQQAQSF PRTFGQGTKVEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSCAASGFTFNKYAMNWVRQAPGKG LEWVARIRSKYNNYATYYADSVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRHGNFGNS YISYWAYWGQGLVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGT VTLTCGSSTGAVTS

				GNYPNWVQQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLSGVQPEDEAEYYCVLWYS NRWVFGGGTKLTVLGGGGDKTHTCPPCPAPEL LGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGSGGGGS GGGGSGGGSGGGSGGGSGGGSDKTHTCPPCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGK
1184.	G4 - Linker	artificial	aa	GGGG
1185.	G5 - Linker	artificial	aa	GGGGG
1186.	SG4Q - Linker	artificial	aa	SGGGGQ
1187.	(G4Q) ₃ - Linker	artificial	aa	GGGGQGGGGQGGGGQ
1188.	scFc_clipopt	artificial	aa	CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEEPEVKFNWYVDGVEVHNAKTK PCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCK KVSNAKALPAPIEKTISKAKGQPREPQVYTLPPS REEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ PENNYKTTTPVLDSDGSFFLYSKLTVDKSRWQ QGNVFSVMHEALHNHYTQKSLSLSPGKGG GGQGGGGQGGGGQGGGGQGGGGQGGGGQCP PCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTC VVVDVSHEEPEVKFNWYVDGVEVHNAKTKPC EEQYGSTYRCVSVLTVLHQDWLNGKEYKCKV SNKALPAPIEKTISKAKGQPREPQVYTLPPSREE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPEN NYKTTTPVLDSDGSFFLYSKLTVDKSRWQQGN VFSCVMHEALHNHYTQKSLSLSPGK
1189.	heFc(A)	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYDTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSVMHEALHNHYTQDSLSLSPG K
1190.	heFc(B)	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTTPVLDSDGSFFLYSKLTVDK SRWQQGNVFSVMHEALHNHYTQKSLSLSP GK
1191.	heFc(A) dDKTHT	artificial	aa	CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEDPEVKFNWYVDGVEVHNAKTK PCEEQYGSTYRCVSVLTVLHQDWLNGKEYK

				CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYDTTPPVLDSDGSFFLYSDLTVDKSRW QQGNVVFSCSVMHEALHNHYTQDSLSLSPGK
1192.	heFc(B) dDKTHT	artificial	aa	CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEDPEVKFNWYVDGVEVHNAKT KPCSEEQYGSTYRCVSVLTVLHQDWLNGKEYK CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SRKEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTTPPVLKSDGSFFLYSKLTVDKSRW QQGNVVFSCSVMHEALHNHYTQKLSLSLSPGK
1193.	I2L scFv - HCDR1	artificial	aa	KYAMN
1194.	I2L scFv - HCDR2	artificial	aa	RIRSKYNNYATYYADAVKD
1195.	I2L scFv - HCDR3	artificial	aa	AGNFGSSYISYFAY
1196.	I2L scFv - LCDR1	artificial	aa	GSSTGAVTSGNYPN
1197.	I2L scFv - LCDR2	artificial	aa	GTKFLAP
1198.	I2L scFv - LCDR3	artificial	aa	VLYYSNRWV
1199.	I2L scFv - VH	artificial	aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLVTVSS
1200.	I2L scFv - VL	artificial	aa	QTVVTQEPSLTVSPGGTVTITCGSSTGAVTSGN YPNWIQKKPGQAPRGLIGGTKFLAPGTPARFSG SLEGGKAALTLSGVQPEDEAEYYCVLYYSNR WVFGSGTKLTVL
1201.	antiMSLN_01 scFv - HCDR1	artificial	aa	SSSYFWG
1202.	antiMSLN_01 scFv - HCDR2	artificial	aa	NIYYSGSSNYNPSLKS
1203.	antiMSLN_01 scFv - HCDR3	artificial	aa	LPRGDRDAFDI
1204.	antiMSLN_01 scFv - LCDR1	artificial	aa	RASQGISNYLA
1205.	antiMSLN_01 scFv - LCDR2	artificial	aa	AASTLQS
1206.	antiMSLN_01 scFv - LCDR3	artificial	aa	QQSYSTPFT
1207.	antiMSLN_01 scFv - VH	artificial	aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSS
1208.	antiMSLN_01 scFv - VL	artificial	aa	EIVMTQSPSSLSASVGDRVTITCRASQGISNYLA WYQQKPGKVPKLLIYAASTLQSGVPSRFSGSG SGTDFTLTISSLQPEDFATYYCQQSYSTPFTFGC GTKVEIK
1209.	antiCDH3_01 scFv - HCDR1	artificial	aa	NYWMN
1210.	antiCDH3_01 scFv - HCDR2	artificial	aa	NIAYGVAGTNYNQKFQG
1211.	antiCDH3_01 scFv - HCDR3	artificial	aa	RYFYVMDY
1212.	antiCDH3_01	artificial	aa	RASQDISNYLN

	scFv - Lcdr1			
1213.	antiCDH3_01 scFv - Lcdr2	artificial	aa	YTSRLHS
1214.	antiCDH3_01 scFv - Lcdr3	artificial	aa	VQYAQFPLT
1215.	antiCDH3_01 scFv - VH	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSS
1216.	antiCDH3_01 scFv - VL	artificial	aa	EIQMTQSPSSLSASVGDRVTITCRASQDISNYLN WYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGS GTDFTLTISLQPEDVATYYCVQY AQFPLTFGC GTKVEIK
1217.	I2L VH-CH1 - HCDR1	artificial	aa	KYAMN
1218.	I2L VH-CH1 - HCDR2	artificial	aa	RIRSKYNNYATYYADAVKD
1219.	I2L VH-CH1 - HCDR3	artificial	aa	AGNFGSSYISYFAY
1220.	I2L VL-CL - LCDR1	artificial	aa	GSSTGAVTSGNYPN
1221.	I2L VL-CL - LCDR2	artificial	aa	GTKFLAP
1222.	I2L VL-CL - LCDR3	artificial	aa	VLYYSNRWV
1223.	I2L VH-CH1 - VH	artificial	aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGM EWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVTVSS
1224.	I2L VL-CL - VL	artificial	aa	QTVVTQEPSLTVSPGGT V TITCGSSTGAVTSGN YPNWIQKKPGQAPRGLIGGTKFLAPGTPARFSG SLEGGKAAL T LSGVQPEDEAEYYCVLYYSNR WVFGSGTKLTVL
1225.	I2L VH-CH1 - CH1	artificial	aa	ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLE SVVTVPSSSLGTQTYICNVNHKPSNTKVDKKV EPKSC
1226.	I2L VL-CL - CL	artificial	aa	GQPKAAPSVTLFPPSSEELQANKATLVCLISDF YPGAVTVAWKADSSPVKAGVETTPSKQSN KYAAKSYLSLTPEQWKSHRSYSCQVTHEGSTV EKTVAPECS
1227.	antiMSLN_01 VH-CH1 - HCDR1	artificial	aa	SSSYFWG
1228.	antiMSLN_01 VH-CH1 - HCDR2	artificial	aa	NIYYSGSSNYNPSLKS
1229.	antiMSLN_01 VH-CH1 - HCDR3	artificial	aa	LPRGDRDAFDI
1230.	antiMSLN_01 VL-CL - LCDR1	artificial	aa	RASQGISNYLA
1231.	antiMSLN_01 VL-CL - LCDR2	artificial	aa	AASTLQS
1232.	antiMSLN_01	artificial	aa	QQSYSTPFT

	VL-CL - Lcdr3			
1233.	antiMSLN_01 VH-CH1 - VH	artificial	aa	QVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSY FWGWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSSVTAADTAVYYCAR LPRGDRDAFDIWGQGTMTVTVSS
1234.	antiMSLN_01 VL-CL - VL	artificial	aa	EIVMTQSPSSLSASVGDRVTITCRASQGISNYLA WYQQKPGKVPKLLIYAASSTLQSGVPSRFSGSG SGTDFLTITSSLPEDFATYYCQSYSTPFTFGC GTKVEIKR
1235.	antiMSLN_01 VH-CH1 - CH1	artificial	aa	ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSLE SVVTVPSSSLGTQTYICNVNHKPSNTKVDKKV EPKSC
1236.	antiMSLN_01 VL-CL - CL	artificial	aa	TVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYP REAKVQWKVDNALQSGNSQESVTEQDSKST YSLKSTLTLSKADYEKHKVYACEVTHQGLSSP VTKSFNRGEC
1237.	antiCDH3_01 VH-CH1 - HCDR1	artificial	aa	NYWMN
1238.	antiCDH3_01 VH-CH1 - HCDR2	artificial	aa	NIAYGVAGTNYNQKFQG
1239.	antiCDH3_01 VH-CH1 - HCDR3	artificial	aa	RYFYVMDY
1240.	antiCDH3_01 VL- CL - Lcdr1	artificial	aa	RASQDISNYLN
1241.	antiCDH3_01 VL- CL - Lcdr2	artificial	aa	YTSRLHS
1242.	antiCDH3_01 VL- CL - Lcdr3	artificial	aa	VQYAQFPLT
1243.	antiCDH3_01 VH-CH1 - VH	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGAVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRRLRSDDTA VYYCATRYFYVMDYWGQGTLLTVTVSS
1244.	antiCDH3_01 VL- CL - VL	artificial	aa	EIQMTQSPSSLSASVGDRVTITCRASQDISNYLN WYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGS GTDFTLTITSSLPEDVATYYCVQYAQFPLTFGC GTKVEIKR
1245.	antiCDH3_01 VH-CH1 - CH1	artificial	aa	ASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYF PEPVTVSWNSGALTSGVHTFPAVLQSSGLYSL KSVVTVPSSSLGTQTYICNVNHKPSNTKVDKK VEPKSC
1246.	antiCDH3_01 VL- CL - CL	artificial	aa	TVAAPSVFIFPPSDEQLKSGTASVVCLLNNFYP REAKVQWKVDNALQSGNSQESVTEQDSKST YSLESTLTLSKADYEKHKVYACEVTHQGLSSP VTKSFNRGEC
1247.	antiCDH3_01 scFv x heFc(A) x antiMSLN_01 scFv	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGAVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRRLRSDDTA VYYCATRYFYVMDYWGQGTLLTVTVSSGGGGGQ GGGGQGGGGQEIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFLTITSSLPEDVATYYC

				VQYAQFPLTFGCGTKVEIKGGGGQGGGGQGGGGQDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSVSMHEALHNHYTQDSLSPGKGGGGQGGGGQGGGGQGVQLQESGPGLVKPSSETLSLTCTVSGGSISSSSYFWGWIRQPPGKCLEWIGNIYYSGSSNYPNPSLKSRTISVDTSKNQFSLKLSSVTAADTAVYYCARLPRGDRDAFDIWGQGTMTVTVSSGGGGQGGGGQGGGGQEIIVMTQSPSSLSASVGDRVTITCRASQGISNYLAWYQQKPGKVPKLLIYAASLTQSGVPSRFSGSGSGTDFLTITSSLPEDFATYYCQQSYSTPFTFGCGTKVEIK
1248.	I2L scFv x heFc(B) x I2L scFv	artificial	aa	EVQLVESGGGLVQPGGSLKLSKAASGFTFNKYAMNWRQAPGKGMWVARIRSKYNNYATYYADAVKDRFTISRDDSKNTLYLQMNNLKTEDTAVYYCVRAGNFGSSYISYFAYWGQGLTVTVSSGGGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGKFLAPGTPARFSGSLEGGKAALTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVLGGGGQGGGGQGGGGQDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPVLKSDFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYTQKSLSPGKGGGGQGGGGQGGGGQEVQLVESGGGLVQPGGSLKLSKAASGFTFNKYAMNWRQAPGKGMWVARIRSKYNNYATYYADAVKDRFTISRDDSKNTLYLQMNNLKTEDTAVYYCVRAGNFGSSYISYFAYWGQGLTVTVSSGGGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT VITICGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGKFLAPGTPARFSGSLEGGKAALTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL
1249.	antiCDH3 ₀₁ scFv x heFc(A) x I2L scFv	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTNYWMNWRQAPGQCLEWMGNIAYGVAGTNYNQKFQGRVTMTVDTSSSTAYMELSRLSDDTAVYYCATRYFYVMDYWGQGLTVTVSSGGGGQGGGGQGGGGQEIQMTQSPSSLSASVGDRVTITCRASQDISNYLNWYQQKPGKVPKLLIYTSRLHSGVPSRFSGSGSGTDFLTITSSLPEDVATYYCVQYAQFPLTFGCGTKVEIKGGGGQGGGGQGGGGQDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGSTYRCVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSVSMHEALHNHYTQDSL

				LSLSPGKGGGGQGGGGQGGGGQEVQLVESGG GLVQPGGSLKLSCAASGFTFNKYAMNWVRQA PGKMEWVARIRSKYNNYATYYADAVKDRFT ISRDDSKNTLYLQMNNLKTEDTAVYYCVRAG NFGSSYISYFAYWGQGLVTVSSGGGGQGGGG QGGGGQQT VVTQEPSLTVSPGGTVTITCGSSTG AVTSGNYPNWIKKPGQAPRGLIGGTKFLAPG TPARFSGSLEGGKAALTLSGVQPEDEAEYYCV LYYSNRWVFGSGTKLTVL
1250.	I2L scFv x heFc(B) x antiMSLN_01 scFv	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKMEWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTAV YYCVRAGNFGSSYISYFAYWGQGLVTVSSG GGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT VTITCGSSTGAVTSGNYPNWIKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTLSGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLGGGGQ GGGGQGGGGQDKTHTCPPCPAPELLGGPSVFL FPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKF NWFYVDGVEVHNAKTKPCEEQYGSTYRCVSVL TVLHQQDWLNGKEYKCKVSNKALPAPIEKTISK AKGQPREPQVYTLPPSRKEMTKNQVSLTCLVK GFYPSDIAVEWESNGQPENNYKTPPVLKSDG SFFLYSKLTVDKSRWQQGNVVFSCVMHEALH NHYTQKSLSLSPGKGGGGQGGGGQGGGGQGV QLQESGPGLVKPSSETLSLTCTVSGGSSSSSYFW GWIRQPPGKCLEWIGNIYYSGSSNYNPSLKS RVTISVDTSKNQFSLKLSVTAADTAVYYCARL PRGDRDAFDIWGQGTMTVTVSSGGGGQGGGGQ GGGQEIIVMTQSPSSLSASVGDRTVITCRASQ GISNYLAWYQQKPGKVPKLLIYAASLTQSGV PSRFSGSGSGTDFTLTISLQPEDFATYYCQ QSYSTPFTFGCGTKVEIKSGGGGQEVQLVESGG GLVQPGGSLKLSCAASGFTFNKYAMNWVRQA PGKMEWVARIRSKYNNYATYYADAVKDRFT ISRDDSKNTLYLQMNNLKTEDTAVYYCVRAG NFGSSYISYFAYWGQGLVTVSSGGGGQGGGG QGGGGQQT VVTQEPSLTVSPGGTVTITCGSSTG AVTSGNYPNWIKKPGQAPRGLIGGTKFLAPG TPARFSGSLEGGKAALTLSGVQPEDEAEYYCV LYYSNRWVFGSGTKLTVL
1251.	heFc(B)dDKTHT x antiMSLN_01 scFv x I2L scFv	artificial	aa	CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEDPEVKFNWFYVDGVEVHNAK TKPCEEQYGSTYRCVSVLTVLHQQDWLNGKEY KCKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SRKEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTPPVLKSDGSFFLYSKLTVDKSRW QQGNVVFSCVMHEALHNHYTQKSLSLSPGK GGGQVQLQESGPGLVKPSSETLSLTCTVSGG SSSYFWGWIRQPPGKCLEWIGNIYYSGSSNY NPSLKSRTISVDTSKNQFSLKLSVTAADTAV YYCARLPRGDRDAFDIWGQGTMTVTVSSGG GGQGGGGQGGGGQEIIVMTQSPSSLSASVGD RTVITCRASQGISNYLAWYQQKPGKVPKLLI YAASLTQSGVPSRFSGSGSGTDFTLTISLQPE DFATYYCQQSYSTPFTFGCGTKVEIKSGGGG QEVQLVESGGGLVQPGGSLKLSCAASGFTFN KYAMNWVRQAPGKMEWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDT AVYYCVRAGNFGSSYISYFAYWGQGLVTVSS GGGGQGGGGQGGGGQQT VVTQEPSLTVSPG GTVTITCGSSTGAVTSGNYPNWIKKPGQAPR GLIGGTKFLAPGTPARFSGSLEGGKAALTL SGVQPEDEAEYYCVLYYSNRWVFGSGTKLTVL

1252.	antiCDH3_01 VH-CH1 x heFc(A)	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIAYG VAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSGVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG K</p>
1253.	antiCDH3_01 VL- CL	artificial	aa	<p>EIQMTQSPSSLSASVGDRVTITCRASQDISNYLN WYQQKPGKVPKLLIYYTSRLHSGVPSRFSGSGS GTDFLTLISSLPEDVATYYCVQYAQFPLTFGC GTKVEIKRTVAAPSVFIFPPSDEQLKSGTASVV CLLNFPYPREAKVQWKVDNALQSGNSQESVT EQDSKDSTYSLESTLTLSKADYEEKHKVYACEV THQGLSSPVTKSFNRGEC</p>
1254.	antiCDH3_01 VH-CH1 x heFc(A) x I2L scFv	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIAYG VAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSGVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG KGGGGQGGGGQGGGGQEVQLVESGGGLVQP GGSLKLSCAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADAVKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGLTVTVSSGGGGQGGGGQGG GGQQT VVTQEPSLTVSPGGT VTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTL SGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVL</p>
1255.	I2L scFv x heFc(B) x antiMSLN_01 scFv	artificial	aa	<p>EVQLVESGGGLVQP GGSLKLSCAASGFTFNKY AMNWVRQAPGKGM EWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVTVSSG GGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTL SGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLGGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG</p>

				KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDK SRWQQGNVFCFSVMHEALHNHYTQKSLSLSP GKGGGGQGGGGQGGGGQVQLQESGPGLVK PSETLSLTCTVSGGSISSSSYFWGWIRQPPGKCL EWIGNIYYSGSSNYPNPSLKSRTISVDTSKNQFS LKLSSVTAADTAVYYCARLPRGDRDAFDIWG QGTMTVTVSSGGGGQGGGGQGGGGQEIVMTQS PSSLSASVGDRTITCRASQGISNYLAWYQQKP GKVPKLLIYAASLTQSGVPSRFSGSGSGTDFTL TISSLQPEDFATYYCQQSYPFTFGCGTKVEIK
1256.	I2L VL-CL	artificial	aa	QTVVTQEPSLTVSPGGTVTITCGSSTGAVTSGN YPNWIQKKPGQAPRGLIGGKFLAPGTPARFSG SLEGGKAALTLSGVQPEDEAEYYCVLYYSNR WVFGSGTKLTVLGQPKAAPSVTLPFSSSEELQA NKATLVCLISDFYPGAVTVAWKADSSPVKAGV ETTTPSKQSNKYAAKSYLSLTPEQWKSRSY SCQVTHEGSTVEKTVAPTECS
1257.	I2L VH-CH1 x heFc(B) x antiMSLN_01 scFv x I2L scFv	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY DAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVVSSA STKGPSVFPLAPSSKSTSGGTAALGCLVKDYFP EPVTVSWNSGALTSGVHTFPAVLQSSGLYSLES VVTVPSSSLGTQTYICNVNHKPSNTKVDKKE PKSCDKTHTCPPCPAPELGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLKSDGSFFLYSK LTVDKSRWQQGNVFCFSVMHEALHNHYTQKS LSLSPGKGGGGQVQLQESGPGLVKPSETLSLT TVSGGSISSSSYFWGWIRQPPGKCLEWIGNIYY SGSSNYPNPSLKSRTISVDTSKNQFSLKLSSVTA ADTAVYYCARLPRGDRDAFDIWGQGTMTVTVS SGGGGQGGGGQGGGGQEIVMTQSPSSLSASV GDRVTITCRASQGISNYLAWYQQKPKVPKLLI YAASLTQSGVPSRFSGSGSGTDFTLTISSLQPED FATYYCQQSYPFTFGCGTKVEIKSGGGGQE VQLVESGGGLVQPGGSLKLSCAASGFTFNKYA MNWVRQAPGKGMWVARIRSKYNNYATYYA DAVKDRFTISRDDSKNTLYLQMNNLKTEDTAV YYCVRAGNFGSSYISYFAYWGQGLTVVSSG GGQGGGGQGGGGQQTQVVTQEPSLTVSPGGTV TITCGSSTGAVTSGNYPNWIQKPGQAPRGLIG GKFLAPGTPARFSGSLEGGKAALTLSGVQPED EAEYYCVLYYSNRWVFGSGTKLTVL
1258.	I2L VH-CH1 x heFc(B) x antiMSLN_01 scFv	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY DAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVVSSA STKGPSVFPLAPSSKSTSGGTAALGCLVKDYFP EPVTVSWNSGALTSGVHTFPAVLQSSGLYSLES VVTVPSSSLGTQTYICNVNHKPSNTKVDKKE

				PKSCDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNGKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLKSDGSFFLYSK LTVDKSRWQQGNVFSVCSVMHEALHNHYTQKS LSLSPGKGGGGQGGGGQGGGGQGVQLQESGP GLVKPSETLSLTCTVSGGSISSSSYFWGWIRQPP GKCLEWIGNIYYSGSSNYPNPSLKSRTISVDTS KNQFSLKLSSVTAADTAVYYCARLPRGDRDAF DIWGQGTMTVTVSSGGGGQGGGGQGGGGQEI MTQSPSSLSASVGDRVTITCRASQGISNYLAWY QQKPGKVPKLLIYAASLTQSGVPSRFSGSGGT DFTLTISSLQPEDFATYYCQQSYSTPFTFGCGTK VEIK
1259.	antiCDH3_01 scFv x I2L scFv x heFc(A)dDKTHT	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGQ GGGGQGGGGQEIQMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYSRL HSGVPSRFSGSGGTDFLTISSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGLTVTVSSGGGGQGG GGGGGGQQTVVVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGKFL APGTPARFSGSLEGGKAALTLVGGGGCPCPAP ELGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDNLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYD TTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFS CSVMHEALHNHYTQDSLSLSPGK
1260.	antiCDH3_01 VH-CH1 x heFc(A) x antiMSLN_01 scFv x I2L scFv	artificial	aa	QVQLVQSGAEVKKPGASVKVSKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSGVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDNLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVY LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSVCSVMHEALHNHYTQDSLSLSPG KGGGGQVQLQESGPGLVKPSETLSLTCTVSGG SISSSSYFWGWIRQPPGKCLEWIGNIYYSGSSN YNPSSLKSRTISVDTSKNQFSLKLSSVTAADTA

				VYYCARLPRGDRDAFDIWGQGTMTVTVSSGGG GQGGGGQGGGGQEIIVMTQSPSSLSASVGDRVT ITCRASQGISNYLAWYQQKPGKVPKLLIYAAS LQSGVPSRFSGSGSGTDFTLTISSLQPEDFATYY CQQSYSTPFTFGCGTKVEIKSGGGGGQEVQLVES GGGLVQPGGSLKLSCAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGGQTLTVTVSSGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGKFL APGTPARFSGSLEGGKAALTLGSGVQPEDEAEY YCVLYYSNRWVFGSGTKLTVL
1261.	I2L scFv x heFc(B)	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGGQTLTVTVSSG GGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTLGSGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLGGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTTPVVKSDGSFFLYSKLTVDK SRWQQGNVFSCSVMHEALHNHYTQKSLSLSP GK
1262.	antiCDH3_01 VH-CH1 x heFc(A) x antiMSLN_01 scFv	artificial	aa	QVQLVQSGAEVKKPGASVKVCKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRLSDDTA VYYCATRYFYVMDYWGQGTLLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQDWLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG KGGGGQGGGGQGGGGQVQLQESGPGLVKPS ETLSLTCTVSGGSISSSSYFWGWIRQPPGKCLE WIGNIYSGSSNYPNPSLKSRTISVDTSKNQFSL KLSSVTAADTAVYYCARLPRGDRDAFDIWGQ GTMVTVSSGGGGQGGGGQGGGGQEIIVMTQSP SSLSASVGDRVTITCRASQGISNYLAWYQQK GKVPKLLIYAASLQSGVPSRFSGSGSGTDFTL TISSLQPEDFATYYCQQSYSTPFTFGCGTKVEIK
1263.	I2L scFv x heFc(B) x I2L scFv	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGGQGTLLTVTVSSG GGGQGGGGQGGGGQQT VVTQEPSLTVSPGGT

				<p>VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTLSGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLGGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDK SRWQQGNVFSCSVMHEALHNHYTQKSLSLSP GKGGGGQGGGGQGGGGQEVQLVESGGGLVQ PGGSLKLSCAASGFTFNKYAMNWVRQAPGKG MEWVARIRSKYNNYATYYADA VKDRFTISR DSKNTLYLQMNNLKTEDTAVYYCVRAGNFGS SYISYFAYWGQGTLVTVSSGGGGQGGGGQGG GGQQT VVTQEPSLTVSPGGT VTITCGSSTGAVT SGNYPNWIQKKPGQAPRGLIGGTKFLAPGTPA RFSGSLEGGKAALTLSGVQPEDEAEYYCVLYY SNRWVFGSGTKLTVL</p>
1264.	I2L VH-CH1 x heFc(B)	artificial	aa	<p>EVQLVESGGGLVQP GGS LKLSCAASGFTFNKY AMNWVRQAPGKGMEWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGTLVTVSSA STKGPSVFPLAPSSKSTSGGTAALGCLVKDYFP EPVTVSWNSGALTSGVHTFPAVLQSSGLYSLES VVTVPSSSLGTQTYICNVNHKPSNTKVDKKE PKSCDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLKSDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGK</p>
1265.	I2L VH-CH1 x heFc(B) x I2L scFv	artificial	aa	<p>EVQLVESGGGLVQP GGS LKLSCAASGFTFNKY AMNWVRQAPGKGMEWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGTLVTVSSA STKGPSVFPLAPSSKSTSGGTAALGCLVKDYFP EPVTVSWNSGALTSGVHTFPAVLQSSGLYSLES VVTVPSSSLGTQTYICNVNHKPSNTKVDKKE PKSCDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPVLKSDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGQGGGGQGGGGQEVQLVESGG GLVQP GGS LKLSCAASGFTFNKYAMNWVRQA PGKGMEWVARIRSKYNNYATYYADAVKDRFT ISRDDSKNTLYLQMNNLKTEDTAVYYCVRAG NFGSSYISYFAYWGQGTLVTVSSGGGGQGGGG QGGGGQQT VVTQEPSLTVSPGGT VTITCGSSTG AVTSGNYPNWIQKKPGQAPRGLIGGTKFLAPG TPARFSGSLEGGKAALTLSGVQPEDEAEYYCV</p>

				LYYSNRWVFGSGTKLTVL
1266.	antiCDH3_01 VH-CH1 x I2L scFv x heFc(A)	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSGVHTFPAVLQSSGLYSLSKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCS GGGGQEVQLVESGGGLVQPGGSLKLSCAASGF TFNKYAMNWVRQAPGKGMWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTLYLQMNLL KTEDTAVYYCVRAGNFGSSYISYFAYWGQGLT VTVSSGGGGQGGGGQGGGGQQT VVTQEPSLT VSPGGTVTITCGSSTGAVTSGNYPNWIQKKPG QAPRGLIGGTKFLAPGTPARFSGSLEGGKAALT LSGVQPEDEAEYYCVLYYSNRWVFGSGTKLT VLGGGGGDKTHTCPPCPAPELLGGPSVFLFPPK PKDTLMISRTPEVTCVVVDVSHEDPEVKFNWY VDGVEVHNAKTKPCEEQYGSTYRCVSVLTVL HQDWLNGKEYKCKVSNKALPAPIEKTISKAKG QPREPQVYTLPPSREEMTKNQVSLTCLVKGFY PSDIAVEWESNGQPENNYDTTPPVLDSDGSFFL YSDLTVDKSRWQQGNVVFSCVMHEALHNHYT QDSLSPGK
1267.	heFc(B) x antiMSLN_01 VH-CH1	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDK SRWQQGNVVFSCVMHEALHNHYTQKSLSLSP GKGGGGQVQLQESGPGLVKPSSETLSLTCTVSG GSISSSSYFWGWIRQPPGKCLEWIGNIYYSGSS NYNPSLKSRTISVDTSKNQFSLKLSVTAADT AVYYCARLPRGDRDAFDIWGQGMVTVSSAS TKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPE PVTVSWNSGALTSGVHTFPAVLQSSGLYSLES VVTVPSSSLGTQTYICNVNHKPSNTKVDKKVE PKSC
1268.	antiMSLN_01 VL-CL x I2L scFv	artificial	aa	EIVMTQSPSSLSASVGDRTITCRASQGISNYLA WYQQKPGKVPKLLIYAASTLQSGVPSRFSGSG SGTDFLTITSSLPEDFATYYCQSYSTPFTFGC GTKVEIKRTVAAPSVFIFPPSDEQLKSGTASV CLLNNFYPREAKVQWKVDNALQSGNSQESVT EQDSKDSTYSLKSTLTLSKADYEKHKVYACEV THQGLSSPVTKSFNRGECSSGGGGQEVQLVESG GGLVQPGGSLKLSCAASGF TFNKYAMNWVRQ APGKGMWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNLLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSGGGGQGGG GQGGGGQQT VVTQEPSLT VSPGGTVTITCGSST GAVTSGNYPNWIQKKPGQAPRGLIGGTKFLAP GTPARFSGSLEGGKAALT LSGVQPEDEAEYYC VLYYSNRWVFGSGTKLTVL
1269.	I2L scFv x antiCDH3_01	artificial	aa	EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY

	VH-CH1 x heFc(A)			ADAVKDRFTISRDDSKNTLYLQMNNLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVTVSSG GGGQGGGGGGGGGGQQT VVTQEP SLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTLSGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLSSGGGG QQVQLVQSGAEVKKPGASVKVSCKASGYTFT NYWMNWVRQAPGQCLEWMGNIA YGVAGTN YNQKFQGRVTMTVDTSSSTAYMELSR LRSDDT AVYYCATRYFYVMDYWGQGLTVTVSSASTKG PSVFPLAPSSKSTSGGTAALGCLVKDYFPEPVT VSWNSGALTSGVHTFPAVLQSSGLYSLKSVVT VPSSSLGTQTYICNVNHKPSNTKVDKKVEPKSC DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG K
1270.	heFc(B) x I2L scFv x antiMSLN_01 VH-CH1	artificial	aa	DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTTPPV LKSDGSFFLYSKLTVDK SRWQQGNVFSCSVMHEALHNHYTQKSLSLSP GKGGGGEVQLVESGGGLVQPGGSLKLSKAAS GFTFNKYAMNWVRQAPGKGM EWVARIRSKY NNYATYYADAVKDRFTISRDDSKNTLYLQMN NLKTEDTAVYYCVRAGNFGSSYISYFAYWGQ GTLTVTVSSGGGGGGGGGGQQT VVTQEP SLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKK PGQAPRGLIGGTKFLAPGTPARFSGSLEGGKAA LTLSGVQPEDEAEYYCVLYYSNRWVFGSGTKL TVLSSGGGGQQVQLQESGPLVKPSETLSLTCT VSGGSISSSSYFWGWIRQPPGKCLEWIGNIYYS GSSNYNPSLKSRTISVDTSKNQFSLKLSSVTA ADTAVYYCARLPRGDRDAFDIWGQGMVTVS SASTKGPSVFPLAPSSKSTSGGTAALGCLVKDY FPEPVTVSWNSGALTSGVHTFPAVLQSSGLYSL ESVVTVPSSSLGTQTYICNVNHKPSNTKVDKK VEPKSC
1271.	antiMSLN_01 VL-CL	artificial	aa	EIVMTQSPSSLSASVGDRVTITCRASQGISNYLA WYQQKPGKVPKLLIYAASTLQSGVPSRFSGSG SGTDFTLTISLQPEDFATYYCQSYSTPFTFGC GTKVEIKRTVAAPSVFIFPPSDEQLKSGTASVV CLLNNFYPREAKVQWKVDNALQSGNSQESVT EQDSKDSTYSLKSTLTLSKADYEKHKVYACEV THQGLSSPVTKSFNRGEC
1272.	antiCDH3_01 scFv x I2L scFv x scFc_clipopt x antiMSLN_01 scFv x I2L scFv	artificial	aa	QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGGQ GGGGQGGGGQEIQMTQSPSSLSASVGDRVTIT

				<p>CRASQDISNYLNWYQQKPGKVPKLLIYYTSRL HSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKSGGGGQEVQLVES GGGLVQPGGSLKLSAASGFTFNKYAMNWVR QAPGKGMWVARIRSKYNNYATYYADAVKD RFTISRDDSKNTLYLQMNNLKTEDTAVYYCVR AGNFGSSYISYFAYWGQGLTVTVSSGGGGQGG GGQGGGGQQT VVTQEPSLTVSPGGTVTITCGS STGAVTSGNYPNWIQKKPGQAPRGLIGGTKFL APGTPARFSGSLEGGKAALTLGVPPEDEAEY YCVLYYSNRWVFGSGTKLTVLGGGGCPCPAP ELGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEEPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGKGGGGQGGG GQGGGGQGGGGQGGGGQGGGGQCPPCPAP ELGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS HEEPEVKFNWYVDGVEVHNAKTKPCEEQYGS TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGGQVQLQE SGPGLVKPSETLSLTCTVSGGSISSSYFWGWIR QPPGKCLEWIGNIYYSGSSNYPNLSKSRVTISV DTSKNQFSLKLSVTAADTAVYYCARLPRGDR DAFDIWGQGTMTVTVSSGGGGQGGGGQGGGG QEVMTQSPSSLSASVGDRTITCRASQGISNYL AWYQQKPGKVPKLLIYAASLQSGVPSRFSGS GSGTDFTLTISSLQPEDFATYYCQQSYSTPFTFG CGTKVEIKSGGGGQEVQLVESGGGLVQPGGSL KLSAASGFTFNKYAMNWVRQAPGKGMWV ARIRSKYNNYATYYADAVKDRFTISRDDSKNT LYLQMNNLKTEDTAVYYCVRAGNFGSSYISYF AYWGQGLTVTVSSGGGGQGGGGQGGGGQQT VVTQEPSLTVSPGGTVTITCGSSTGAVTSGNYP NWIQKKPGQAPRGLIGGTKFLAPGTPARFSGSL EGGKAALTLGVPPEDEAEYCVLYYSNRWV FGSGTKLTVL</p>
1273.	antiCDH3_01 VH-CH1 x I2L scFv x scFc_clipopt x antiMSLN_01 VH-CH1	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYTFTN YWMNWVRQAPGQCLEWMGNIAYGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRDRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCS GGGGQEVQLVESGGGLVQPGGSLKLSAASGF TFNKYAMNWVRQAPGKGMWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTLYLQMNNL KTEDTAVYYCVRAGNFGSSYISYFAYWGQGL TVTVSSGGGGQGGGGQGGGGQQT VVTQEPSL VSPGGTVTITCGSSTGAVTSGNYPNWIQKKPG QAPRGLIGGTKFLAPGTPARFSGSLEGGKAAL</p>

				<p>LSGVQPEDEAEYYCVLYYSNRWVFGSGTKLT VLGGGGCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEEPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSDGSFFLYSKLTV DKSRWQQGNVFSCSVMHEALHNHYTQKSLSL SPGKGGGGQGGGGQGGGGQGGGGQGGGGQGG GGGQCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEEPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWE SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKS RWQQGNVFSCSVMHEALHNHYTQKSLSLSPG KGGGGQVQLQESGPGLVKPSSETLSLTCTVSGG SSSSSYFWGWIRQPPGKCLEWIGNIYYSGSSN YNPSLKSRTISVDTSKNQFSLKLSVTAADTA VYYCARLPRGDRDAFDIWGQGTMTVSSAST KGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEP VTVSWNSGALTSGVHTFPAVLQSSGLYSLESV VTPSSSLGTQTYICNVNHKPSNTKVDKKVEP KSC</p>
1274.	antiCDH3_01 scFv x I2L VH- CH1 x heFc(A)	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSKCASGYTFTN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGQ GGGGQGGGGQEIQMTQSPSSLSASVGDRVIT CRASQDISNYLNWYQQKPKGKPKLLIYYTSRL HSGVPSRFSGSGSGTDFTLTISSLQPEDVATYYC VQYAQFPLTFGCGTKVEIKGGGGQEVQLVESG GGLVQPGGSLKLSCAASGFTFNKYAMNWVRQ APGKGMWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSASTKGPSVF PLAPSSKSTSGGTAALGCLVKDYFPEPVTVSW NSGALTSGVHTFPAVLQSSGLYSLESVTVPS SLGTQTYICNVNHKPSNTKVDKKVEPKSCKT HTPPCPAPELLGGPSVFLFPPKPKDTLMISRT P ETCVVVDVSHEDPEVKFNWYVDGVEVHNA KTKPCEEQYGSTYRCVSVLTVLHQDWLNGKE YKCKVSNKALPAPIEKTISKAKGQPREPQVYTL PPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTTPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG K</p>
1275.	heFc(B) x antiMSLN_01 scFv x I2L VH- CH1	artificial	aa	<p>DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW E SNGQPENNYKTTTPVLDSDGSFFLYSKLTVDK SRWQQGNVFSCSVMHEALHNHYTQKSLSLSP GKGGGGQVQLQESGPGLVKPSSETLSLTCTVSG GSISSSSYFWGWIRQPPGKCLEWIGNIYYSGSS</p>

				<p>NYNPSLKS RV TISVDT SKNQFSLKLSSVTAADT AVYYCARLPRGDRDAFDIWGQGTMTVTVSSGG GGQGGGGQGGGGQEI VMTQSPSSLSASVGDR VTITCRASQGISNYLAWYQQKPGKVPKLLIYA ASTLQSGVPSRFSGSGSGTDFTLTISLQPEDFA TYYCQQSYS TPFTFGCGTKVEIKGGGGQEVQL VESGGGLVQPGGSLKLSCAASGFTFNKYAMN WVRQAPGKGM EWVARIRSKYNNYATYYADA VKDRFTISRDDSKNTLYLQMNNLKTEDTAVYY CVRAGNFGSSYISYFAYWGQGLTVTVSSASTK GPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPV TVSWNSGALTSGVHTFPAVLQSSGLYSLESVV TVPSSSLGTQTYICNVNHKPSNTKVDK KVEPKS C</p>
1276.	<p>antiCDH3_01 scFv x 12L VH- CH1 x scFc_clipopt x antiMSLN_01 scFv x 12L VH- CH1</p>	artificial	aa	<p>QVQLVQSGAEVKKPGASVKV SCKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSGGGGQ GGGGQGGGGQEI QMTQSPSSLSASVGDRVTIT CRASQDISNYLNWYQQKPGKVPKLLIYYSRL HSGVPSRFSGSGSGTDFTLTISLQPEDVATYYC VQY AQFPLTFGCGTKVEIKGGGGQEVQLVESG GGLVQPGGSLKLSCAASGFTFNKYAMNWVRQ APGKGM EWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSASTKGPSVF PLAPSSKSTSGGTAALGCLVKDYFPEPVTVSW NSGALTSGVHTFPAVLQSSGLYSLESVVTVPSS SLGTQTYICNVNHKPSNTKVDK KVEPKSCGGG GCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPE VTCVVVDVSHEEPEVKFNWYVDGVEVHNAKT KPCEEQYGSTYRCVSVLTVLHQDWLNGKEYK CKVSNKALPAPIEKTISKAKGQPREPQVYTLPP SREEMTKNQVSLTCLVKGFYPSDIAVEWESNG QPENNYKTTTPVLDSDGSFFLYSKLTVDKSRW QQGNVFSCSVMHEALHNHYTQKSLSLSPGKGG GGGQGGGGQGGGGQGGGGQGGGGQGGGGQ CPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSHEEPEVKFNWYVDGVEVHNAKT PCEEQYGSTYRCVSVLTVLHQDWLNGKEYK KVS NKALPAPIEKTISKAKGQPREPQVYTLPPS REEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ PENNYKTTTPVLDSDGSFFLYSKLTVDKSRWQ QQGNVFSCSVMHEALHNHYTQKSLSLSPGKGG GGQVQLQESGPGLVKPSETLSLTCTVSGGSISS SYFWGWIRQPPGKCLEWIGNIYYSGSSNYNPSL KSRVTISVDT SKNQFSLKLSSVTAADTAVYYC ARLPRGDRDAFDIWGQGTMTVTVSSGGGGQGG GGQGGGGQEI VMTQSPSSLSASVGDRVTITCR ASQGISNYLAWYQQKPGKVPKLLIYA ASTLQSG VPSRFSGSGSGTDFTLTISLQPEDFATYYCQ SYSTPFTFGCGTKVEIKGGGGQEVQLVESGGG LVQPGGSLKLSCAASGFTFNKYAMNWVRQAP GKGM EWVARIRSKYNNYATYYADAVKDRFTI SRDDSKNTLYLQMNNLKTEDTAVYYCVRAGN FGSSYISYFAYWGQGLTVTVSSASTKGPSVFPL</p>

				APSSKSTSGGTAALGCLVKDYFPEPVTVSWNS GALTSGVHTFPAVLQSSGLYSLESVVTVPSSSL GTQTYICNVNHKPSNTKVKDKKVEPKSC
1277.	I2L scFv x heFc(B) x antiMSLN_01 scFv x I2L scFv	artificial	aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDA VYYCVRAGNFGSSYISYFAYWGQGLVTVSSG GGGQGGGGQGGGGQQT VVTQEPLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTLSGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLGGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTPPVLKSDGSFFLYSKLTVDK SRWQQGNVFSCVMHEALHNHYTQKSLSLSP GKGGGGQVQLQESGPGLVKPSSETLSLTCTVSG GSISSSYFWGWIRQPPGKCLEWIGNIYYSGSS NYNPSLKSRTISVDTSKNQFSLKLSVTAADT AVYYCARLPRGDRDAFDIWGQGMVTVSSGG GGQGGGGQGGGGQEI VMTQSPSSLSASVGD VTITCRASQGISNYLAWYQKPKGPKLLIYA ASTLQSGVPSRFSGSGSGTDFLTISLQPEDFA TYYCQQSYSTPFTFGCGTKVEIKSGGGGQEVQ LVESGGGLVQPGGSLKLSAASGFTFNKYAMN WVRQAPGKGMWVARIRSKYNNYATYYADA VKDRFTISRDDSKNTLYLQMNNLKTEDAVYY CVRAGNFGSSYISYFAYWGQGLVTVSSGGGG QGGGGQGGGGQQT VVTQEPLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLIGGT KFLAPGTPARFSGSLEGGKAALTLSGVQPEDEA EYYCVLYYSNRWVFGSGTKLTVL
1278.	I2L scFv x heFc(B) x antiMSLN_01 VH-CH1	artificial	aa	EVQLVESGGGLVQPGGSLKLSAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMNNLKTEDA VYYCVRAGNFGSSYISYFAYWGQGLVTVSSG GGGQGGGGQGGGGQQT VVTQEPLTVSPGGT VTITCGSSTGAVTSGNYPNWIQKKPGQAPRGLI GGTKFLAPGTPARFSGSLEGGKAALTLSGVQPE DEAEYYCVLYYSNRWVFGSGTKLTVLGGGGG DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMIS RTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH NAKTKPCEEQYGSTYRCVSVLTVLHQDWLNG KEYKCKVSNKALPAPIEKTISKAKGQPREPQVY TLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEW ESNGQPENNYKTPPVLKSDGSFFLYSKLTVDK SRWQQGNVFSCVMHEALHNHYTQKSLSLSP GKGGGGQGGGGQGGGGQVQLQESGPGLVK PSETLSLTCTVSGGSISSSYFWGWIRQPPGKCL EWIGNIYYSGSSNYNPSLKSRTISVDTSKNQFS LKLSSVTAADTAVYYCARLPRGDRDAFDIWG QGMVTVSSASTKGPSVFPLAPSSKSTSGGTA ALGCLVKDYFPEPVTVSWNSGALTSGVHTFPAV LQSSGLYSLESVVTVPSSSLGTQTYICNVNHK

				SNTKVDKKVEPKSC
1279.	antiCDH3_01 VH-CH1 x heFc(A) x antiMSLN_01 VH-CH1	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSGVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCD KTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISR TPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN AKTKPCEEQYGSTYRCVSVLTVLHQD WLNGK EYKCKVSNKALPAPIEKTISKAKGQPREPQVYT LPPSREEMTKNQVSLTCLVKGFYPSDIAVEWES NGQPENNYDTTPPVLDSDGSFFLYSDLTVDKS RWQQGNVFSCSVMHEALHNHYTQDSLSLSPG KGGGGQGGGGQGGGGQGVQLQESGPGLVKPS ETLSLTCTVSGGSISSSSYFWGWRQPPGKCLE WIGNIYSGSSNYPNPSLKSRTISVDTSKNQFSL KLSSVTAADTAVYYCARLPRGDRDAFDIWGQ GTMVTVSSASTKGPSVFPLAPSSKSTSGGTAAL GCLVKDYFPEPVTVSWNSGALTSGVHTFPAVL QSSGLYSLESVTVPSSSLGTQTYICNVNHKPS NTKVDKKVEPKSC</p>
1280.	I2L VH-CH1 x heFc(B) x I2L VH-CH1	artificial	aa	<p>EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGMWVARIRSKYNNYATYY ADAVKDRFTISRDDSKNTLYLQMN NLKTEDTA VYYCVRAGNFGSSYISYFAYWGQGLTVTVSSA STKGPSVFPLAPSSKSTSGGTAALGCLVKDYFP EPVTVSWNSGALTSGVHTFPAVLQSSGLYSLES VTVPSSSLGTQTYICNVNHKPSNTKVDKKVE PKSCDKTHTCPPCPAPELLGGPSVFLFPPKPKD TLMISRTPPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNGKEYKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTPPV LKSDGSFFLYSK LTVDKSRWQQGNVFSCSVMHEALHNHYTQKS LSLSPGKGGGGQGGGGQGGGGQEVQLVESGG GLVQPGGSLKLSCAASGFTFNKYAMNWVRQA PGKGMWVARIRSKYNNYATYYADAVKDRFT ISRDDSKNTLYLQMN NLKTEDTAVYYCVRAG NFGSSYISYFAYWGQGLTVTVSSASTKGPSVFP LAPSSKSTSGGTAALGCLVKDYFPEPVTVSWN SGALTSGVHTFPAVLQSSGLYSLESVTVPSSS LGTQTYICNVNHKPSNTKVDKKVEPKSC</p>
1281.	antiCDH3_01 VH-CH1 x I2L VH-CH1 x heFc(A)	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSR LRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSGVHTFPAVLQSSGLYSLKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCS GGGGQEVQLVESGGGLVQPGGSLKLSCAASGF TFNKYAMNWVRQAPGKGMWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTLYLQMN NL KTEDTAVYYCVRAGNFGSSYISYFAYWGQGL</p>

				<p>VTVSSASTKGPSVFPLAPSSKSTSGGTAALGCL VKDYFPEPVTVSWNSGALTSQVHTFPAVLQSS GLYSLESVVTVPSSSLGTQTYICNVNHKPSNTK VDKKVEPKSCDKTHTCPPCPAPELLGGPSVFLF PPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPDI AVEWESNGQPENNYDTTPPVLDSDGSF FLYSDLTVDKSRWQQGNV FSCSV MHEALHNH YTQDSLSPGK</p>
1282.	antiMSLN_01 VL-CL x I2L VH- CH1	artificial	aa	<p>EIVMTQSPSSLSASVGDRTITCRASQGISNYLA WYQQKPGKVPKLLIYAASLTQSGVPSRFSGSG SGTDFLTISLQPEDFATYYCQSYSTPFTFGC GTKVEIKRTVAAPSVFIFPPSDEQLKSGTASV CLLNFPYPREAKVQWKVDNALQSGNSQESVT EQDSKDSTYSLKSTLTLKADYEEKHKVYACEV THQGLSSPVTKSFNRGECSSGGGGQEVQLVESG GGLVQPGGSLKLSAASGFTFNKYAMNWVRQ APGKGMEWVARIRSKYNNYATYYADAVKDR FTISRDDSKNTLYLQMNNLKTEDTAVYYCVRA GNFGSSYISYFAYWGQGLTVTVSSASTKGPSVF PLAPSSKSTSGGTAALGCLVKDYFPEPVTVSW NSGALTSQVHTFPAVLQSSGLYSLESVVTVPSS SLGTQTYICNVNHKPSNTKVDKKVEPKSC</p>
1283.	antiCDH3_01 VH-CH1 x I2L VH-CH1 x scFc_clipopt x antiMSLN_01 VH-CH1	artificial	aa	<p>QVQLVQSGAEVKKPGASVKVSCKASGYFTFN YWMNWVRQAPGQCLEWMGNIA YGVAGTNY NQKFQGRVTMTVDTSSSTAYMELSRLRSDDTA VYYCATRYFYVMDYWGQGLTVTVSSASTKGP SVFPLAPSSKSTSGGTAALGCLVKDYFPEPVTV SWNSGALTSQVHTFPAVLQSSGLYSLSKSVTV PSSSLGTQTYICNVNHKPSNTKVDKKVEPKSCS GGGGQEVQLVESGGGLVQPGGSLKLSAASGF TFNKYAMNWVRQAPGKGMEWVARIRSKYNN YATYYADAVKDRFTISRDDSKNTLYLQMNNL KTEDTAVYYCVRAGNFGSSYISYFAYWGQGL TVTVSSASTKGPSVFPLAPSSKSTSGGTAALGCL VKDYFPEPVTVSWNSGALTSQVHTFPAVLQSS GLYSLESVVTVPSSSLGTQTYICNVNHKPSNTK VDKKVEPKSCGGGGCPPCPAPELLGGPSVFLF PPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFN WYVDGVEVHNAKTKPCEEQYGSTYRCVSVLT VLHQDWLNGKEYKCKVSNKALPAPIEKTISKA KGQPREPQVYTLPPSREEMTKNQVSLTCLVKG FYPDI AVEWESNGQPENNYKTTTPPVLDSDGSF FLYSKLTVDKSRWQQGNV FSCSV MHEALHNH YTQKSLSPGKGGGGQGGGGQGGGGQGGGG QGGGGQGGGGQCPPCPAPELLGGPSVFLFPPK KDTLMISRTPEVTCVVVDVSHEDPEVKFNWYV DGVEVHNAKTKPCEEQYGSTYRCVSVLTVLH QDWLNGKEYKCKVSNKALPAPIEKTISKAKGQ PREPQVYTLPPSREEMTKNQVSLTCLVKGFYPS DIAVEWESNGQPENNYKTTTPPVLDSDGSFFLYS KLTVDKSRWQQGNV FSCSV MHEALHNHYTQK SLSPGKGGGGQVQLQESGPGLVKPSETLSLT CTVSGGSISSSYFWGWIRQPPGKCLEWIGNIY</p>

				YSGSSNYNPSLKSRTISVDTSKNQFSLKLSSTV AADTAVYYCARLPRGDRDAFDIWGGTMVTV SSASTKGPSVFPLAPSSKSTSGGTAALGCLVKD YFPEPVTVSWNSGALTSGVHTFPAVLQSSGLYS LESVVTVPSSSLGTQTYICNVNHKPSNTKVDKK VEPKSC
1284.	CH3-G8A_6- B12x I2Cx scFc	artificial	aa	EVQLLESGGGLVQPGGSLRLSCAASGFSFSSYPI NWVRQAPGKGLEWVGVIWTGGGTNYASSVK GRFTISRDNKNTVYLQMNSLRAEDTAVYYCA KSRGVYDFDGRGAMDYWGQGLTVTVSSGGG GSGGGGSGGGGSDIVMTQSPDSLAVSLGERATI NCKSSQSLLYSSNQKNYFAWYQQKPGPPKLL IYWASTRESGVPDRFSGSGSGTDFLTISLQAE DVAVYYCQQYYSYPYTFGQGTKLEIKSGGGGS EVQLVESGGGLVQPGGSLKLSCAASGFTFNKY AMNWVRQAPGKGLEWVARIRSKYNNYATYY ADSVKDRFTISRDDSKNTAYLQMNNLKTEDTA VYYCVRHGNFGNSYISYWAYWGQGLTVTVSS GGGGSGGGGSGGGGSQTVVTQEPSLTVSPGGT VTLTCGSSTGAVTSGNYPNWVQQKPGQAPRG LIGGTKFLAPGTPARFSGSLLGGKAALTLGTVQ PEDEAEYYCVLWYSNRWVFGGGTKLTVLGGG GDKTHTCPPCPAPELLGGPSVFLFPPKPKDTLM ISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEV HNAKTKPCEEQYGSTYRCVSVLTVLHQDWLN GKEYCKCKVSNKALPAPIEKTISKAKGQPREPQV YTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE WESNGQPENNYKTTTPVLDSDGSFFLYSKLTV DKSRWQQGNVFCFSVMHEALHNHYTQKLSL SPGKGGGGSGGGGSGGGGSGGGGSGGGGSGG GGSDKTHTCPPCPAPELLGGPSVFLFPPKPKDT LMISRTPEVTCVVVDVSHEDPEVKFNWYVDG VEVHNAKTKPCEEQYGSTYRCVSVLTVLHQD WLNKEYCKCKVSNKALPAPIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSDGSFFLYSK LTVDKSRWQQGNVFCFSVMHEALHNHYTQKS LSLSPGK
1285.	MS 5-F11 x I2C x scFc	artificial	aa	QVQLVESGGGLVKPGGSLRLSCAASGFTFSDY YMTWIRQAPGKGLEWLSYISSSGSTIYYADSV KGRFTISRDNKNSLFLQMNSLRAEDTAVYYC ARDRNSHFDYWGQGLTVTVSSGGGGSGGGGS GGGGSDIQMTQSPSSVSASVGDRTITCRASQG INTWLAWYQQKPGKAPKLLIYGASGLQSGVPS RFGSGSGTDFLTISLQPEDFATYYCQQAQSF PRTFGQGTKVEIKSGGGGSEVQLVESGGGLVQ PGGSLKLSCAASGFTFNKYAMNWVRQAPGK LEWVARIRSKYNNYATYYADSVKDRFTISRDD SKNTAYLQMNNLKTEDTAVYYCVRHGNFGNS YISYWAYWGQGLTVTVSSGGGGSGGGGSGGG GSQTVVTQEPSLTVSPGGTVTLTCGSSTGAVTS GNYPNWVQQKPGQAPRGLIGGTKFLAPGTPAR FSGSLLGGKAALTLGTVQPEDEAEYYCVLWYS NRWVFGGGTKLTVLGGGGDKTHTCPPCPAPELL GGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SHEDPEVKFNWYVDGVEVHNAKTKPCEEQYGS

				TYRCVSVLTVLHQDWLNGKEYKCKVSNKALP APIEKTISKAKGQPREPQVYTLPPSREEMTKNQ VSLTCLVKGFYPSDIAVEWESNGQPENNYKTT PPVLDSDGSFFLYSKLTVDKSRWQQGNVFS VMHEALHNHYTQKSLSLSPGKGGGSGGGGS GGGSGGGSGGGSGGGSGGGSDKTHTCPAP ELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSHEDPEVKFNWYVDGVEVHNAKTKPCEEQY GSTYRCVSVLTVLHQDWLNGKEYKCKVSNKA LPAPIEKTISKAKGQPREPQVYTLPPSREEMTK NQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS CSVMHEALHNHYTQKSLSLSPGK
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Claims

1. A molecule comprising at least two polypeptide chains, wherein the molecule comprises
 - (i.) a first binding domain which binds to a first target cell surface antigen (TAA1),
 - (ii.) a second binding domain which binds to an extracellular epitope of the human and/or the Macaca CD3 chain,
 - (iii.) a third binding domain which binds to a second target cell surface antigen (TAA2), and
 - (iv.) a fourth binding domain which binds to an extracellular epitope of the human and/or the Macaca CD3 chain,

wherein the first binding domain and the second binding domain form a first bispecific entity, and the third and the fourth binding domain form a second bispecific entity, and wherein the molecule further comprises a spacer entity selected from

- (1.) a dimerizing domain selected from
 - (a.) an Fc domain comprising a first and a second polypeptide monomer comprising a hinge, a CH2 domain and a CH3 domain, respectively, wherein the first and second polypeptide monomer form a heterodimer; wherein the heterodimer is formed by
 - charged pair mutations selected from (i.) D399K, K409D, K392D, and E356K, (ii.) D399K, K409D, K392D, E357K, K370D, and E356K, (iii.) D399K, K409D, K392D, E356K and K439D, (iv.) D399K, K409D, and K392D, (v.) D399K, K409D, K392D, E357K and E370K, (vi.) D399K, K409D, K392D, E357K, K370E and K360E, (vii.) D399K, K409D, K392D, E357K, K370E, E356K, and K439E, and (viii.) D399K, K409D, K392D, E357K, K370E, K360E, E356K, and K439D, preferably comprising a K392D, K409D and/or K439D mutation in the CH3 domain of the first polypeptide monomer and comprising a E356K and/or D399K mutation in the CH3 domain of the second polypeptide monomer, wherein the positions are according to EU numbering; or
 - knobs-into-holes mutations comprising preferably a T366S, L368A and Y407V mutation in the first polypeptide monomer and a T366W mutation in the second monomer, wherein the positions are according to EU numbering;

(b.) a human serum albumin (HSA) domain comprising a first and a second polypeptide monomer, wherein the first and the second polypeptide monomer correspond to an HSA subdomain, respectively, wherein the first and second polypeptide monomer form a native HSA-like heterodimer; and

(c.) a Fab comprising a first and a second polypeptide monomer, wherein preferably the first polypeptide monomer comprises a VL and CL domain and the second polypeptide monomer comprises a VH and CH1 domain, wherein the CL and CH1 domains are linked by a disulfide bridge;

wherein the dimerizing domain comprises two N-termini and two C-termini, respectively, whereof at least one N-terminus and one C-terminus, respectively, is linked to a bispecific entity, wherein any of the first, second, third and fourth domain can be selected from any form of binding domain, preferably selected from Fab and single chain domain, the single chain domain preferably selected from single chain Fv (scFv) and scFab;

(2.) a single chain domain selected from ubiquitin , beta 2 microglobulin , , VH-only domain , PSI domain from Met-receptor, Fibronectin type III domain from tenascin , Granulocyte-macrophage colony-stimulating factor (GM-CSF) , interleukin-4 , , Interleukin-2 , PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1) , Tim-3 (AS 24-130), MiniSOG , a programmed cell death protein 1 (PD1) domain, human serum albumin (HSA), or a single chain Fc (scFc) domain comprising two polypeptide monomers comprising each a hinge, a CH2 and a CH3 domain a hinge and a further CH2 and a CH3 domain, wherein said two polypeptide monomers are fused to each other via a peptide linker, wherein the single chain domain comprises one N-terminus and one C-terminus, which are, respectively, linked to a bispecific entity, wherein at least one of the first, second, third and fourth binding domain is a two-chained Fab, and any of the remaining at least three binding domains can be selected from any form of binding domain, preferably selected from Fab and single chain domain, the single chain domain preferably selected from scFv and scFab;

wherein the distance between the C alpha atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus of the spacer entity are spaced apart by at least 30 Å, wherein the spacer entity spaces apart the first and the second bispecific entity by at least a distance of about 50 Å, wherein the indicated distance is preferably understood as the distance between centers of mass (i.) of the first and the third binding domain or (ii.) the first and the second bispecific entity, and which spacer entity is positioned between the first and the second bispecific entity.

2. The molecule according to claim 1 which is an antigen-binding molecule, preferably a bispecific antigen-binding molecule, more preferably a multichain multitargeting bispecific antigen-binding molecule.

3. The antigen-binding molecule of claim 2, wherein when the spacer is a single chain domain, the arrangement of the binding domains in an amino to carboxyl order is selected from the group consisting of
(i.) first and second domain, spacer, third and fourth domain

(ii.) first and second domain, spacer, fourth and third domain

(iii.) second and the first domain, spacer, third and fourth domain, and

(iv.) second and first domain, spacer, fourth and third domain.

4. The antigen-binding molecule according to any of the preceding claims, wherein when the spacer is a single chain domain, the arrangement of the binding domains in an amino to carboxyl order is selected from the group consisting of

(i.) first domain in the format of Fab, second domain in the format of a single chain domain, preferably scFv, spacer, third domain in the format of Fab and fourth domain in the format of a single chain domain, preferably scFv;

(ii.) first domain in the format of Fab, second domain in the format of Fab, spacer, third domain in the format of Fab and fourth domain in the format of Fab;

(iii.) first domain in the format of a single chain domain, preferably scFv, second domain in the format of Fab, spacer, third in the format of scFv and fourth domain in the format of Fab;

(iv.) first domain in the format of a single chain domain, preferably scFv, second domain in the format of scFv, spacer, third in the format of scFv and fourth domain in the format of Fab;

(v.) first domain in the format of a single chain domain, preferably scFv, second domain in the format of a single chain domain, preferably scFv, spacer, third in the format of Fab and fourth domain in the format of a single chain domain, preferably scFv;

(vi.) first domain in the format of Fab, second domain in the format of a single chain domain, preferably scFv, spacer, third in the format of a single chain domain, preferably scFv and fourth domain in the format of a single chain domain, preferably scFv; and

(vii.) first domain in the format of a single chain domain, preferably scFv, second domain in the format of Fab, spacer, third in the format of a single chain domain, preferably scFv and fourth domain in the format of a single chain domain, preferably scFv,

wherein each scFv comprises in an amino to carboxyl order VH, linker and VL or VL, linker and VH, preferably VH, linker and VL.

5. The antigen-binding molecule according to any of the preceding claims, wherein when the spacer is a dimerizing domain, the arrangement of the binding domains in an amino to carboxyl order is selected from the group consisting of

(i.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, second domain in the format of scFv, first polypeptide monomer of the spacer dimerizing domain, and a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain, and the fourth domain in the format of a scFv;

(ii.) first domain in the format of Fab, second domain in the format of Fab, spacer comprising first and second polypeptide monomer of the spacer dimerizing domain, third domain in the format of Fab and fourth domain in the format of Fab;

(iii.) a first chain comprising the second domain in the format of scFv, the VH and CH1 of the first domain, forming a Fab together with the second chain, first polypeptide monomer of the spacer dimerizing domain, a second chain comprising the VL and CL of the first domain, a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain, and the fourth domain in the format of a scFv;

(iv.) a first chain comprising the second domain in the format of scFv, the VH and CH1 of the first domain, forming a Fab together with the second chain, first polypeptide monomer of the spacer dimerizing domain, a second chain comprising the VL and CL of the first domain, a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the fourth domain in the format of a scFv, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain;

(v.) a first chain comprising the second domain in the format of scFv, the VH and CH1 of the first domain, forming a Fab together with the second chain, first polypeptide monomer of the spacer dimerizing domain,

a second chain comprising the VL and CL of the first domain, a third chain comprising the second polypeptide monomer of the spacer dimerizing domain, the fourth domain in the format of a scFv, the third domain comprising the VH and CH1 of the third domain, forming a Fab together with the VL and CL of the third domain on the fourth chain, and a fourth chain comprising the VL and CL of the third domain;

wherein each scFv comprises in an amino to carboxyl order VH, linker and VL or VL, linker and VH, preferably VH, linker and VL.

6. The antigen-binding molecule according to any of the preceding claims, wherein when the spacer is a dimerizing domain, the arrangement of the binding domains is in an amino to carboxyl order and is selected from the group consisting of

(i.) a first chain comprising the first domain in the format of scFv, first polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv and a second chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, fourth domain in the format of scFv;

(ii.) a first chain comprising the first domain in the format of scFv, first polypeptide monomer of the spacer dimerizing domain, second domain in the format of scFv and a second chain comprising the third domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, fourth domain in the format of scFv;

(iii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv, and a third chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, fourth domain in the format of scFv;

(iv.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, second domain in the format of scFv, and a third chain comprising the fourth domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv;

(v.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the

spacer dimerizing domain, and a third chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv, and fourth domain in the format of scFv;

(vi.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, third domain in the format of scFv, and fourth domain in the format of scFv, and a third chain comprising the second domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain;

(vii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, VH and CH1 of the third domain forming a Fab together with the third chain, a third chain comprising the VL and CL of the third domain, a fourth chain comprising the VH and CH1 of the second domain, forming a Fab together with the fifth chain, second polypeptide monomer of the spacer dimerizing domain, VH and CH1 of the third domain forming a Fab together with the sixth chain, a fifth chain comprising the VL and CL of the second domain, and a sixth chain comprising the VL and CL of the fourth domain;

(viii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, second domain in the format of scFv, a third chain comprising the fourth domain in the format of scFv, second polypeptide monomer of the spacer dimerizing domain, VH and CH1 of the third domain, forming a Fab together with the fourth chain, and a fourth chain comprising the VL and CL of the third domain;

(ix.) a first chain comprising the first domain in the format of a scFv, the first polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the third chain, the second polypeptide monomer of the spacer dimerizing domain, the VH and CH1 of the fourth domain, forming a Fab together with the fourth chain, a third chain comprising the VL and CL of the second domain, and a fourth chain comprising the VL and CL of the fourth domain;

(x.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the fourth chain, second polypeptide monomer of

the spacer dimerizing domain, the fourth domain in the format of a scFv, and a fourth chain comprising the VL and CL of the second domain;

(xi.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, the second domain in the format of scFv, a third chain comprising the VH and CH1 of the fourth domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing domain, the third domain in the format of a scFv, and a fourth chain comprising the VL and CL of the fourth domain;

(xii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, the fourth domain in the format of a scFv, and a fourth chain comprising the VL and CL of the second domain;

(xiii.) a first chain comprising the VL and CL of the first domain, a second chain comprising the VH and CH1 of the first domain, forming a Fab together with the first chain, first polypeptide monomer of the spacer dimerizing domain, the third domain in the format of scFv, the fourth domain in the format of a scFv, a third chain comprising the VH and CH1 of the second domain, forming a Fab together with the fourth chain, second polypeptide monomer of the spacer dimerizing domain, and a fourth chain comprising the VL and CL of the second domain;

wherein each scFv comprises in a N to C orientation VH, linker and VL or VL, linker and VH, preferably VH, linker and VL.

7. The antigen-binding molecule according to any of the preceding claims, wherein the spacer entity is a globular protein, wherein the distance between the C alpha atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus are spaced apart by at least 20 Å, preferably at least 30 Å, more preferably at least 50 Å, in order to effectively space apart the first and the second bispecific entity by preferably at least 50 Å.

8. The antigen-binding molecule according to any of the preceding claims, wherein said spacer entity which sufficiently spaces apart the first and the second bispecific entity and is single-chained is selected from a group consisting of ubiquitin, beta 2 microglobulin, SAND domain, Green fluorescent protein (GFP), VHH antibody lama domain, PSI domain from Met-receptor, Fibronectin type III domain from

tenascin , Granulocyte-macrophage colony-stimulating factor (GM-CSF) , interleukin-4 , CD137L Ectodomain , Interleukin-2 , PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1) , Tim-3 (AS 24-130), MiniSOG, a programmed cell death protein 1 (PD1) domain, human serum albumin (HSA) or a derivate of any of the foregoing spacer entities, a multimer of a rigid linker, and a Fc domain or dimer or trimer thereof, each Fc domain comprising two polypeptide monomers comprising each a hinge, a CH2 and a CH3 domain a hinge and a further CH2 and a CH3 domain, wherein said two polypeptide monomers are fused to each other via a peptide linker or wherein the two polypeptide monomers are linked together by non-covalent CH3-CH3 interactions and/or covalent disulfide bonds to form a heterodimer.

9. The antigen-binding molecule according to any of the preceding claims, wherein said spacer entity when single chained is at least one Fc domain, preferably one domain or two or three covalently linked domains, which or each of which comprises in an amino to carboxyl order:

hinge-CH2-CH3-linker-hinge-CH2-CH3.

10. The antigen-binding molecule according to any of the preceding claims, wherein each of said polypeptide monomers in the spacer entity has an amino acid sequence that is at least 90% identical to a sequence selected from the group consisting of: SEQ ID NO: 17-24, wherein preferably each of said polypeptide monomers has an amino acid sequence selected from SEQ ID NO: 17-24.

11. The antigen-binding molecule according to any of the preceding claims, wherein the CH2 domains in the spacer comprises an intra domain cysteine disulfide bridge.

12. The antigen-binding molecule according to any of the preceding claims, wherein the spacer entity comprises an amino acid sequence selected the group consisting of SEQ ID NO: 13 and 15 to 16 and 25 to 34 ubiquitin (SEQ ID NO: 1081), beta 2 microglobulin (SEQ ID NO: 1083), SAND domain (SEQ ID NO: 1084), Green fluorescent protein (GFP) (SEQ ID NO: 1085), VHH antibody lama domain (SEQ ID NO: 1086), PSI domain from Met-receptor (SEQ ID NO: 1087), Fibronectin type III domain from tenascin (SEQ ID NO: 1088), Granulocyte-macrophage colony-stimulating factor (GM-CSF) (SEQ ID NO: 1089), interleukin-4 (SEQ ID NO: 1090), CD137L Ectodomain (SEQ ID NO: 1091), Interleukin-2 (SEQ ID NO: 1092), PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1) (SEQ ID NO: 1093), Tim-3 (AS 24-130) (SEQ ID NO: 1094), MiniSOG (SEQ ID NO: 1095), a programmed cell death protein 1 (PD1) domain (SEQ ID NO: 16), human serum albumin (has, SEQ ID NO: 15) or an amino acid with at least 90%, preferably 95% or even 98% sequence identity thereof, preferably scFc (SEQ ID NO: 25).

13. The antigen-binding molecule according to any of the preceding claims, wherein the first peptide monomer of the first peptide chain in the dimerizing spacer is SEQ ID NO 35 and the second peptide monomer of the second peptide chain in the dimerizing spacer is SEQ ID NO 36, wherein the two peptide monomers preferably form a heterodimer.

14. The antigen-binding molecule according to any of the preceding claims, wherein the antigen-binding molecule is characterized by

(i) the first and third domain comprise two antibody-derived variable domains and the second and the fourth domain comprises two antibody-derived variable domains;

(ii) the first and third domain comprise one antibody-derived variable domain and the second and the fourth domain comprises two antibody-derived variable domains;

(iii) the first and third domain comprise two antibody-derived variable domains and the second and the fourth domain comprises one antibody-derived variable domain; or

(iv) the first domain comprises one antibody-derived variable domain and the third domain comprises one antibody-derived variable domain.

15. The antigen-binding molecule according to any of the preceding claims 1 to 7, wherein the antigen-binding molecule comprises two polypeptide chains, wherein

the first polypeptide chain comprises a VH of the first domain, a VH second domain, the first polypeptide monomer comprising preferably a hinge, a CH2 and a CH3 domain, a VH of the third domain, and a VH of the fourth domain; and

the second polypeptide chain comprises a VL of the first domain, a VL second domain, the first polypeptide monomer comprising preferably a hinge, a CH2 and a CH3 domain, a VL of the third domain, and a VL of the fourth domain,

wherein preferably the first and second polypeptide monomer form a heterodimer, thereby connecting the first and the second polypeptide chain.

16. The antigen-binding molecule according to any of the preceding claims, wherein the antigen-binding molecule, wherein the first, second, third and fourth binding domain each comprise in an amino to carboxyl order a VH domain and a VL domain, wherein the VH and VL within each domain is connected by a peptide linker, preferably a flexible linker which comprises serine, glutamine and/or glycine as amino

acid building blocks, preferably only serine (Ser, S) or glutamine (Gln, Q) and glycine (Gly, G), more preferably (G4S)_n or (G4Q)_n, even more preferably SEQ ID NO: 1 or 3.

17. The antigen-binding molecule according to any of the preceding claims, wherein the peptide linker comprises or consists of S(G4X)_n and (G4X)_n, wherein X is selected from the group consisting of Q, T, N, C, G, A, V, I, L, and M, and wherein n is an integer selected from integers 1 to 20, preferably wherein n is 1, 2, 3, 4, 5 or 6, preferably wherein X is Q, wherein preferably the peptide linker is (G4X)_n, n is 3, and X is Q.

18. The antigen-binding molecule according to any of the preceding claims, wherein the peptide linker between the first binding domain and the second binding domain and the third binding domain and the fourth binding domain is preferably a flexible linker which comprises serine, glutamine and/or glycine or glutamic acid, alanine and lysine as amino acid building blocks, preferably selected from the group consisting of SEQ ID NO: 1 to 4, 6 to 12 and 1125.

19. The antigen-binding molecule according to any of the preceding claims, wherein the peptide linker between the first binding domain or the second binding domain and the spacer, and/or the third binding domain and the fourth binding domain and the spacer, respectively, is preferably a short linker rich in small and/or hydrophilic amino acids, preferably glycine and preferably SEQ ID NO: 5.

20. The antigen-binding molecule according to any of the preceding claims, wherein any of the first target cell surface antigen and the second target cell surface antigen is selected from the group consisting of CS1, BCMA, CDH3, FLT3, CD123, CD20, CD22, EpCAM, MSLN and CLL1.

21. The antigen-binding molecule according to any of the preceding claims, wherein the first target cell surface antigen and the second target cell surface antigen are not identical.

22. The antigen-binding molecule according to any of the preceding claims 1 to 20, wherein the first target cell surface antigen and the second target cell surface antigen are identical.

23. The antigen-binding molecule according to any of the preceding claims, wherein the first binding domains is capable of binding to the first target cell surface antigen and the third binding domain is capable of binding to the second target cell surface antigen simultaneously, preferably wherein the first target cell surface antigen and the second target cell surface antigen are on the same target cell.

24. The antigen-binding molecule according to any of the preceding claims, wherein the first target cell surface antigen and the second target cell surface antigen, respectively, are selected from the group consisting of CS1 and BCMA, BCMA and CS1, FLT3 and CD123, CD123 and FLT3, CD20 and CD22,

CD22 and CD20, EpCAM and MSLN, MSLN and EpCAM, MSLN and CDH3, CDH3 and MSLN, FLT3 and CLL1, and CLL1 and FLT3.

25. The antigen-binding molecule according to any of the preceding claims, wherein the first target cell surface antigen and/or the second target cell surface antigen is human MSLN (selected from SEQ ID NOs: 1181, 1182 and 1183), and wherein the first and/or third binding domain of the antigen-binding molecule of the invention binds to human MSLN epitope cluster E1 (SEQ ID NO: 1175, aa 296-346 position according to Kabat) as determined by murine chimere sequence analysis as described herein, but preferably not to human MSLN epitope cluster E2 (SEQ ID NO: 1176, aa 247-384 position according to Kabat), E3 (SEQ ID NO: 1177, aa 385-453 position according to Kabat), E4 (SEQ ID NO: 1178, aa 454-501 position according to Kabat) and/or E5 (SEQ ID NO: 1179 aa 502-545 position according to Kabat).

26. The antigen-binding molecule according to any of the preceding claims wherein the first target cell surface antigen and/or the second target cell surface antigen is human CDH3 (SEQ ID NOs: 1170), and wherein the first and/or third binding domain of the antigen-binding molecule of claim 1 binds to human CDH3 epitope cluster D2B (SEQ ID NO: 1171, aa 253-290 position according to Kabat), D2C (SEQ ID NO: 1172 aa 291-327 position according to Kabat), D3A (SEQ ID NO: 1173 aa 328-363 position according to Kabat) and D4B (SEQ ID NO: 1174, aa 476-511 position according to Kabat), preferably D4B (SEQ ID NO: 1174, aa 476-511 position according to Kabat), as determined by murine chimere sequence analysis as described herein.

27. The antigen-binding molecule according to any of the preceding claims, wherein the second and the fourth binding domain (CD3 binding domains) both have (i.) an affinity lower than characterized by a KD value of about 1.2×10^{-8} M measured by surface plasmon resonance (SPR), or (ii.) an affinity characterized by a KD value of about 1.2×10^{-8} M measured by SPR.

28. The antigen-binding molecule according to any of the preceding claims, wherein the second and the fourth binding domain (CD3 binding domains) have an affinity characterized by a KD value of about 1.0×10^{-7} to 5.0×10^{-6} M measured by SPR, preferably about 1.0 to 3.0×10^{-6} M, more preferably about 2.5×10^{-6} M measured by SPR.

29. The antigen-binding molecule according to any of the preceding claims, wherein the second and the fourth binding domain (CD3 binding domains) have an affinity characterized by a KD value of about 1.0×10^{-7} to 5.0×10^{-6} M measured by SPR, preferably about 1.0 to 3.0×10^{-6} M, more preferably about 2.5×10^{-6} M measured by SPR.

30. The antigen-binding molecule according to any of the preceding claims, wherein each of the second and the fourth binding domain (CD3 binding domains) individually has an at least about 10-fold, preferably at least about 50-fold or more preferably at least about 100-fold lower activity than one CD3 binding domain comprising a VH according to SEQ ID NO 43 and a VL according to SEQ ID NO 44 (i.e. in a mono targeting context in contrast to a dual targeting context).

31. The antigen-binding molecule according to any of the preceding claims, wherein the second and the fourth binding domain comprise a VH region comprising CDR-H 1, CDR-H2 and CDR-H3 selected from SEQ ID NOs 37 to 39, 45 to 47, 53 to 55, 61 to 63, 69 to 71, 436 to 438, 1126 to 1128, 1136 to 1138, 1142 to 1144, 1148 to 1150, and 1217 to 1219 and a VL region comprising CDR-L1, CDR-L2 and CDR-L3 selected from SEQ ID NOs 40 to 42, 48 to 50, 56 to 58, 64 to 66, 72 to 74, 439 to 441, 1129 to 1131, 1139 to 1141, 1145 to 1147, 1151 to 1153, and 1220 to 1222, preferably 61 to 63 and 64 to 66 or 1217 to 1219 and 1220 to 1222.

32. The antigen-binding molecule according to any of the preceding claims, wherein the second and fourth binding domain comprise a VH region selected from SEQ ID NOs 43, 51, 59, 67, 75, 442, 1132 and 1223, preferably 67 or 1223.

33. The antigen-binding molecule according to any of the preceding claims, wherein the second and fourth binding domain comprise a VL region selected from SEQ ID NOs 44, 52, 60, 68, 76, 443 1133 and 1224, preferably 68 or 1224.

34. The antigen-binding molecule according to any of the preceding claims, wherein the second and fourth binding domain comprising a VH region selected from SEQ ID NOs 43, 51, 59, 67, 75, 442, 1132 and 1223, preferably 67, and a VL region selected from SEQ ID NOs 44, 52, 60, 68, 76, 443, 1133 and 1224, preferably 68, wherein when the VH region is 1132 and the VL region is 1133, the second and/or fourth binding domain as scFab domain additionally comprises a CH1 domain of SEQ ID NO: 1134 and a CLK domain of SEQ ID NO: 1135, and wherein the VH and VL region are linked to each other by a linker preferably selected from SEQ ID NO 1, 3 and 1125, or wherein the VH of the VH-CH1 of the second and fourth domain is SEQ ID NO 1223, and the CH1 of the VH-CH1 of the second and fourth domain is SEQ ID NO 1224, and the VL of the VL-CL of the second and fourth domain is SEQ ID NO 1225, and the CL of the VL-L of the second and fourth domain is SEQ ID NO 1226.

35. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or the third (target) binding domain bind to CDH3 and comprise a VH region comprising SEQ ID NO: 1154 as CDR-H 1 wherein X1 (the number behind the "X" indicates the numerical order of the "X" in

respective amino acid sequence in N- to C-orientation in the sequence table) is S or N, X2 is Y or S, X3 is P or W, X4 is I or M and X5 is Y, N or H; SEQ ID NO: 1155 as CDR-H2 wherein X1 is K, V, N or R; X2 is A, D, R, Y, S, W or H; X3 is Y, S, P, G or T; X4 is S, G or K; X5 is A, V, D, K, G, or T; X6 is A, V, D, K, S, G or H; X7 is Y, G, or E; X8 is K, I, or N; X9 is A, S, or N; X10 is S, Q or G; X11 is S or K; X12 is F or V; and X13 is K or Q; and SEQ ID NO: 1156 as CDR-H3, wherein X1 is F or Q; X2 is R,K,S or W; X3 is G or D; X4 is Y, P or R; X5 is R, S, G, N or T; X6 is Y, A or H; X7 is F, L or M; X8 is A or V; and X9 is Y or V; and wherein the first and/or the third (target) binding domain bind to CDH3 and comprise a VL region comprising SEQ ID NO: 1158 as CDR-L 1 wherein X1 is K or R, X2 is A or S; X3 is Q,D,S,G or E; X4 is S, D or N; X5 is V, L or I; X6 is ,K, Y, S, or H; X7 is S or N; X8 is F, L or M; and X9 is A,N or H; SEQ ID NO: 1159 as CDR-L 2 wherein X1 is Y, G, W, or N; X2 is T or A; X3 is S or K; X4 is T, N or R; X5 is L or R; X6 is E, A, V or H; and X7 is S or E; and SEQ ID NO: 1160 as CDR-L3 wherein X1 is Q or V; X2 is Q, N or H; X3 is F, L, Y, W, N, or H; X4 is A, D, Y, S or N; X5 is Q, R, S, G, W or M; X6 is T, Y or F; and X7 is F,Y or L.

36. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or the third (target) binding domain bind to MSLN and comprise a VH region comprising SEQ ID NO: 1162 as CDR-H 1 wherein X1 (the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence table) is S, G or D; X2 is Y, A, G or F; X3 is I, W, or M; and X4 is V, S, G, T, or H; SEQ ID NO: 1163 as CDR-H 2 wherein X1 is A, S, N, W, Y, or V; X2 is Y, S or N; X3 is Y, G, P, or S; X4 is D, H, S, or N; X5 is G or S; X6 is E, G or S; X7 is G, S, N, F, T or Q; X8 is S, W, K, D, I or T; X9 is Y or N; X10 is A or N; X11 is A, P, N, D, E, I or Q; X12 is D, A, S or K; X13 is V, L, or F; X14 is K or Q; and X15 is G or S; and SEQ ID NO: 1164 as CDR-H 3 wherein X1 is D, E or V; X2 is R, G, or E; X3 is Y, A, or N; X4 is S,Y,V, or H; X5 is A,P,F,Y, or H; X6 is R or S; X7 is E or G; X8 is Y or L; X9 is R,Y or L; X10 is Y or G; X11 is D or Y; X12 is R,Y, or F; X13 is M,S,F,D or Y; X14 is A,G,S,or T; X15 is L, M,or F; and X16 is Y,I or V; and wherein the first and/or the third (target) binding domain bind to MSLN and comprise a VL region comprising SEQ ID NO: 1166 as CDR-L 1 wherein X1 is A or S; X2 is G or S; X3 is E or Q; X4 is G,S or K; X5 is I,L,V or F; X6 is R,G or S; X7 is D,S,N or T; X8 is A,S,K or T; X9 is Y or W; X10 is V or L; and X11 is Y or A; SEQ ID NO 1167 as CDR-L2 wherein X1 is A,G or Q; X2 is A or S; X3 is S or T; X4 is G,S,K,I or T; X5 is R or L; X6 is A,P or Q; and X7 is S or T; and SEQ ID NO 1168 as CDR-L 3 wherein X1 is A or Q; X2 is Y, S, A, or T; X3 is G, E, Y, H or Q; X4 is A or S; X5 is S, T or F; X6 is P or T; X7 is R, A, L or F; and X8 is V or T.

37. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or the third (target) binding domain bind to CDH3 and comprise a VH region of SEQ ID NO: 1157 wherein

(the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence table) X1 is Q or E; X2 is V,L; X3 is Q,E ;X4 is A or G; X5 is G or E; X6 is V or L; X7 is K or V X8 is K or Q, X9 is A or G, X10 is V or L, X11 is K or R, X12 is V or L, X13 is A or K, X14 is Y or F, X15 is T or S, X16 is T or S, X17 is S or N, X18 is Y or S, X19 is P or W, X20 is I or M, X21 is Y, N or H, X22 is T or A, X23 is Q or K, X24 is V or M, X25 is S or G, X26 is K, V, N or R, X27 is A, D, R, Y, S, W or H, X28 is Y, S, P, Gr or T, X29 is S, K, or G, X30 is A, V, D, K, or ,T; X31 is A, D, K, S, G, or H; X32 is Y,G, or E, X33 is K,I, or N, X34 is A,S, or N, X35 is S,Q, or G, X36 is S or K, X37 is F or V, X38 is Q or K, X39 is F or V, X40 is I or M, X41 is T or S, X42 is V,I or R, X43 is T,K or N, X44 is T,A,S or K, X45 is S or N, X46 is A,V or L, X47 is L or M, X48 is Q or E, X49 is L or M, X50 is S or N, X51 is S or R, X52 is T or R, X53 is A or S, X54 is G, D or E; X55 is T or S, X56 is T, K, or R, X57 is S, Q, W, or R, X58 is D, or G, X59 is Y, P, or R; X60 is F,S,G,N or T, X61 is Y, A, or H, X62 is A,-,or V, X63 is F or M, X64 is Y or V; X65 is T,L or M ; and a VL region of SEQ ID NO 1161 wherein X1 is D or E; X2 Q or V; X3 is L,M; X4 is A,S or D; X5 is F,S or T; X6 is A or S; X7 is A or V; X8 is P,V or L; X9 is D or E; X10 is A or V; X11 is I or L; X12 is T, S, or N; X13 is K or R; X14 is A,S; or X15 is Q,D,S,G or E; X16 is S, D or N; X17 is V, I or L; X18 is K, Y, S or H; X19 is S or N; X20 is F, L or M; X21 is A, N or H; X22 is K or Q; X23 is A, P or V; X24 is K or R; X25 is I or V; X26 is Y, G, W or N; X27 is T or A; X28 is S or K; X29 is T, N or R; X30 is L or R; X31 is E, A, V or H; X32 is S or E; X33 is A, S, V or D; X34 is D or E; X35 is T or K; X36 is S or R; X37 is A,S or P; X38 is F or V; X39 is A,G; X40 is T or V; X41 is Q or V; X42 is Q, N, H; X43 is F, L, Y, W, N or H; X44 is A, D, Y, S or N; X45 is Q, R, S, G, W or M; X46 is F, Y or T; X47 is F, Y or L; X48 is V or L; and X49 is D or E (wherein all aa per position are preferably meant to be in the alternative “or” even if not explicitly stated).

38. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or the third (target) binding domain bind to MSLN and comprise a VH region of SEQ ID NO: 1165 wherein (the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence table) X1 is E or Q; X2 is V,L or Q; X3 is E or Q; X4 is A,G or P; X5 is E or G; X6 is V or L; X7 is V or K; X8 is K or Q; X9 is G or S; X10 is E, A, G or R; X11 is S or T; X12 is V or L; X13 is R, S or K; X14 is V or L; X15 is S or T; X16 is A,K or T; X17 is A or V; X18 is Y, I or F; X19 is S or T; X20 is S or F; X21 is S or T; X22 is D, G or S; X23 is Y, G, A or F; X24 is I, W or M; X25 is G, S, V, T or H; X26 is I or V; X27 is A or P; X28 is M, K or Q; X29 is G or C; X30 is I, M, V or L; X31 is A, G or S; X32 is A, S, N, W, Y or V; X33 is Y, S or N; X34 is Y, G, P or S; X35 is D, H, S or N; X36 is G or S; X37 is E, G or S; X38 is G, S, N, F, T or Q; X39 is S,

K, W, D, I, or T; X40 is Y or N; X41 is A or N; X42 is A, P, N, E, D, I or Q; X43 is D, A, S or K; X44 is V,L or F; X45 is K,Q; X46 is G or S; X47 is V or F; X48 is I or M; X49 is S or T; X50 is R or V; X51 is N or T; X52 is A or S; X53 is I or K; X54 is S or N; X55 is S, T or Q; X56 is A, L or F; X57 is Y, S or F; X58 is L or M; X59 is E, K or Q; X60 is M or L; X61 is S or N; X62 is R or S; X63 is V or L; X64 is R or T; X65 is A or S; X66 is D, A or E; X67 is R or K; X68 is D, E, V or L; X69 is E, R, G or P; X70 is R, A, N or Y; X71 is G, S, Y, V or H; X72 is A, P, F, D or Y; X73 is R or G; X74 is M, R, S or D; X75 is E or G; X76 is Y or L; X77 is Y or F; X78 is Y, S or F; X79 is A, G, S, T or H; X80 is L, M or F; X81 is Y, I or V; and X82 is L, M or T; and a VL region of SEQ ID NO 1169 (the number behind the “X” indicates the numerical order of the “X” in respective amino acid sequence in N- to C-orientation in the sequence table) X1 is E,S or D; X2 is Y,I or L; X3 is E,V or T; X4 is V,L or M; X5 is P or S; X6 is G or S; X7 is S or T; X8 is V or L; X9 is A, V or L; X10 is P or V; X11 is E, Q or D; X12 is R or T; X13 is A or V; X14 is S or T; X15 is I or L; X16 is S or T; X17 is A or S; X18 is G or S; X19 is E or Q; X20 is G,S or K; X21 is I, V, L or F; X22 is R, G or S; X23 is D or S; X24 is A, S, N, K or T; X25 is Y, W or M; X26 is V or L; X27 is Y or A; X28 is K or Q; X29 is A,S or V; X30 is R,V or K; X31 is V or L; X32 is A,G or Q; X33 is A or S; X34 is S or T; X35 is G,S,K,I or T; X36 is R or L; X37 is A,P or Q; X38 is S or T; X39 is I or V; X40 is E,S or D; X41 is G or N; X42 is N or T; X43 is D or T; X44 is A or F; X45 is R,G or S; X46 is L or T; X47 is E or Q; X48 is A or P; X49 is E or M; X50 is E or F; X51 is D,V or T; X52 is A or Q; X53 is Y,S,A or T; X54 is G,E,Y,H or Q; X55 is A or S; X56 is S,T or F; X57 is P or T; X58 is R, A, L or F; X59 is V or T; X60 is P or C; X61 is V or L; X62 is E or T; X63 is I or V; and X64 is L or K (wherein all aa per position are preferably meant to be in the alternative “or” even if not explicitly stated).

39. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or the third (target) binding domain comprise a VH region comprising CDR-H 1, CDR-H2 and CDR-H3 selected from SEQ ID NO: 77 to 79, 86 to 88, 95 to 97, 103 to 105, 111 to 113, 119 to 121, 127 to 129, 135 to 137, 143 to 145, 151 to 153, 159 to 161, 168 to 170, 177 to 179, 185 to 187, 194 to 196, 203 to 205, 212 to 214, 221 to 223, 230 to 232, 238 to 240, 334 to 336, 356 to 358, 365 to 367, 376 to 378, 385 to 387 and 194, 432 and 196, 446 to 448, 454 to 456, 462 to 464, 470 to 472, 478 to 480, 486 to 488, 494 to 496, 502 to 504, 510 to 512, 518 to 520, 526 to 528, 534 to 536, 542 to 544, 550 to 552, 558 to 560, 566 to 568, 574 to 576, 582 to 584, 590 to 592, 598 to 600, 606 to 608, 614 to 616, 622 to 624, 630 to 632, 638 to 640, 646 to 648, 654 to 656, 662 to 664, 670 to 672, 678 to 680, 686 to 688, 694 to 696, 702 to 704, 710 to 712, 718 to 720, 726 to 728, 734 to 736, 742 to 744, 750 to 752, 758 to 760, 766 to 768, 774 to 776, 782 to 784, 790 to 792, 798 to 800, 806 to 808, 814 to 816, 822 to 826, 830 to 832, 838 to 840, 846 to 848, 854 to 856, 862 to 864, 870 to 872, 878 to 880, 886 to 888, 894 to 896, 902 to 904, 910 to 912, 918 to 920, 926 to 928,

934 to 936, 942 to 944, 950 to 952, 958 to 960, 966 to 968, 974 to 976, 982 to 984, 990 to 992, 998 to 1000, 1006 to 1008, 1014 to 1016, 1022 to 1024, 1030 to 1032, 1038 to 1040, 1046 to 1048, 1054 to 1056, and 1062 to 1064, or preferably any combination of CDR-H 1, CDR-H2 and CDR-H3 as disclosed together in the sequence table Tab. 6, preferably 86 to 88 and 194, 432 and 196 for the first and the third binding domain, respectively, more preferably 194, 432 and 196 for the first and 86 to 88 for the third binding domain or 1227 to 1229 and 1237 to 1239 for the first and third binding domain.

40. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or third (target) binding domain comprise a VL region comprising CDR-L1, CDR-L2 and CDR-L3 selected from SEQ ID NO: 80 to 82, 89 to 91, 98 to 100, 106 to 108, 114 to 116, 122 to 124, 130 to 132, 138 to 140, 146 to 148, 154 to 156, 162 to 164, 171 to 173, 180 to 182, 188 to 190, 197 to 199, 206 to 208, 215 to 217, 224 to 226, 233 to 235, 241 to 243, 337 to 339, 359 to 361, 368 to 370, 379 to 381, 388 to 390, 449 to 451, 457 to 459, 465 to 467, 473 to 475, 481 to 483, 489 to 491, 497 to 499, 505 to 507, 513 to 515, 521 to 523, 529 to 531, 537 to 539, 545 to 547, 553 to 555, 561 to 563, 569 to 571, 577 to 579, 585 to 587, 593 to 595, 601 to 603, 609 to 611, 617 to 619, 625 to 627, 633 to 635, 641 to 643, 649 to 651, 657 to 659, 665 to 667, 673 to 675, 681 to 683, 689 to 691, 697 to 699, 705 to 707, 713 to 715, 721 to 723, 729 to 731, 737 to 739, 745 to 747, 753 to 755, 761 to 763, 769 to 771, 777 to 779, 785 to 787, 793 to 795, 801 to 803, 809 to 811, 817 to 819, 825 to 829, 833 to 835, 841 to 843, 849 to 851, 857 to 859, 865 to 867, 873 to 875, 881 to 883, 889 to 891, 897 to 899, 905 to 907, 913 to 915, 921 to 923, 929 to 931, 937 to 939, 945 to 947, 953 to 955, 961 to 963, 969 to 971, 977 to 979, 985 to 987, 993 to 995, 1001 to 1003, 1009 to 1011, 1017 to 1019, 1025 to 1027, 1033 to 1035, 1041 to 1043, 1049 to 1051, 1057 to 1059, and 1065 to 1067 or preferably any combination of CDR-L 1, CDR-L2 and CDR-L3 as disclosed together in the sequence table Tab. 6, preferably 89 to 91 and 197 to 199 for the first and the third binding domain, respectively, more preferably 197 to 199 for the first and 89 to 91 for the third binding domain or 1230 to 1232 and 1240 1242 for the first and the third binding domain.

41. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or third (target) binding domain comprise a VH region selected from SEQ ID NO: 83, 92, 101, 109, 117, 125, 133, 141, 149, 157, 165, 174, 183, 191, 200, 209, 218, 227, 236, 244, 340, 362, 371, 382, 391, and 433, 452, 460, 468, 476, 484, 492, 500, 508, 516, 524, 532, 540, 548, 556, 564, 572, 580, 588, 596, 604, 612, 620, 628, 636, 644, 652, 660, 668, 676, 684, 692, 700, 708, 716, 724, 732, 740, 748, 756, 764, 772, 780, 788, 796, 804, 812, 820, 828, 836, 844, 852, 860, 868, 876, 884, 892, 900, 908, 916, 924, 932, 940, 948, 956, 964, 972, 980, 988, 996, 1004, 1012, 1020, 1028, 1036, 1044, 1052, 1060, and 1068 or preferably any VH as disclosed together in the sequence table Tab. 52, preferably 433 and 92 for the first and the third binding domain, respectively, more preferably 433 for the first and 92 for the third binding

domain or 1233 + 1235 and 1243 + 1245 (VH and CH1 in Fab) for the first and third binding domain, respectively.

42. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or third (target) binding domain comprises a VL region selected from SEQ ID NO: 84, 93, 102, 110, 118, 126, 134, 142, 150, 158, 166, 175, 184, 192, 201, 210, 219, 228, 237, 245, 341, 363, 372, 383, 392, 453, 461, 469, 477, 485, 493, 501, 509, 517, 525, 533, 541, 549, 557, 565, 573, 581, 589, 597, 605, 613, 621, 629, 637, 645, 653, 661, 669, 677, 685, 693, 701, 709, 717, 725, 733, 741, 749, 757, 765, 773, 781, 789, 797, 805, 813, 821, 829, 837, 845, 853, 861, 869, 877, 885, 893, 901, 909, 917, 925, 933, 941, 949, 957, 965, 973, 981, 989, 997, 1005, 1013, 1021, 1029, 1037, 1045, 1053, 1061, and 1069 or preferably any VL as disclosed together in the sequence table Tab. 52, preferably 200 and 93 for the first and the third binding domain, respectively, more preferably 200 for the first and 93 for the third binding domain or 1234 + 1236 and 1244 + 1246 (VL and CL in Fab) for the first and third binding domain, respectively.

43. The antigen-binding molecule according to any of the preceding claims, wherein the first and/or third (target) binding domain comprises a VL region of increased stability by a single amino acid exchange (E to I), selected from SEQ ID NO: 85, 94, 193, 202, 211, 220, 229, 364, 384, 393, preferably 94 and 202.

44. The antigen-binding molecule according to any of the preceding claims, which comprises a combination of amino acid sequences selected from the group consisting of SEQ ID NOs: 1259 and 1251, 1247 and 1248, 1249 and 1250, 1254, 1255 and 1253, 1252, 1257, 1253 and 1256, and 1254, 1258, 1253 and 1256, or any other full length multitargeting bispecific antigen—binding molecule as disclosed in the sequence table Tab. 6.

45. A polynucleotide encoding an antigen-binding molecule of any of claims 1 to 44.

46. A vector comprising a polynucleotide of claim 45.

47. A host cell transformed or transfected with the polynucleotide of claim 45 or with the vector of claim 46.

48. A process for the production of an antigen-binding molecule of any of claims 1 to 44, said process comprising culturing a host cell of the present invention under conditions allowing the expression of the antigen-binding molecule and recovering the produced antigen-binding molecule from the culture.

49. A pharmaceutical composition comprising an antigen-binding molecule of any of claims 1 to 38 or produced according to the process of claim 48.

50. The pharmaceutical composition of claim 49 which is stable for at least four weeks at about -20°C.

51. An antigen-binding molecule of claims 1 to 44 or produced according to the process of claim 48, for use in the prevention, treatment or amelioration of a disease selected from a proliferative disease, a tumorous disease, cancer or an immunological disorder.

52. The antigen-binding molecule according to claim 51, wherein the disease preferably is acute myeloid leukemia (AML), Non-Hodgkin lymphoma (NHL), Non-small-cell lung carcinoma (NSCLC), pancreatic cancer and Colorectal cancer (CRC).

53. A method for the treatment or amelioration of a proliferative disease, the method comprising administering to a subject in need thereof a molecule comprising at least one polypeptide chain, wherein the molecule comprises

- (i.) a first binding domain which binds to a first target cell surface antigen (TAA1),
- (ii.) a second binding domain which binds to an extracellular epitope of the human and/or the Macaca CD3 chain,
- (iii.) a third binding domain which binds to a second target cell surface antigen (TAA2), and
- (iv.) a fourth binding domain which binds to an extracellular epitope of the human and/or the Macaca CD3 chain,

wherein the first binding domain and the second binding domain form a first bispecific entity, and the third and the fourth binding domain form a second bispecific entity, and

wherein the molecule further comprises a spacer entity selected from

- (1.) a dimerizing domain selected from
 - (a.) an Fc domain comprising a first and a second polypeptide monomer comprising a hinge, a CH2 domain and a CH3 domain, respectively, wherein the first and second polypeptide monomer form a heterodimer; wherein the heterodimer is formed by
 - charged pair mutations selected from (i.) D399K, K409D, K392D, and E356K, (ii.) D399K, K409D, K392D, E357K, K370D, and E356K, (iii.) D399K, K409D, K392D, E356K and K439D, (iv.) D399K, K409D, and K392D, (v.) D399K, K409D, K392D, E357K and E370K, (vi.) D399K, K409D, K392D,

E357K, K370E and K360E, (vii.) D399K, K409D, K392D, E357K, K370E, E356K, and K439E, and (viii.) D399K, K409D, K392D, E357K, K370E, K360E, E356K, and K439D, preferably comprising a K392D, K409D and/or K439D mutation in the CH3 domain of the first polypeptide monomer and comprising a E356K and/or D399K mutation in the CH3 domain of the second polypeptide monomer, wherein the positions are according to EU numbering; or

- knobs-into-holes mutations comprising preferably a T366S, L368A and Y407V mutation in the first polypeptide monomer and a T366W mutation in the second monomer, wherein the positions are according to EU numbering;

(b.) a human serum albumin (HSA) domain comprising a first and a second polypeptide monomer, wherein the first and the second polypeptide monomer correspond to an HSA subdomain, respectively, wherein the first and second polypeptide monomer form a native HSA-like heterodimer; and

(c.) a Fab comprising a first and a second polypeptide monomer, wherein preferably the first polypeptide monomer comprises a VL and CL domain and the second polypeptide monomer comprises a VH and CH1 domain, wherein the CL and CH1 domains are linked by a disulfide bridge;

wherein the dimerizing domain comprises two N-termini and two C-termini, respectively, whereof at least one N-terminus and one C-terminus, respectively, is linked to a bispecific entity, wherein any of the first, second, third and fourth domain can be selected from any form of binding domain, preferably selected from Fab and single chain domain, the single chain domain preferably selected from single chain Fv (scFv) and scFab;

(2.) a single chain domain selected from ubiquitin, beta 2 microglobulin, VH-only domain, PSI domain from Met-receptor, Fibronectin type III domain from tenascin, Granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-4, Interleukin-2, PD-1 binding domain from human Programmed cell death 1 ligand 1 (PDL1), Tim-3 (AS 24-130), MiniSOG, a programmed cell death protein 1 (PD1) domain, human serum albumin (HSA), or a single chain Fc (scFc) domain comprising two polypeptide monomers comprising each a hinge, a CH2 and a CH3 domain a hinge and a further CH2 and a CH3 domain, wherein said two polypeptide monomers are fused to each other via a peptide linker,

wherein the single chain domain comprises one N-terminus and one C-terminus, which are, respectively, linked to a bispecific entity, wherein at least one of the first, second, third and fourth binding domain is a two-chained Fab, and any of the remaining at least three binding domains can be selected from any form of binding domain, preferably selected from Fab and single chain domain, the single chain domain preferably selected from scFv and scFab;

wherein the distance between the C alpha atoms of the first amino acid located at the N-terminus and the last amino acid at the C-terminus of the spacer entity are spaced apart by at least 30 Å, wherein the spacer entity spaces apart the first and the second bispecific entity by at least a distance of about 50 Å, wherein the indicated distance is preferably understood as the distance between centers of mass (i.) of the first and the third binding domain or (ii.) the first and the second bispecific entity, and which spacer entity is positioned between the first and the second bispecific entity comprising the step of administering to a subject in need thereof the antigen-binding molecule of the present invention, or produced according to the process of the present invention, wherein the disease preferably is , acute myeloid leukemia, Non-Hodgkin lymphoma, Non-small-cell lung carcinoma, pancreatic cancer and/or Colorectal cancer.

54. The method of claim 53, wherein the method comprises addressing a disease-associated target being significantly co-expressed on a pathophysiological and one or more physiological tissues by providing a multitargeting bispecific antigen-binding molecule of the format described herein, wherein the molecule addresses (i.) the target expressed both on the disease-associated and the physiological tissue and (ii.) a further target which is disease associated but not expressed on the physiological tissue under (i.), wherein the method preferably avoids the formation of intra-abdominal adhesions and/or fibrosis where such target is MSLN.

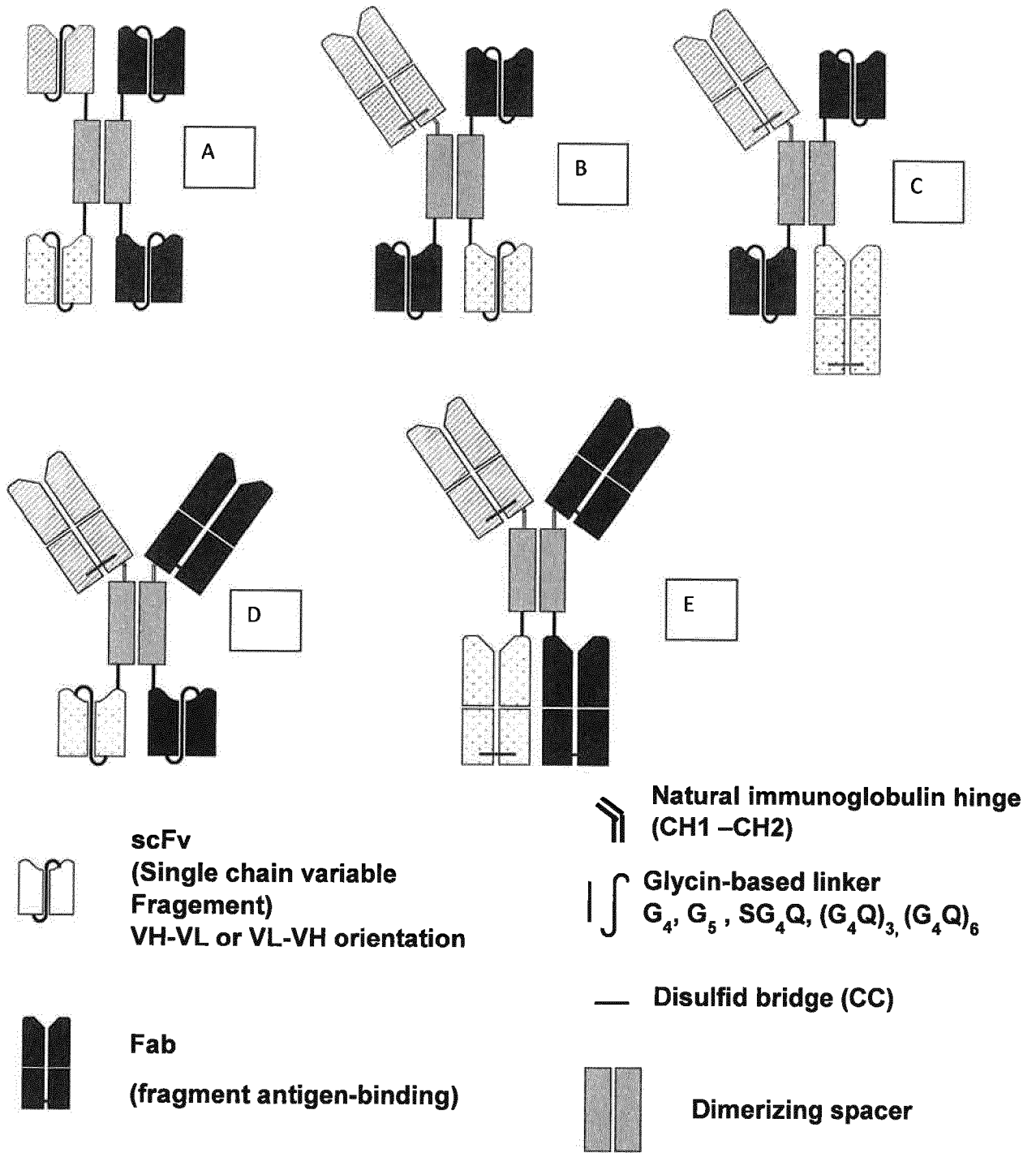
55. The method of claim 53, wherein the disease is a tumorous disease, a cancer, or an immunological disorder.

56. The method of claim 55, wherein the disease preferably is acute myeloid leukemia, Non-Hodgkin lymphoma, Non-small-cell lung carcinoma, pancreatic cancer and/or Colorectal cancer.

57. The method of claim 47, wherein the TAA1 and TAA2 are preferably selected from EpCAM and MSLN, MSLN and EpCAM, MSLN and CDH3, CDH3 and MSLN, FLT3 and CLL1, and CLL1 and FLT3.

58. A kit comprising an antigen-binding molecule of any of claims 1 to 44, or produced according to the process of claim 48, a polynucleotide of claim 45, a vector of claim 46, and/or a host cell of claim 47.

Fig. 1



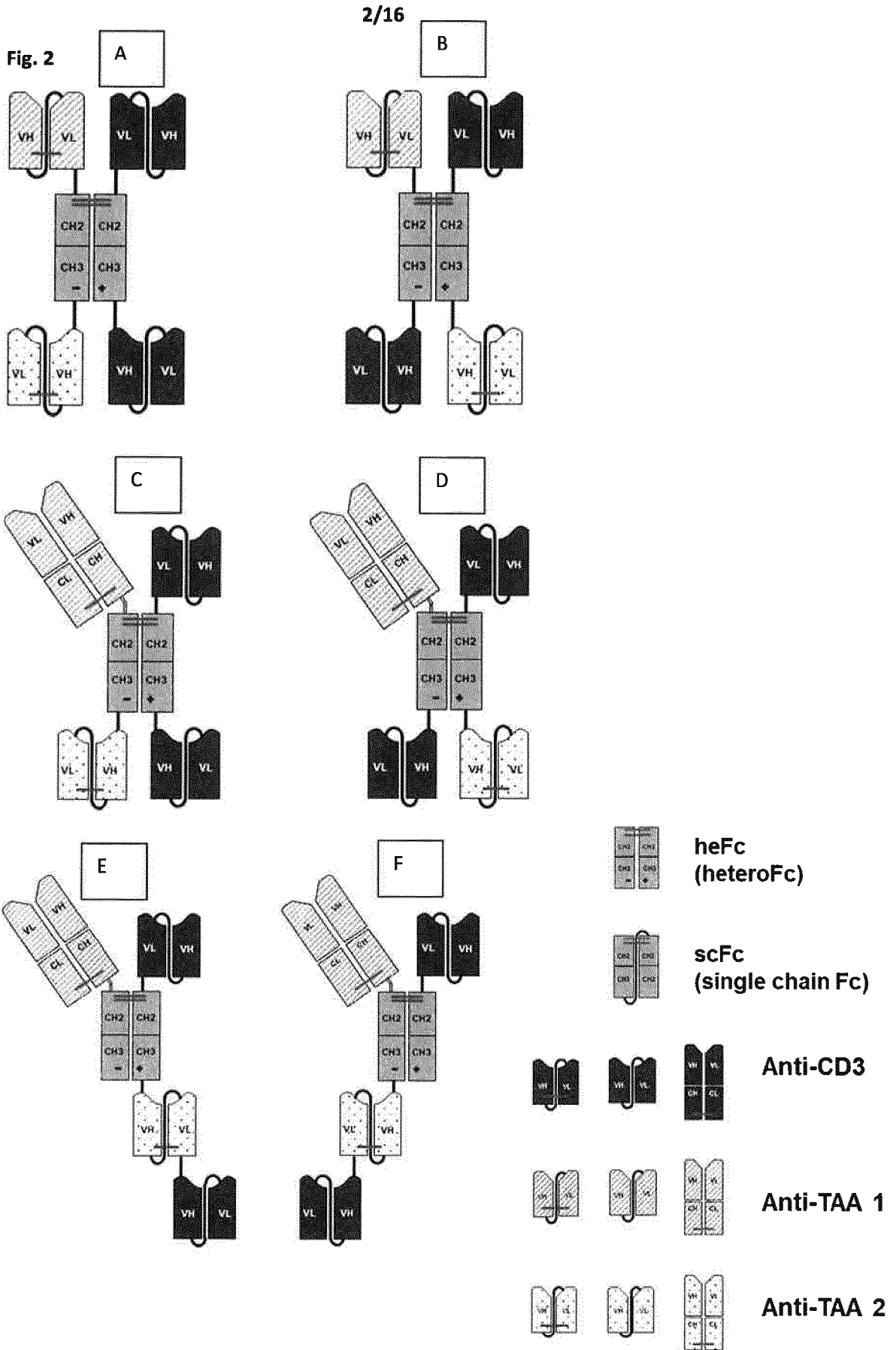


Fig. 2 (cont.)

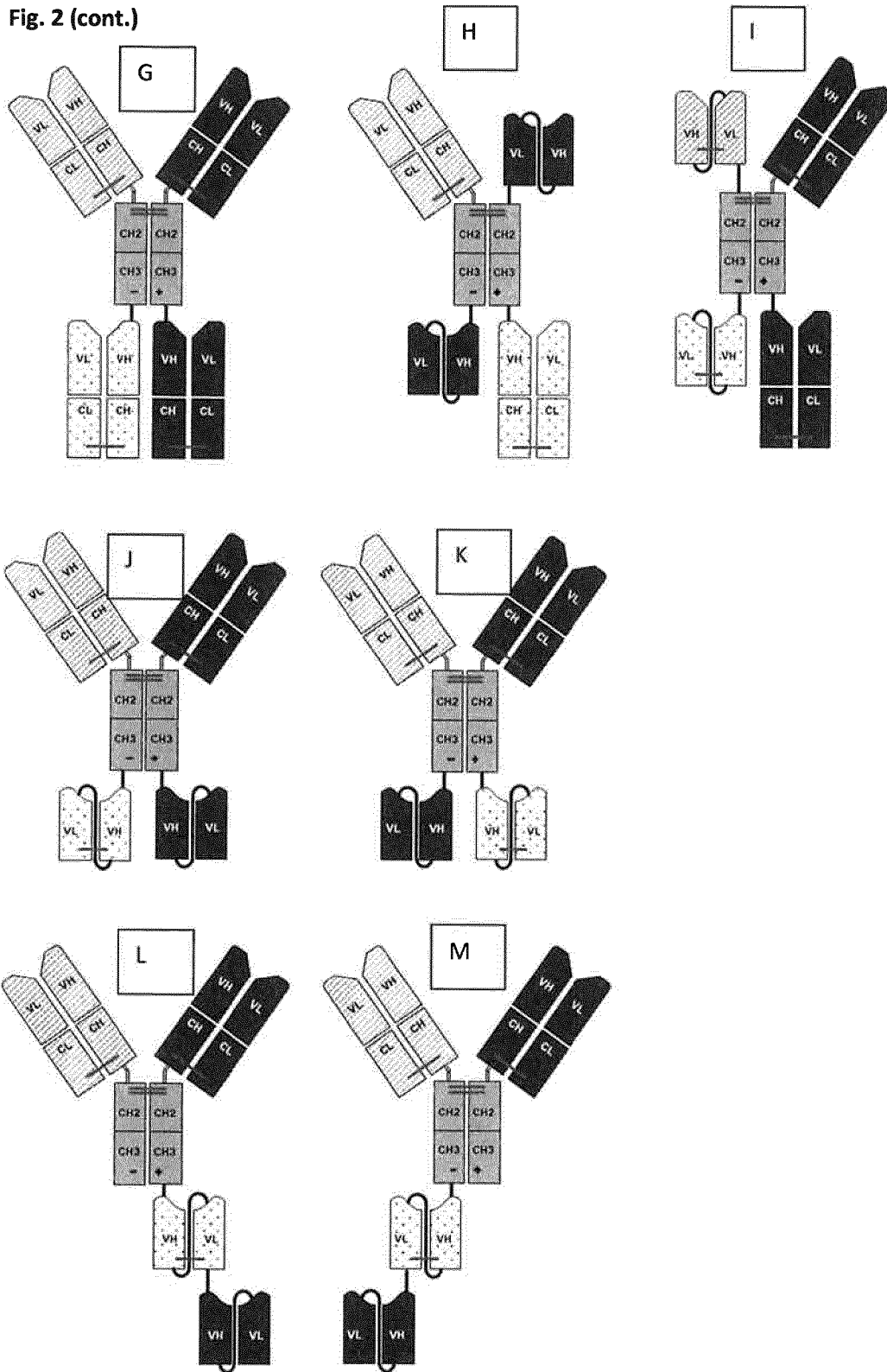


Fig. 3

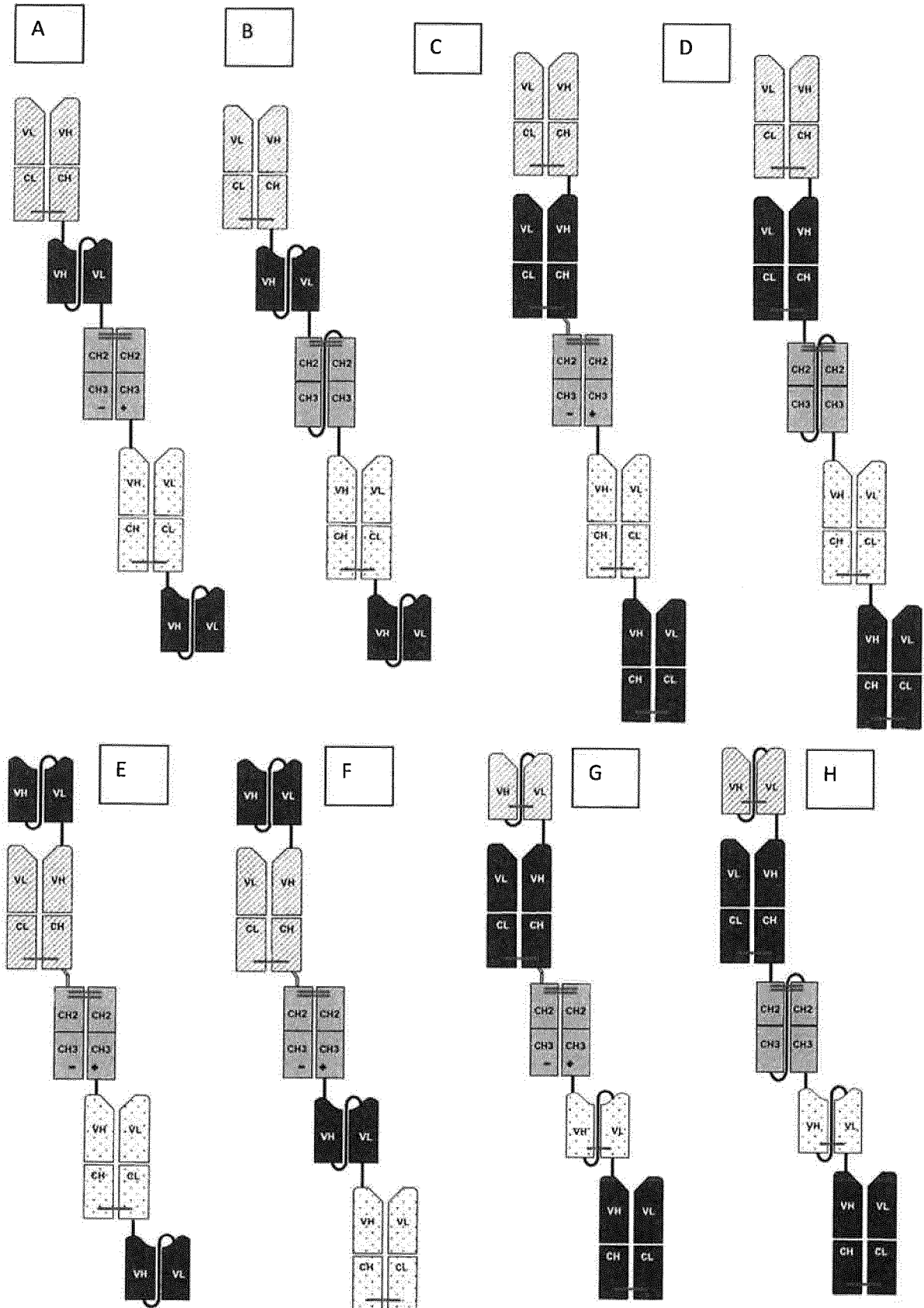


Fig. 3 (cont.)

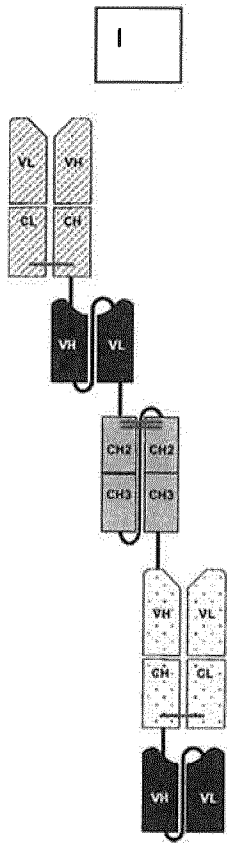


Fig. 4

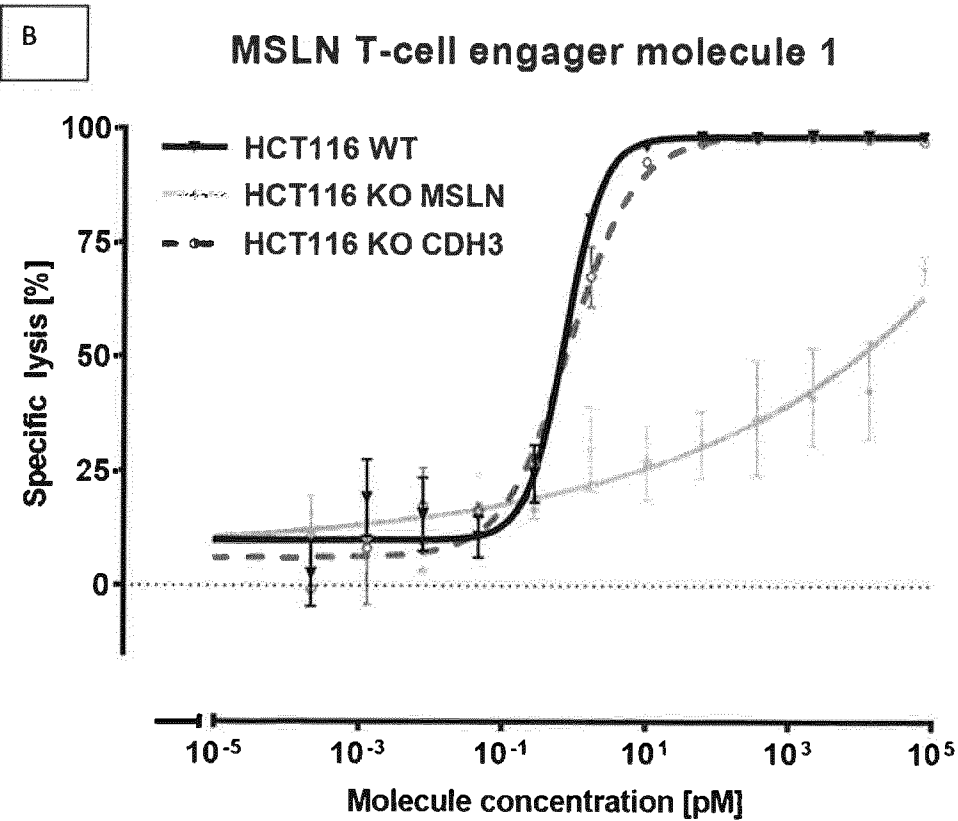
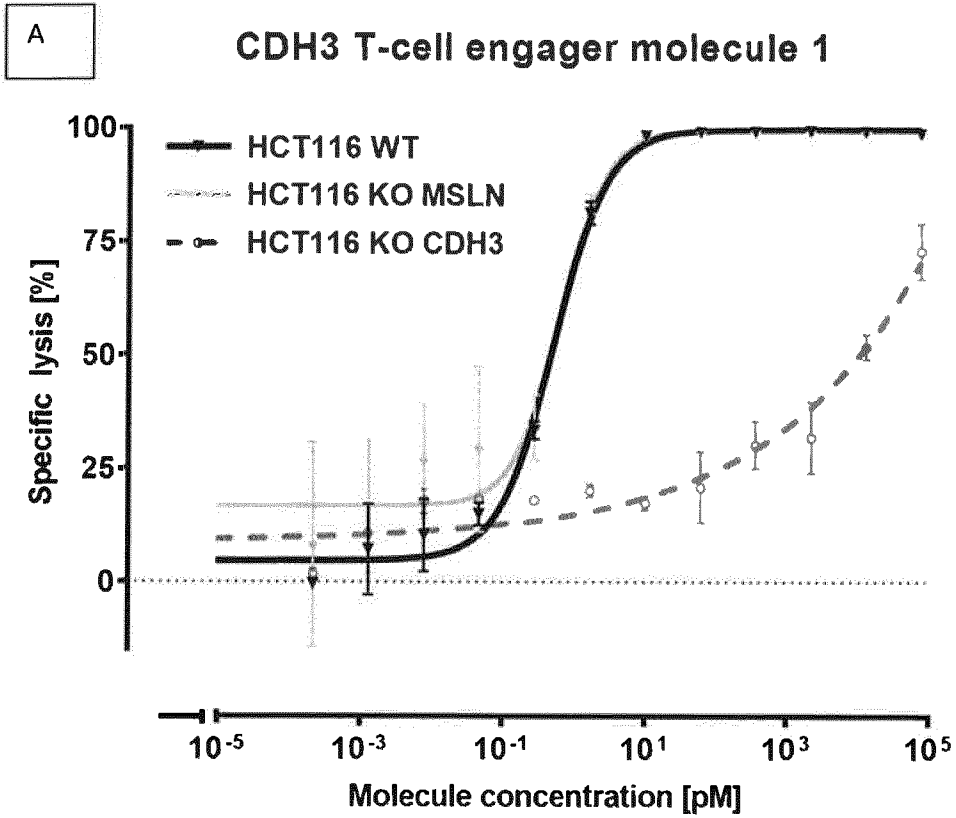
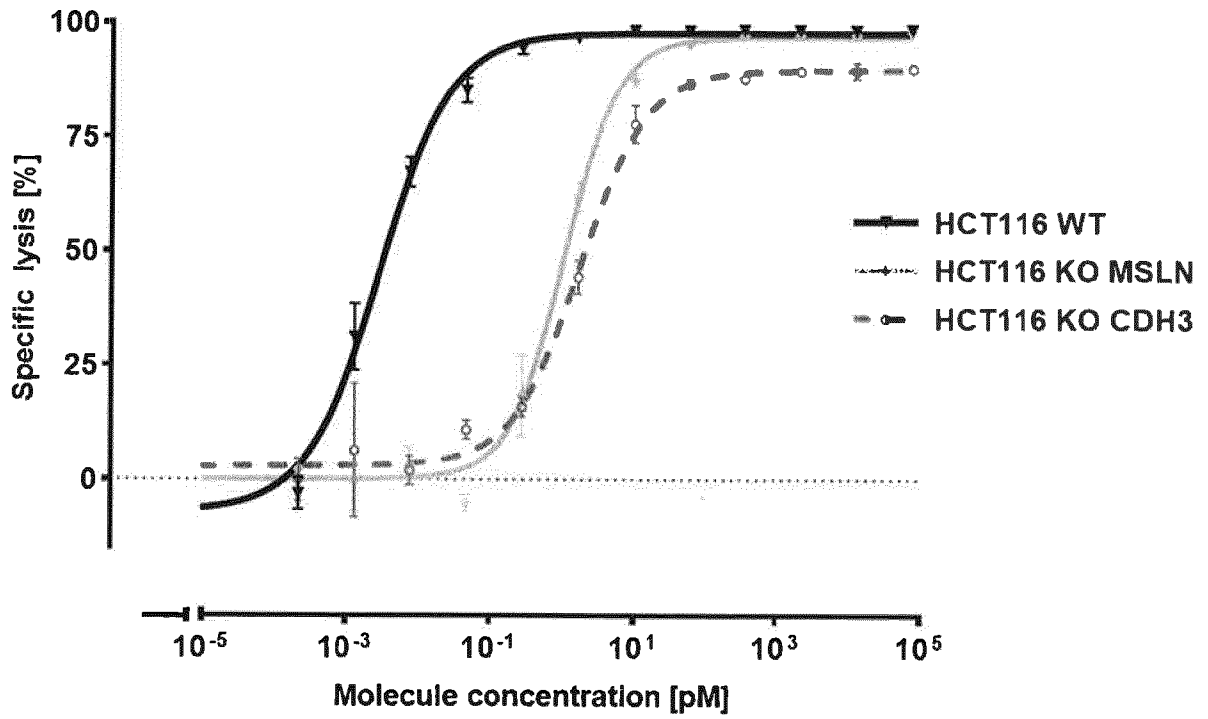


Fig. 4 (cont.)

C

MSLN-CDH3 T-cell engager molecule 1



D

MSLN-CDH3 T-cell engager molecule 2

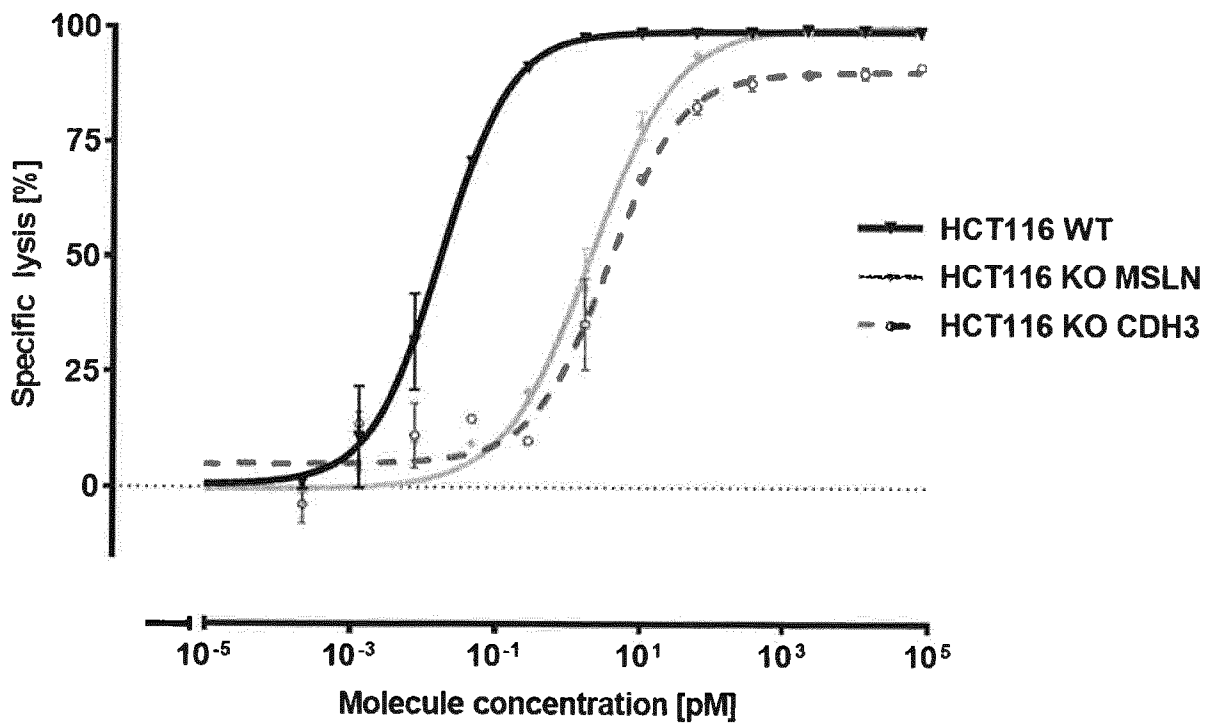
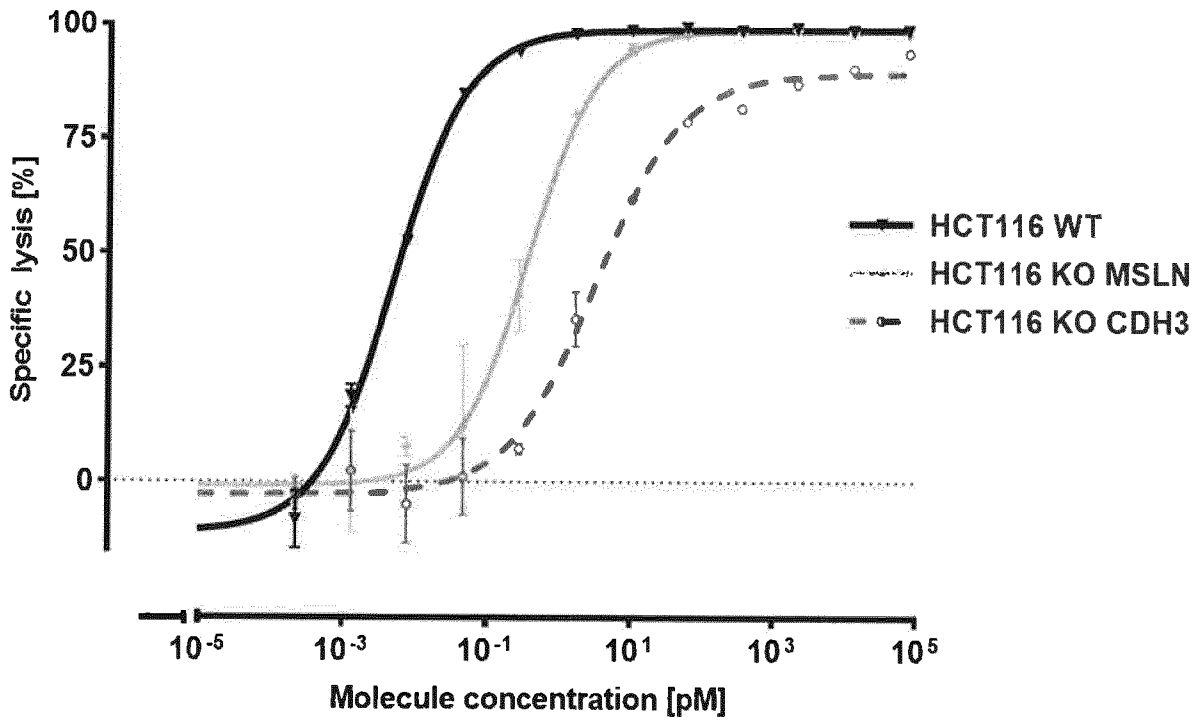


Fig. 4 (cont.)

E

MSLN-CDH3 T-cell engager molecule 3



F

MSLN-CDH3 T-cell engager molecule 4

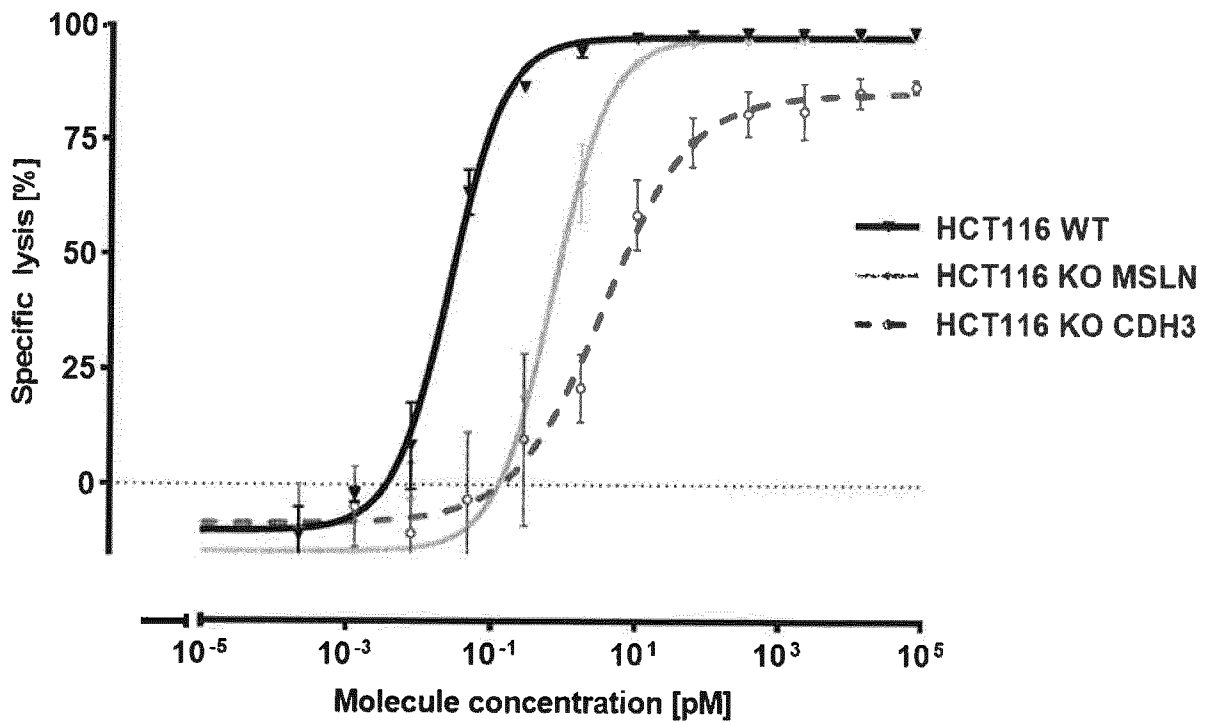
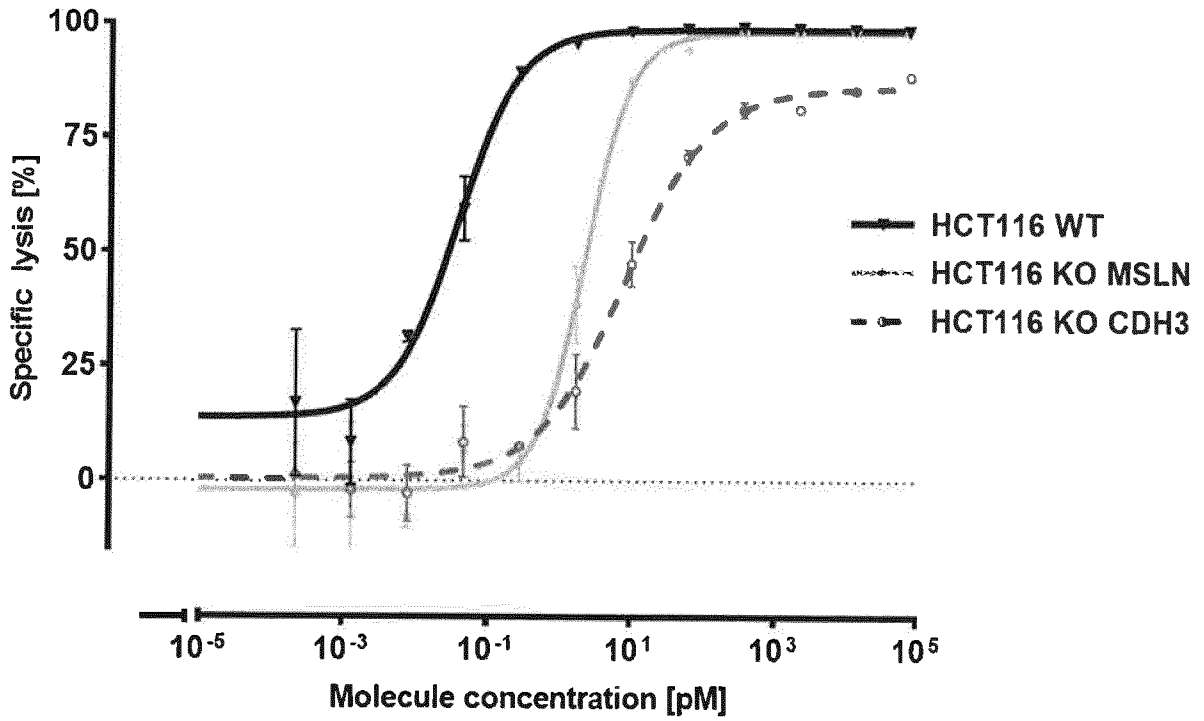


Fig. 4 (cont.)

G

MSLN-CDH3 T-cell engager molecule 5



H

MSLN-CDH3 T-cell engager molecule 6

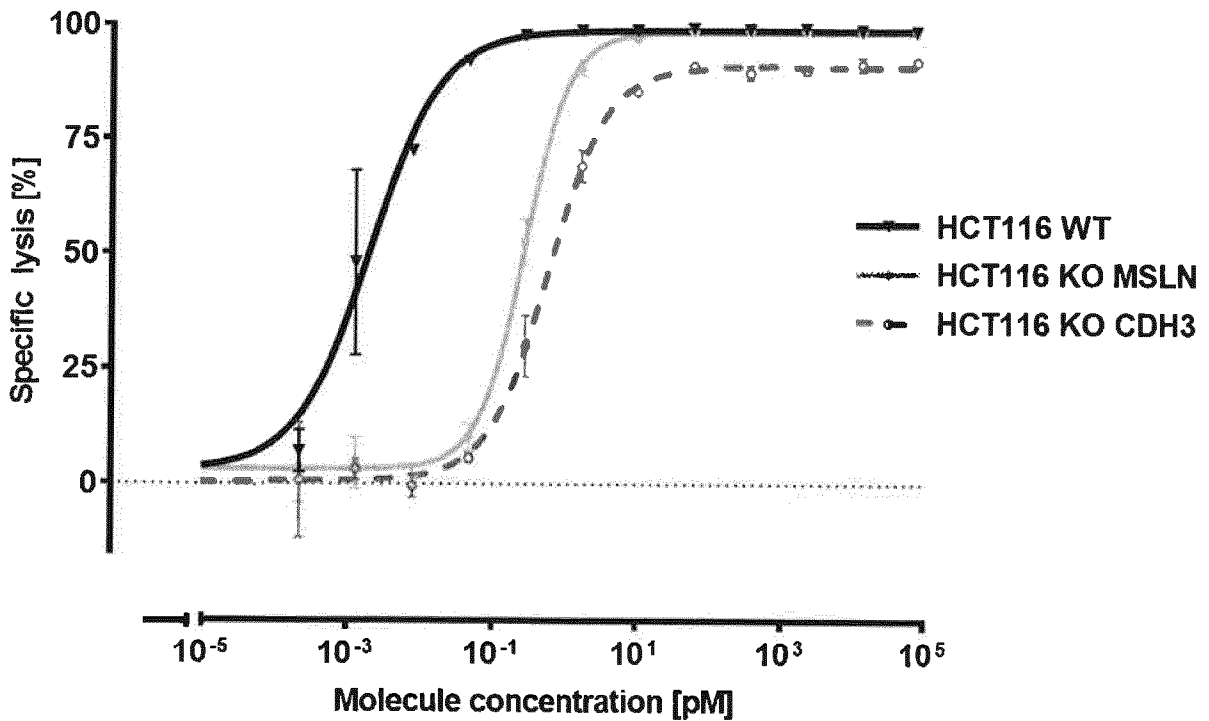


Fig. 4 (cont.)

I

MSLN-CDH3 T-cell engager molecule 7

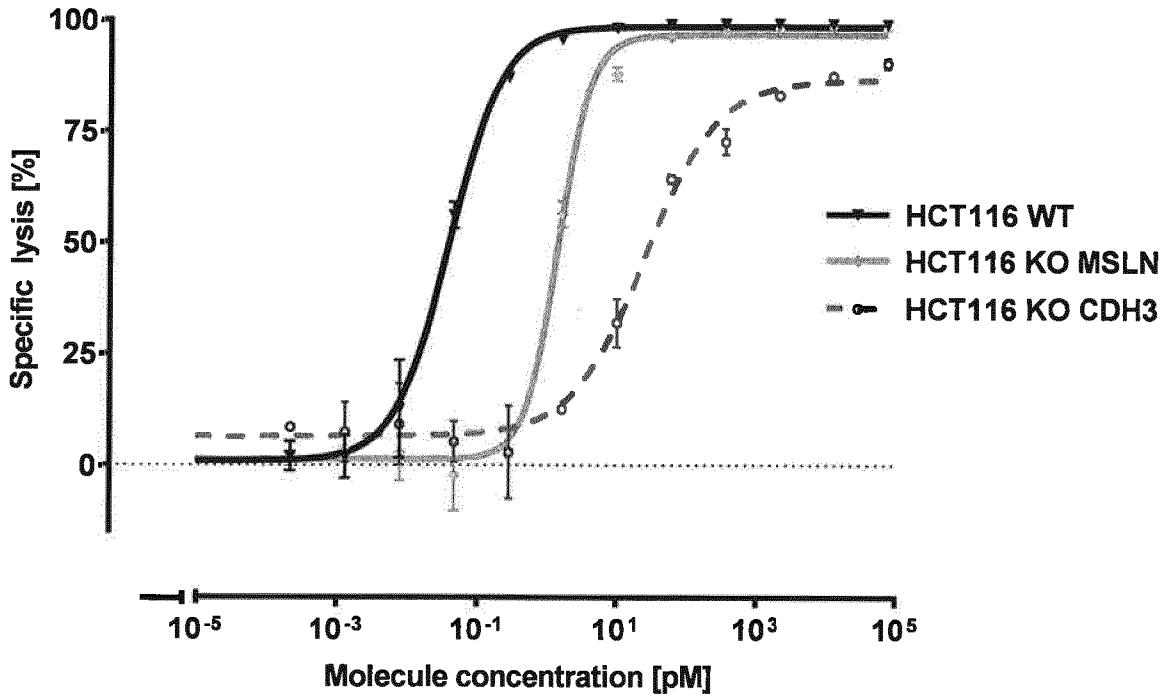


Fig. 5

A

CDH3 T-cell engager molecule 1

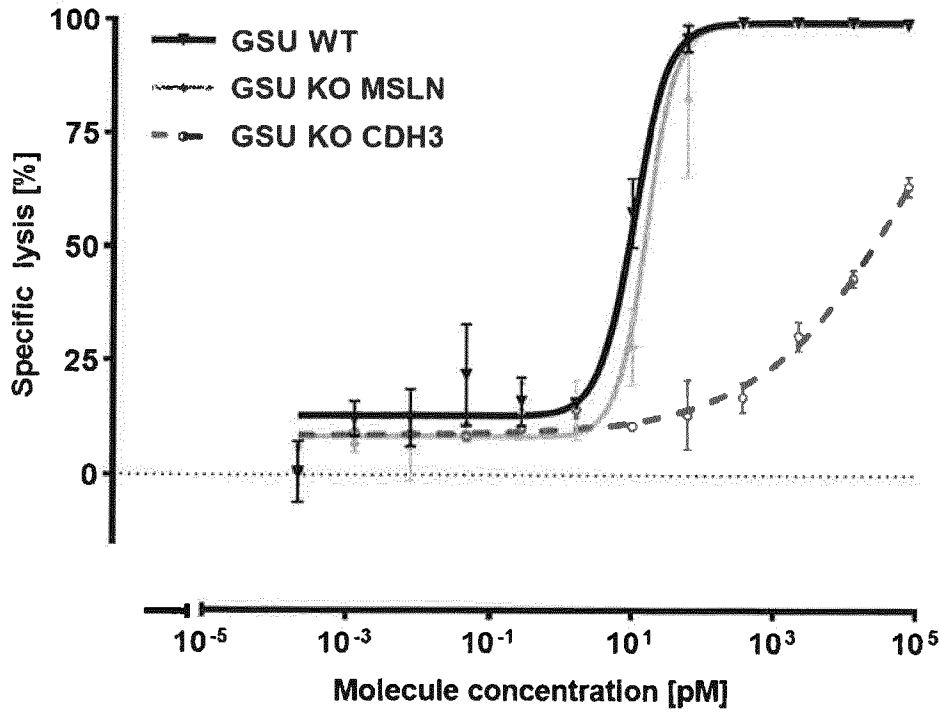
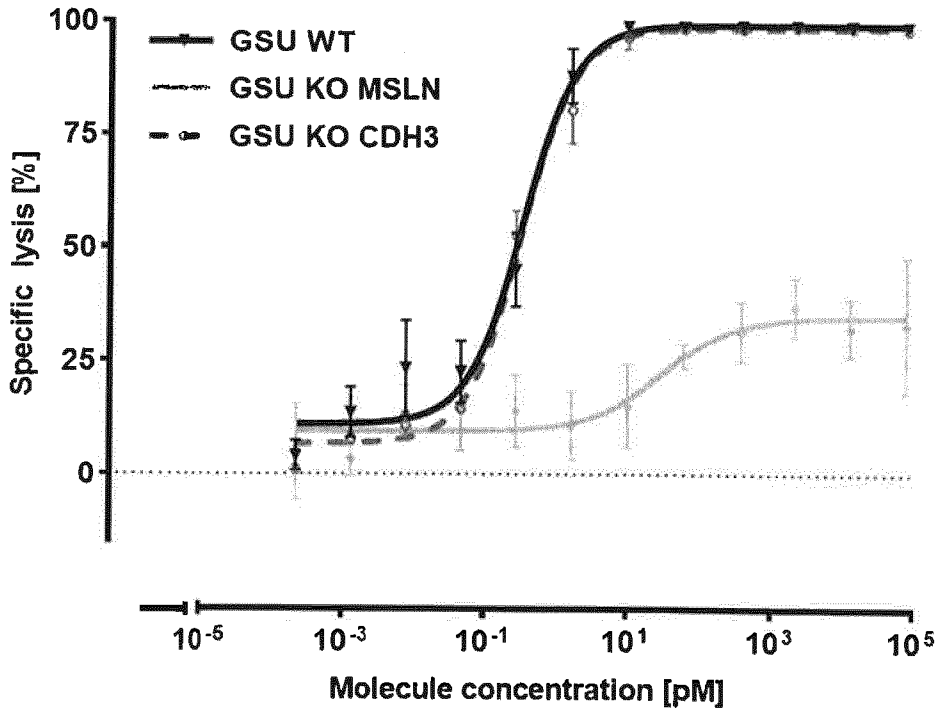


Fig. 5 (cont.)

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B

MSLN T-cell engager molecule 1



C

MSLN-CDH3 T-cell engager molecule 1

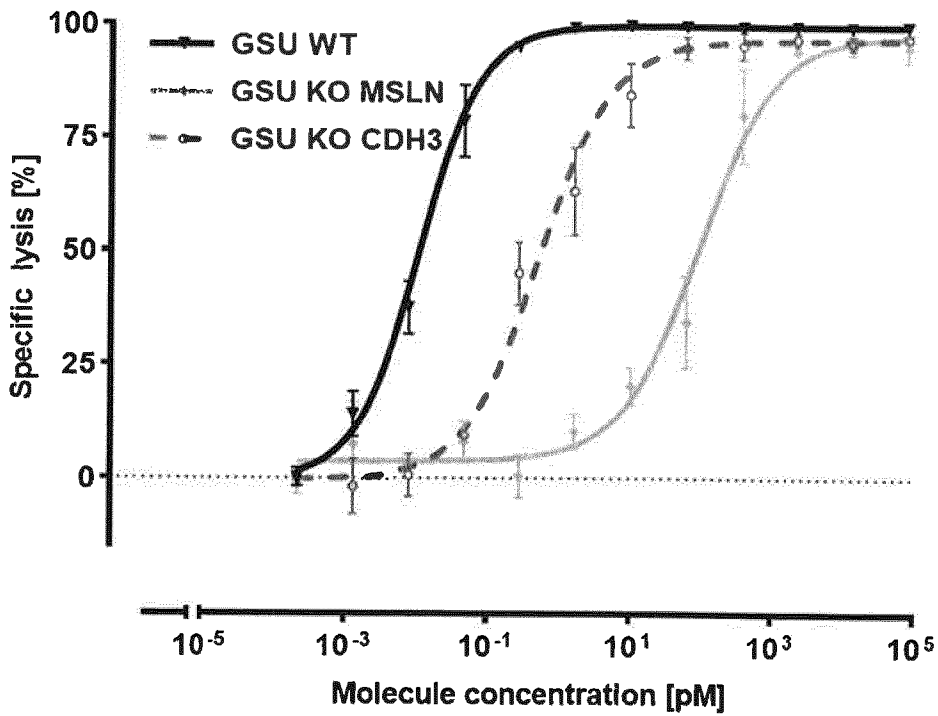
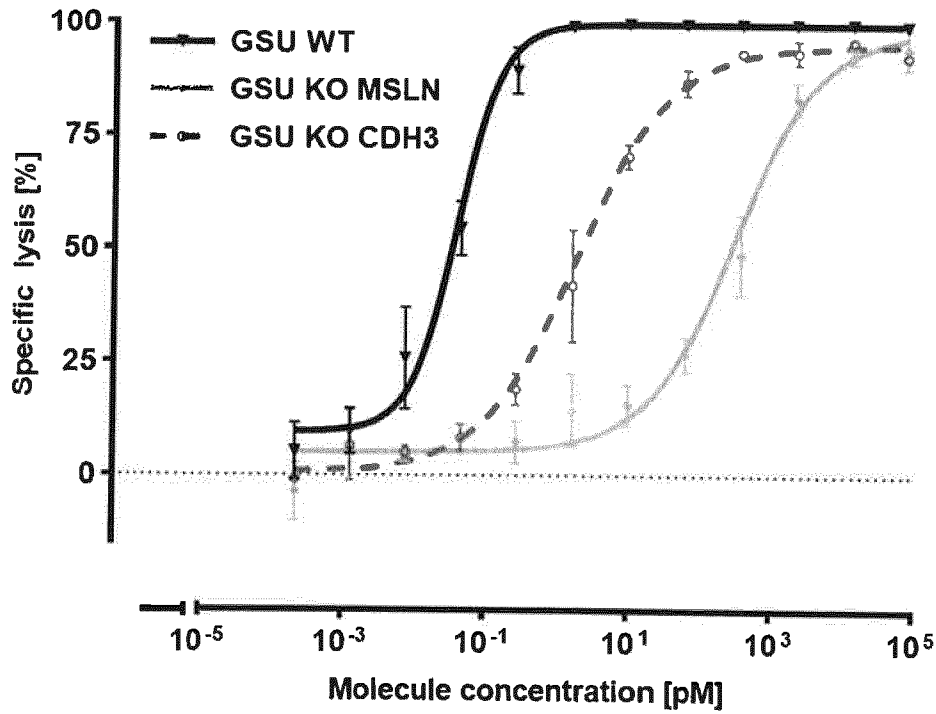


Fig. 5 (cont.)

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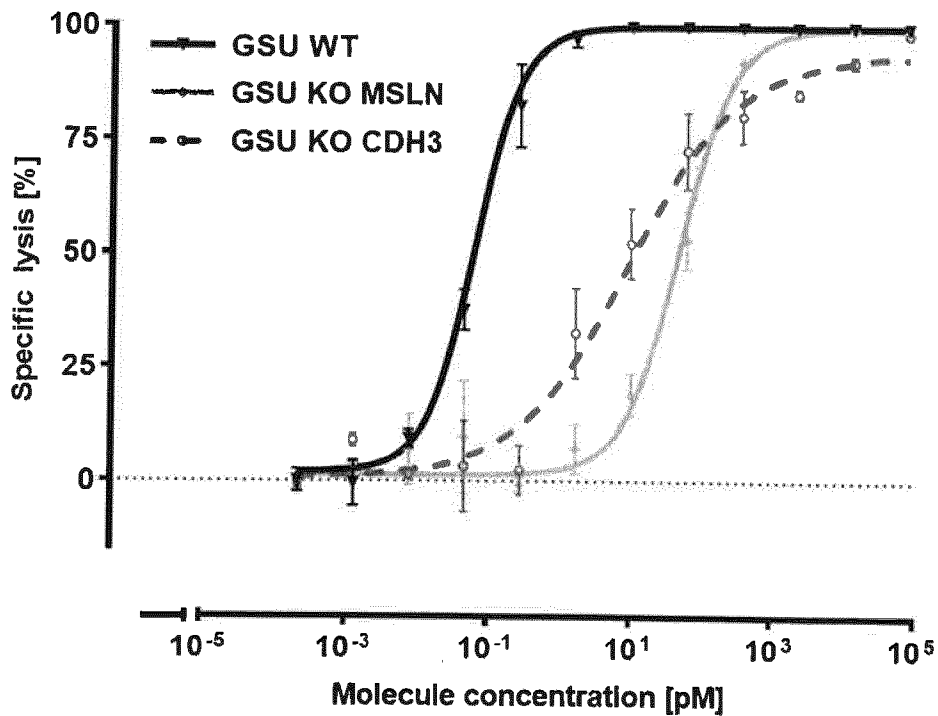
D

MSLN-CDH3 T-cell engager molecule 2



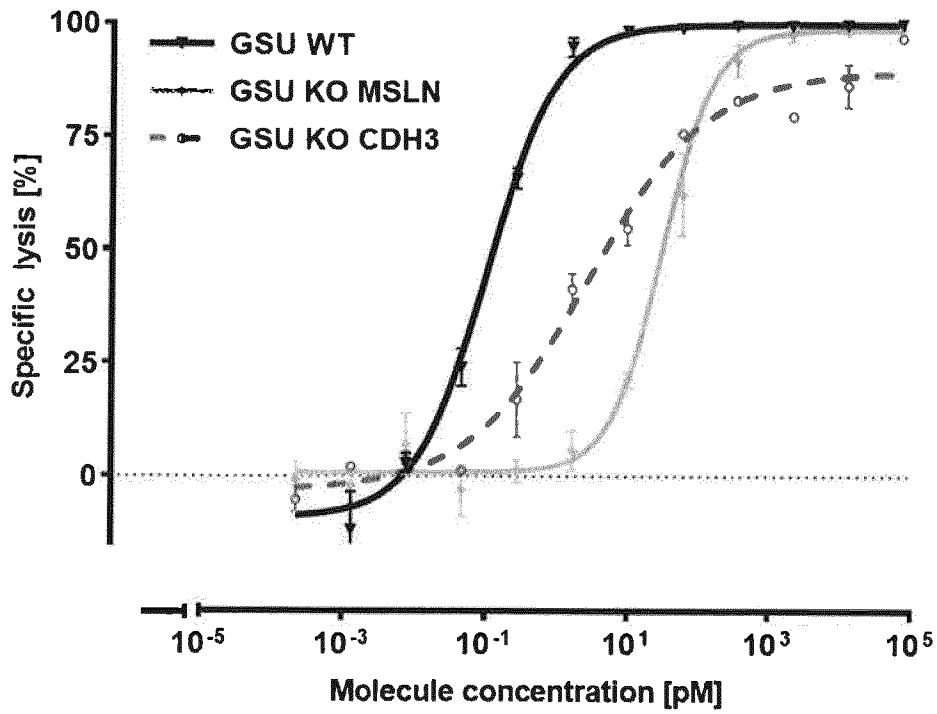
E

MSLN-CDH3 T-cell engager molecule 3



F

MSLN-CDH3 T-cell engager molecule 4



G

MSLN-CDH3 T-cell engager molecule 5

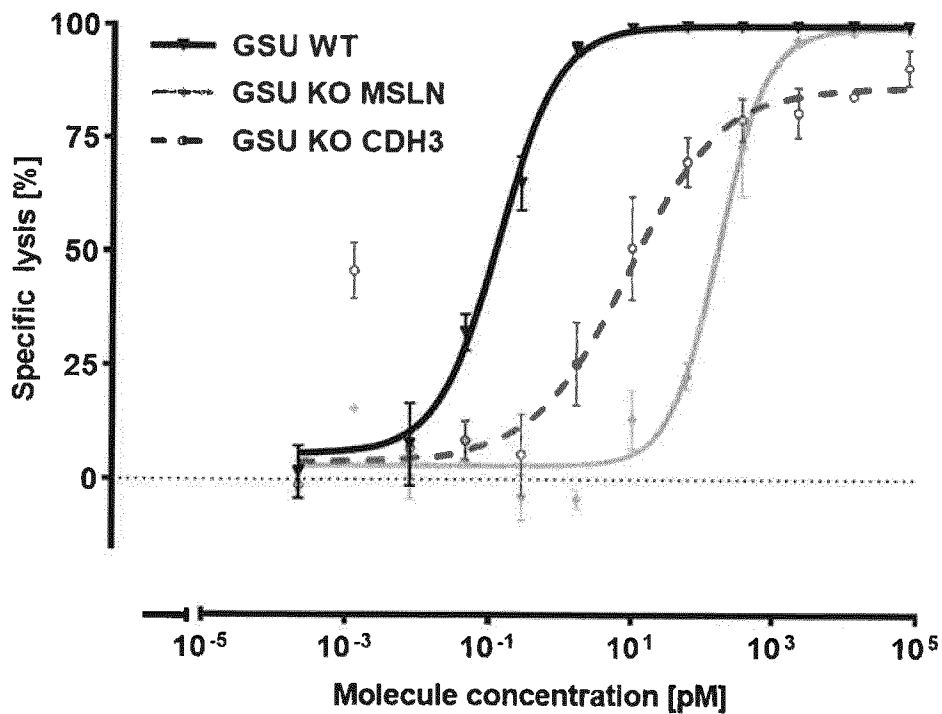
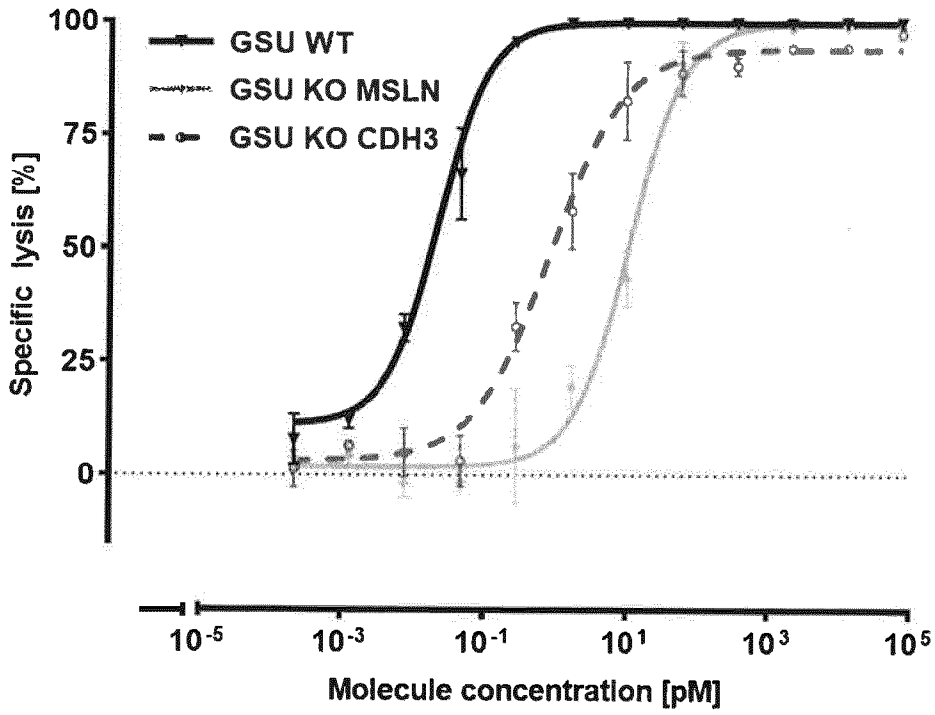


Fig. 5 (cont.)

H

MSLN-CDH3 T-cell engager molecule 6



I

MSLN-CDH3 T-cell engager molecule 7

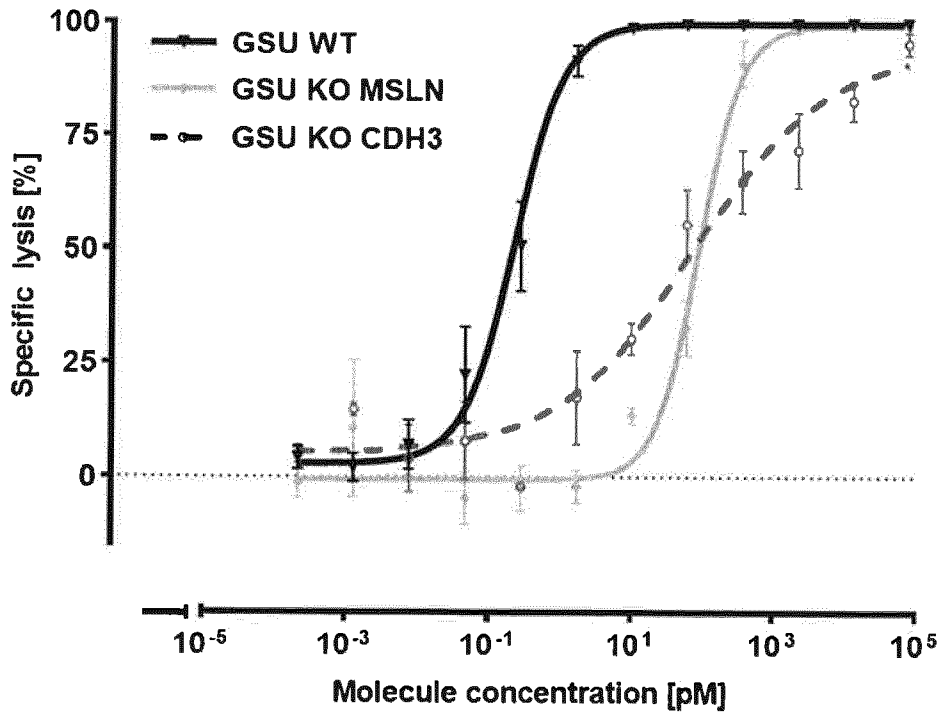


Fig. 6

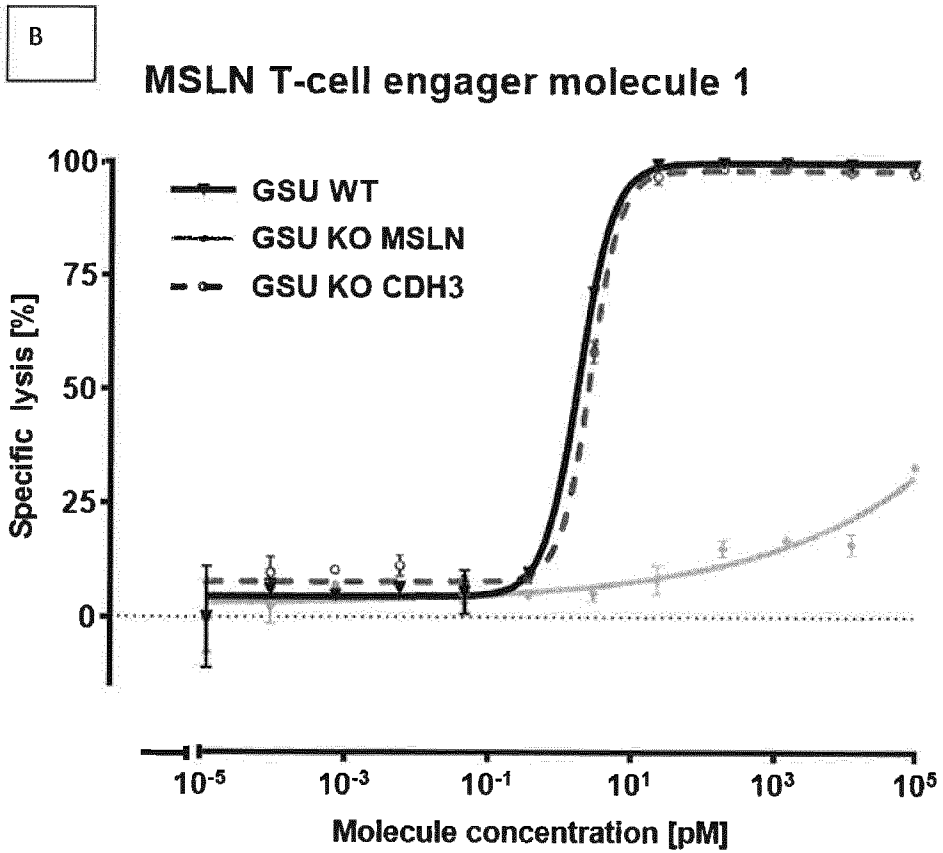
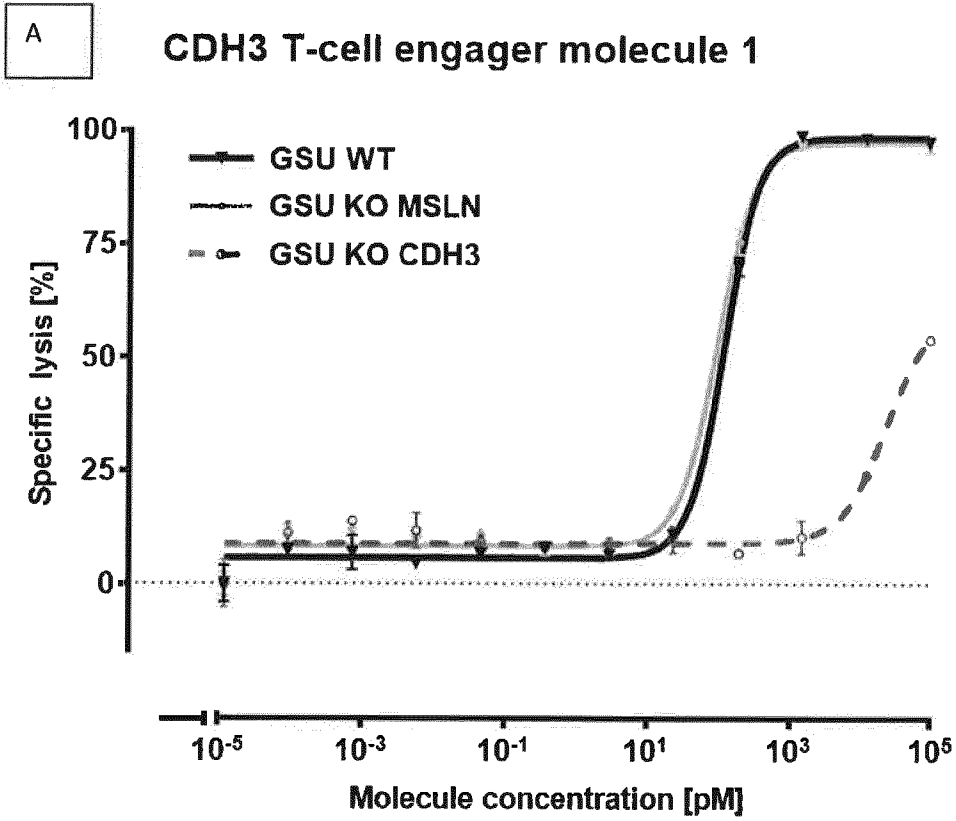
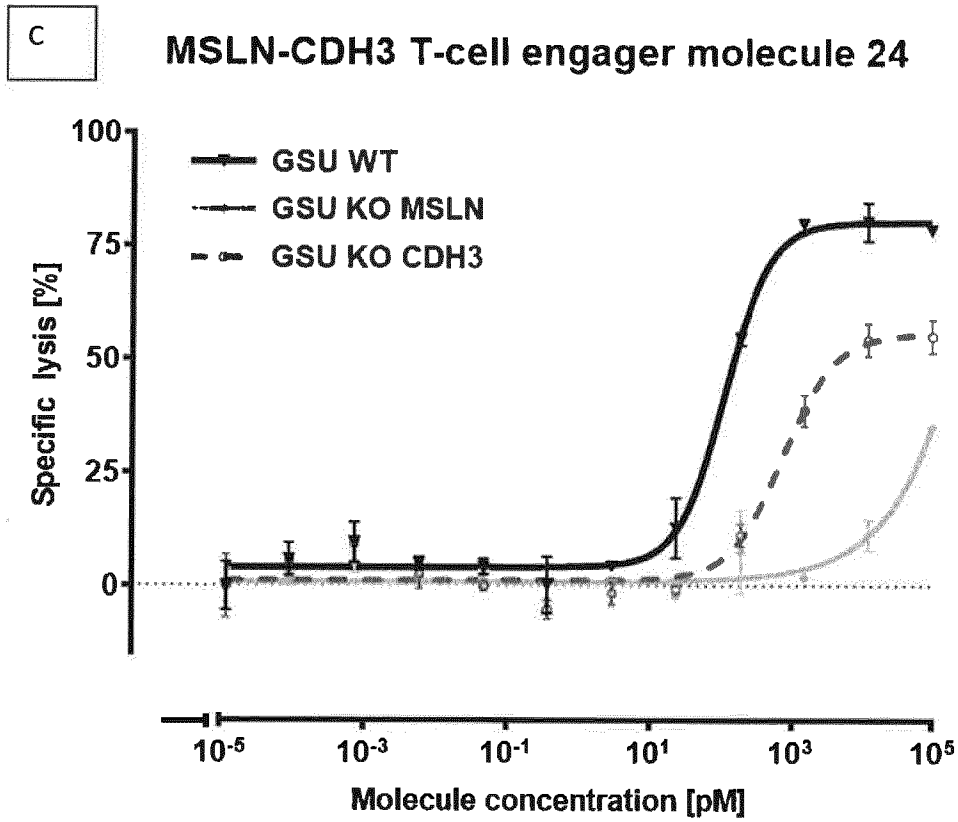


Fig. 6 (cont.)

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INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2023/062750

A. CLASSIFICATION OF SUBJECT MATTER
INV. C07K16/18 C07K16/28 C07K16/46 C07K16/30 C07K16/32
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
 Minimum documentation searched (classification system followed by classification symbols)
C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
EPO-Internal, BIOSIS, Sequence Search, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	<p>YOU GIHOON ET AL: "Bispecific Antibodies: A Smart Arsenal for Cancer Immunotherapies", VACCINES, vol. 9, no. 7, 2 July 2021 (2021-07-02), page 724, XP93000326, DOI: 10.3390/vaccines9070724 the whole document</p> <p align="center">----- -/--</p>	1-58

Further documents are listed in the continuation of Box C. See patent family annex.

* Special categories of cited documents :

"A" document defining the general state of the art which is not considered to be of particular relevance	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E" earlier application or patent but published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"O" document referring to an oral disclosure, use, exhibition or other means	"&" document member of the same patent family
"P" document published prior to the international filing date but later than the priority date claimed	

Date of the actual completion of the international search 29 August 2023	Date of mailing of the international search report 06/09/2023
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Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer Scheffzyk, Irmgard
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INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2023/062750

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	<p>UCKUN FATIH M ET AL: "A Clinical Phase 1B Study of the CD3xCD123 Bispecific Antibody APVO436 in Patients with Relapsed/Refractory Acute Myeloid Leukemia or Myelodysplastic Syndrome", CANCERS, 15 August 2021 (2021-08-15), XP93076592, DOI: 10.3390/cancers Retrieved from the Internet: URL:https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8394899/pdf/cancers-13-04113.pdf> figure 1</p> <p style="text-align: center;">-----</p>	1-58
Y	<p>G. HERNANDEZ-HOYOS ET AL: "MOR209/ES414, a Novel Bispecific Antibody Targeting PSMA for the Treatment of Metastatic Castration-Resistant Prostate Cancer", MOLECULAR CANCER THERAPEUTICS, vol. 15, no. 9, 12 July 2016 (2016-07-12), pages 2155-2165, XP055483056, US ISSN: 1535-7163, DOI: 10.1158/1535-7163.MCT-15-0242 figure 1A</p> <p style="text-align: center;">-----</p>	1-58
Y	<p>MEHTA NAVEEN K ET AL: "A novel IgG-based FLT3xCD3 bispecific antibody for the treatment of AML and B-ALL", JOURNAL FOR IMMUNOTHERAPY OF CANCER, vol. 10, no. 3, 1 March 2022 (2022-03-01), page e003882, XP93076599, DOI: 10.1136/jitc-2021-003882 Retrieved from the Internet: URL:https://jitc.bmj.com/content/jitc/10/3/e003882.full.pdf> figure 1A</p> <p style="text-align: center;">-----</p>	1-58
Y	<p>DAFNE M&#252;LLER ET AL: "Improved Pharmacokinetics of Recombinant Bispecific Antibody Molecules by Fusion to Human Serum Albumin", JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 282, no. 17, 27 April 2007 (2007-04-27), pages 12650-12660, XP055537278, US ISSN: 0021-9258, DOI: 10.1074/jbc.M700820200 figure 1</p> <p style="text-align: center;">-----</p> <p style="text-align: center;">-/--</p>	1-58

INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2023/062750

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
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