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(10) **Pub. No.: US 2025/0250362 A1**(43) **Pub. Date:** Aug. 7, 2025(54) **CYTOTOXICITY-INDUCING THERAPEUTIC AGENT**(71) Applicant: **Chugai Seiyaku Kabushiki Kaisha**, Tokyo (JP)(72) Inventors: **Junichi NEZU**, Shizuoka (JP); **Atsushi NARITA**, Shizuoka (JP); **Takahiro ISHIGURO**, Kanagawa (JP); **Mika SAKURAI**, Shizuoka (JP); **Hirotake SHIRAIWA**, Shizuoka (JP); **Naoka HIRONIWA**, Shizuoka (JP); **Tomoyuki IGAWA**, Shizuoka (JP); **Yumiko KAWAI**, Kanagawa (JP)(21) Appl. No.: **18/887,180**(22) Filed: **Sep. 17, 2024****Related U.S. Application Data**

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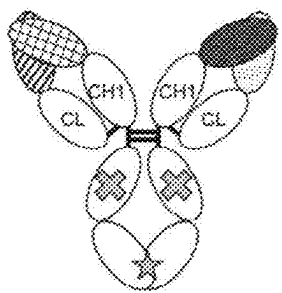
(57)

ABSTRACT

Novel multispecific antigen-binding molecules maintaining excellent cellular cytotoxicity and high stability, which comprise a domain that contains an antibody variable region having glycan 3-binding activity and a domain that contains an antibody variable region having T-cell receptor complex-binding activity, were discovered. Since the molecules of the present invention show a strong cytotoxicity against cells and tissues expressing glycan 3, it is possible to produce novel pharmaceutical compositions for treating or preventing various cancers.

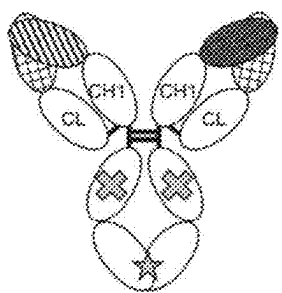
Specification includes a Sequence Listing.

Fig. 1A ERY22



- Anti-cancer antigen (GPC3) antibody H-chain variable region
- Anti-cancer antigen (GPC3) antibody L-chain variable region
- Anti-CD3 antibody H-chain variable region
- Anti-CD3 antibody L-chain variable region
- Antibody constant region
- ❖ Silent Fc mutation
- ★ Mutation for heteromeric Fc association

Fig. 1B ERY27



- Anti-cancer antigen (GPC3) antibody H-chain variable region
- Anti-CD3 antibody H-chain variable region
- Common L-chain variable region
- Antibody constant region
- ❖ Silent Fc mutation
- ★ Mutation for heteromeric Fc association

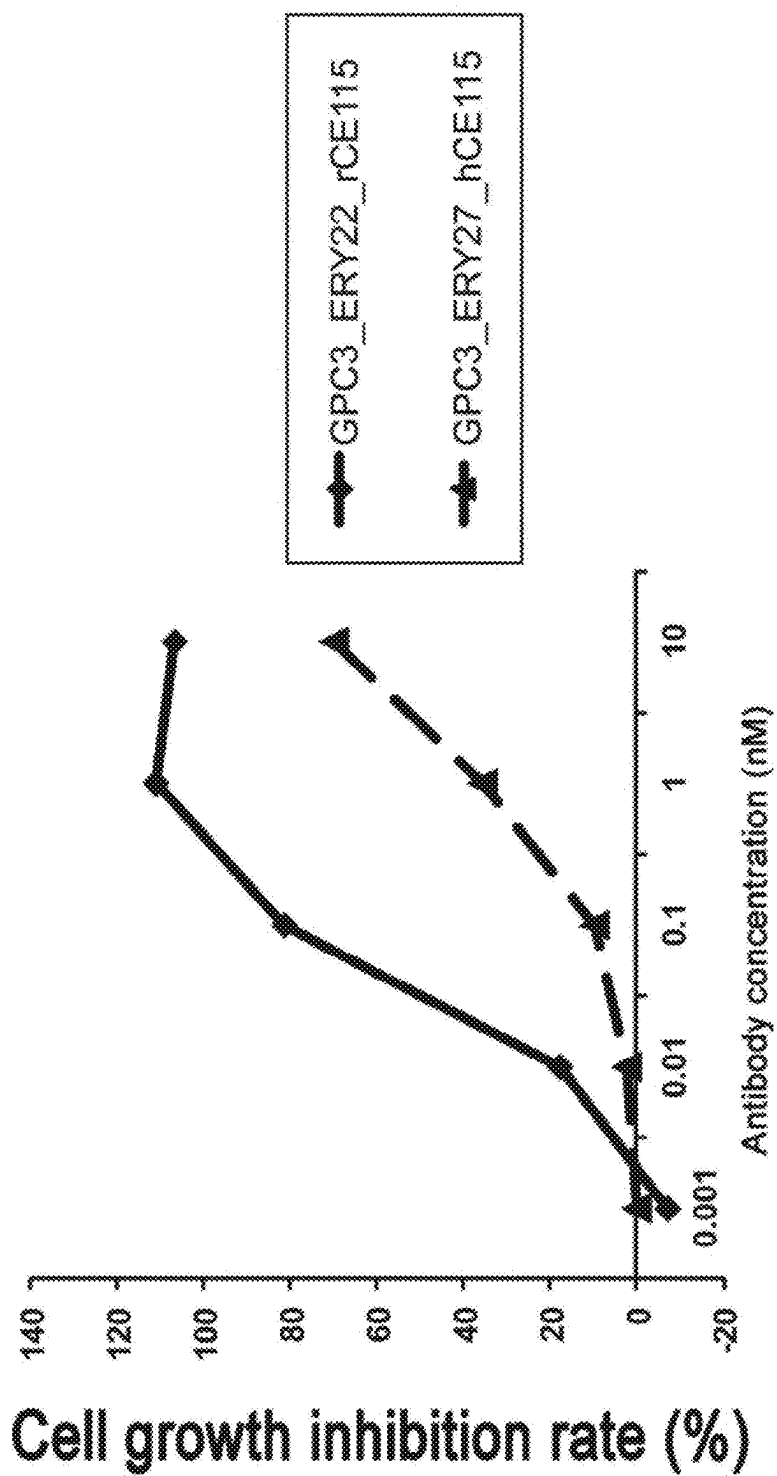


Fig. 2

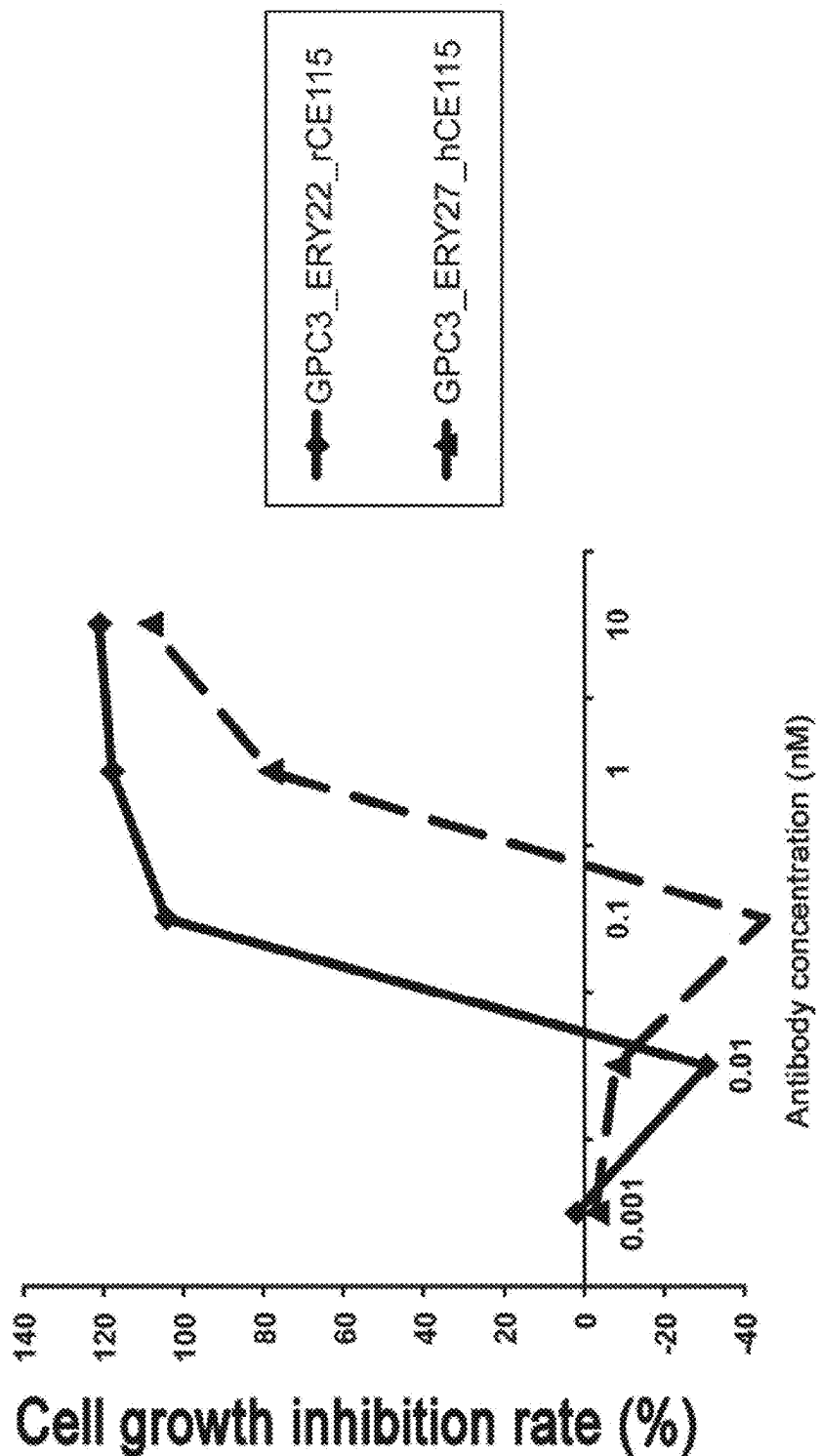


Fig. 3

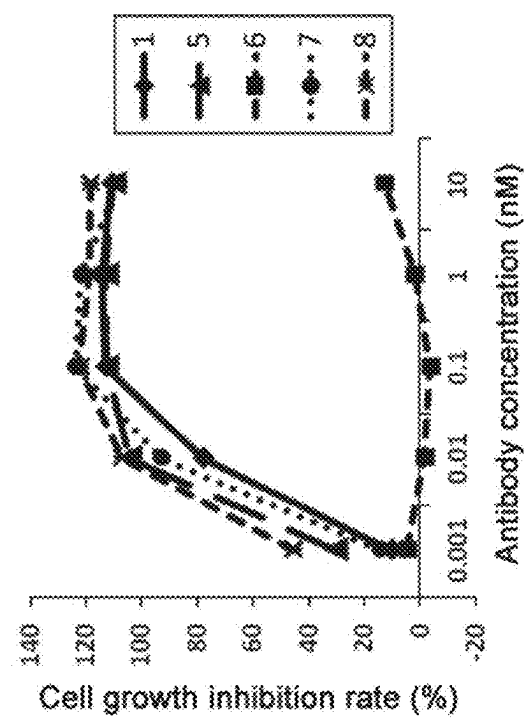


Fig. 4B

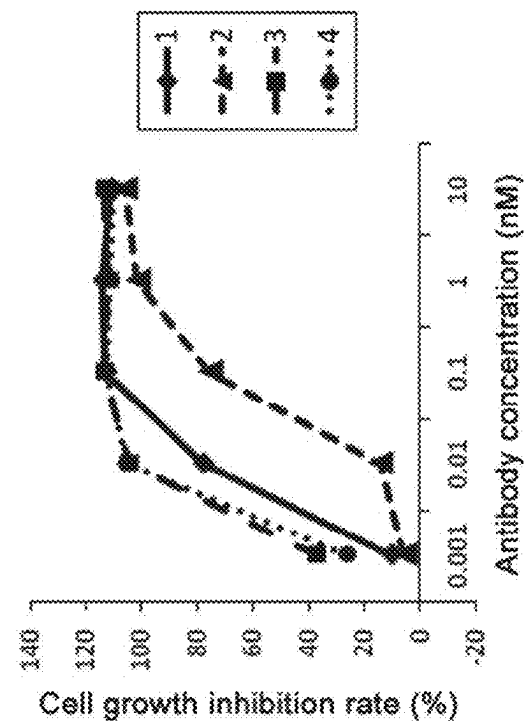


Fig. 4A

Fig. 5A

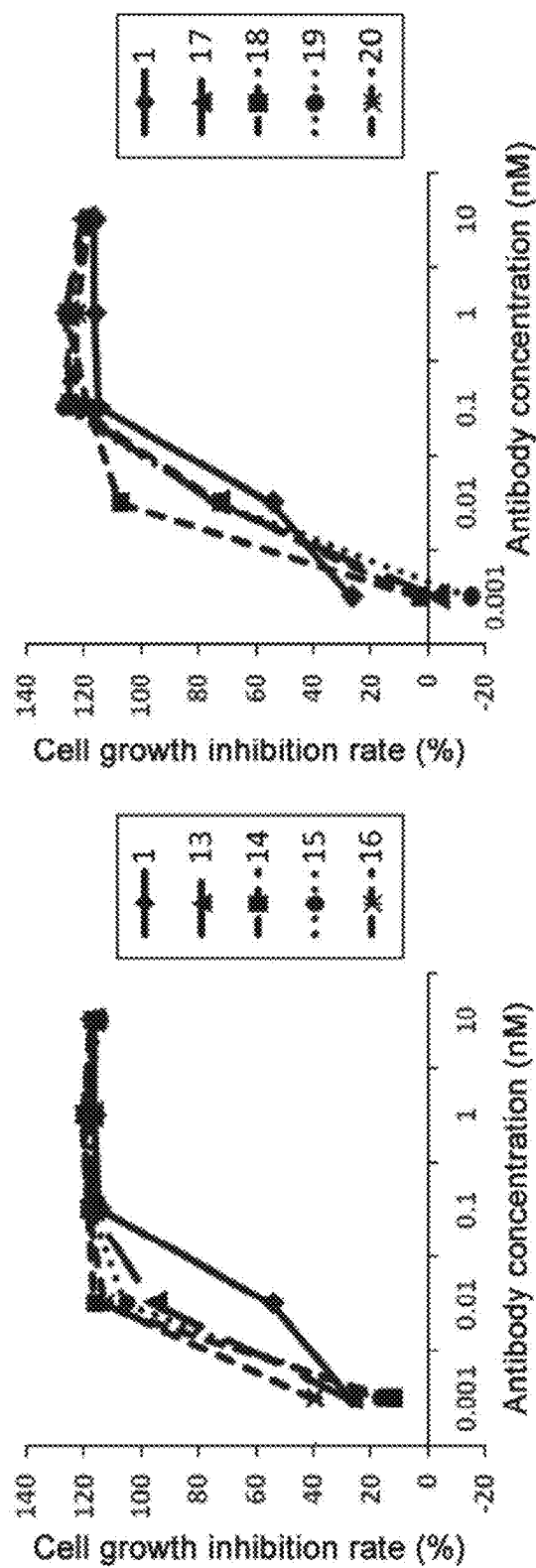


Fig. 5B

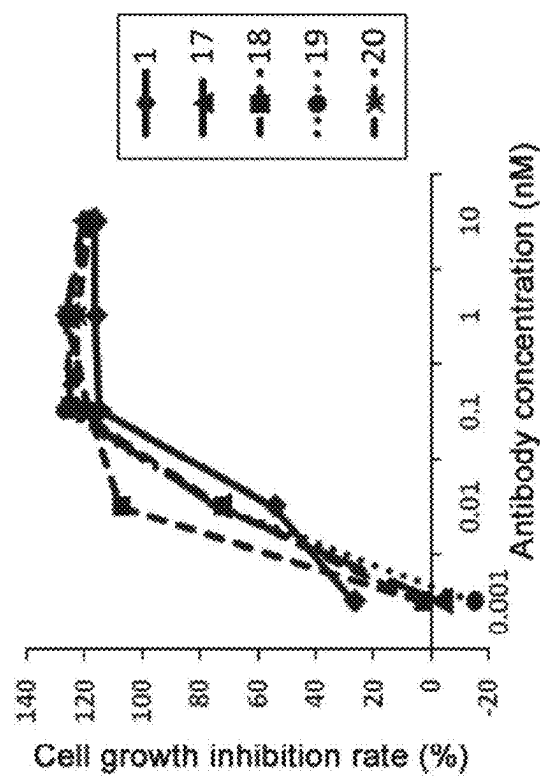


Fig. 6A

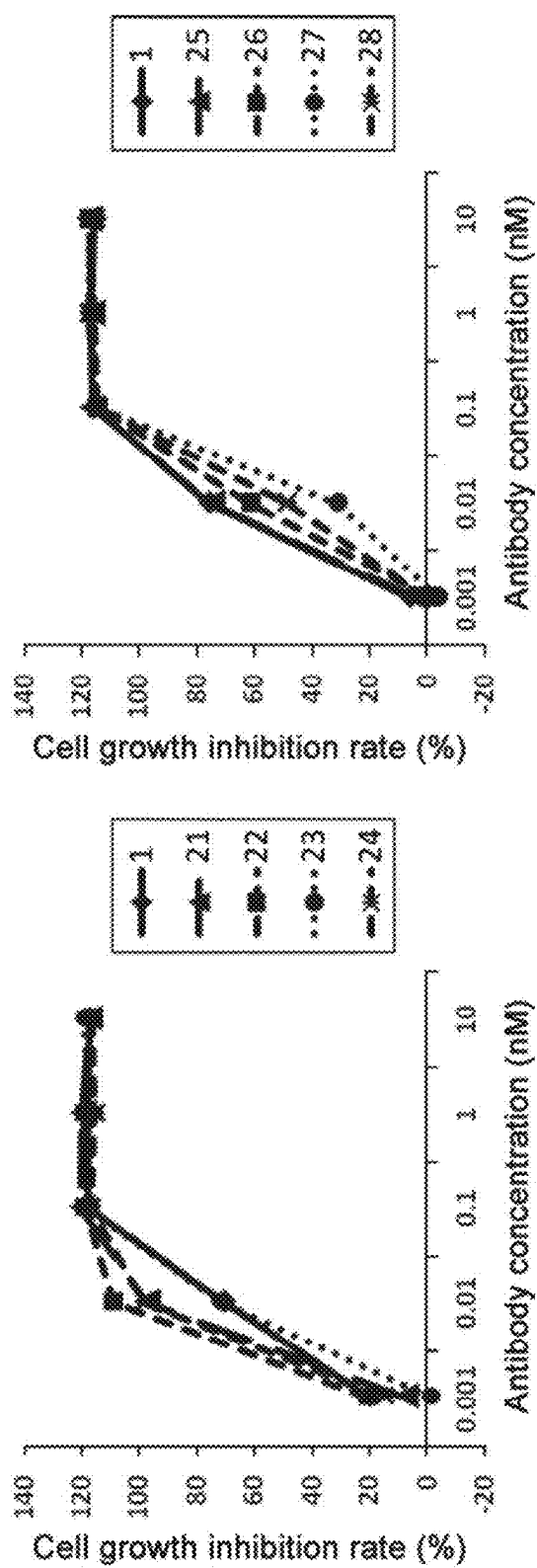


Fig. 6B

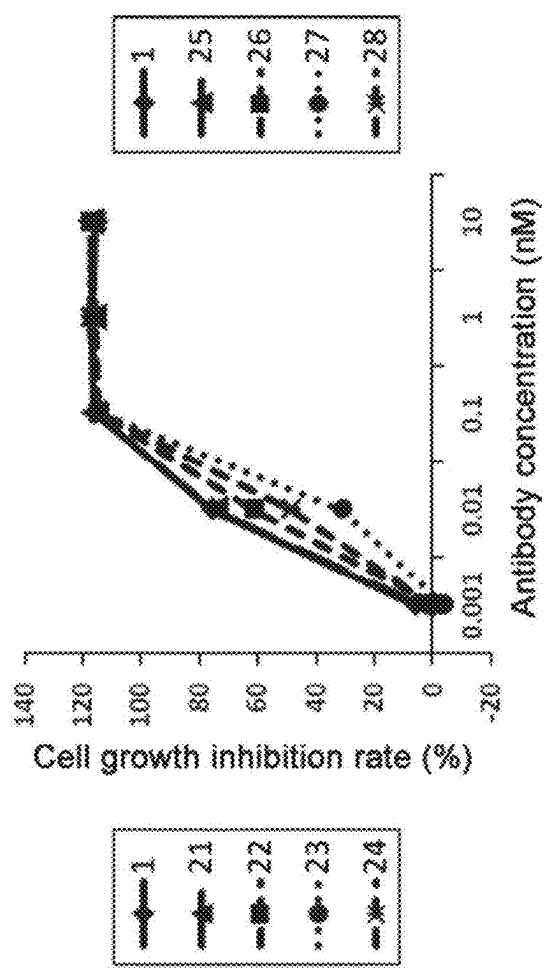


Fig. 7A

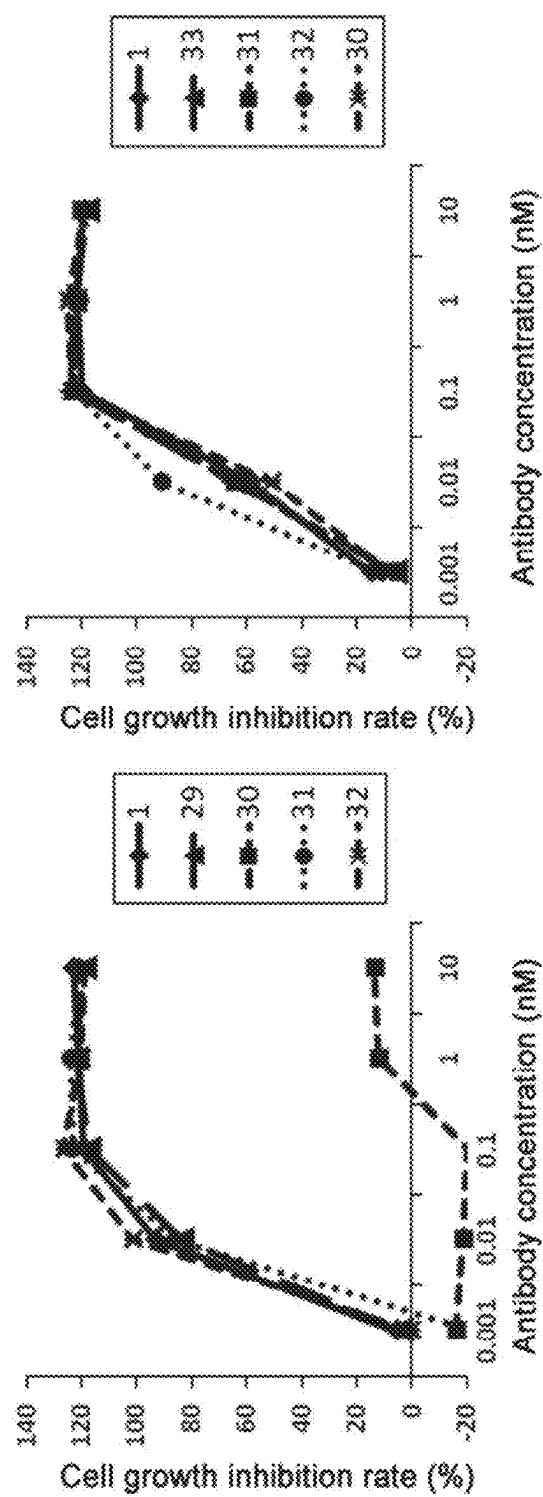


Fig. 7B

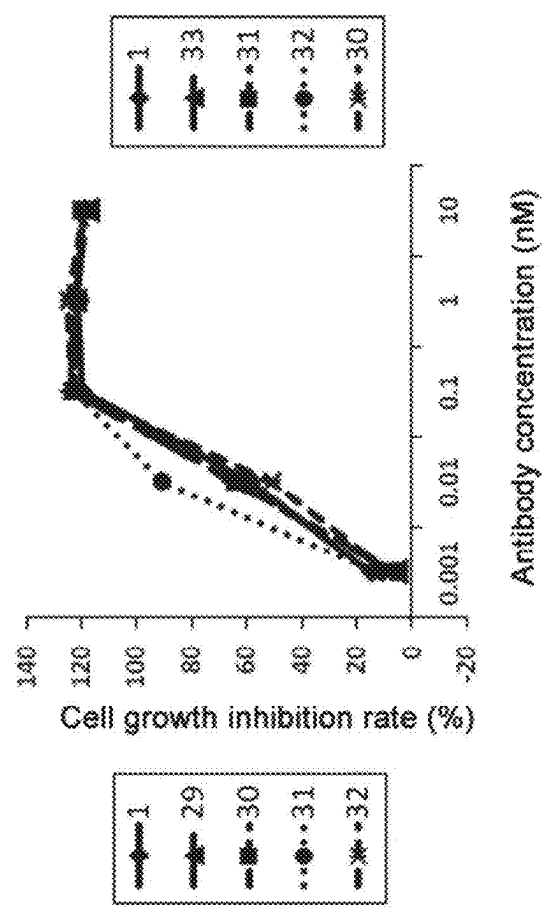


Fig. 8A

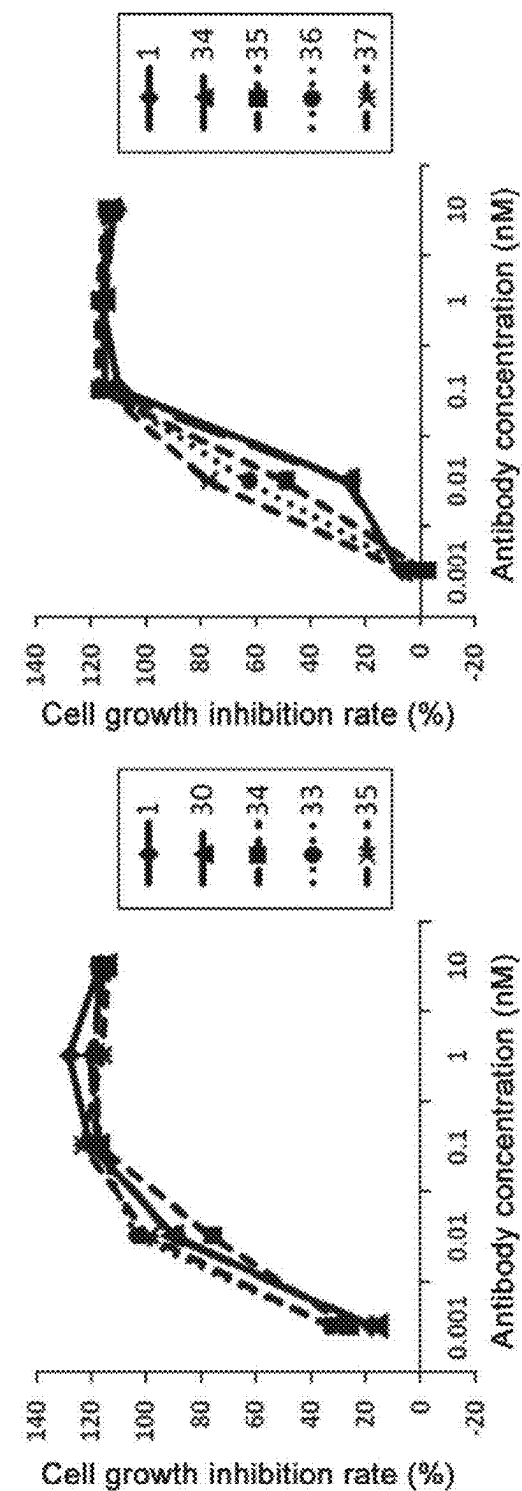


Fig. 8B

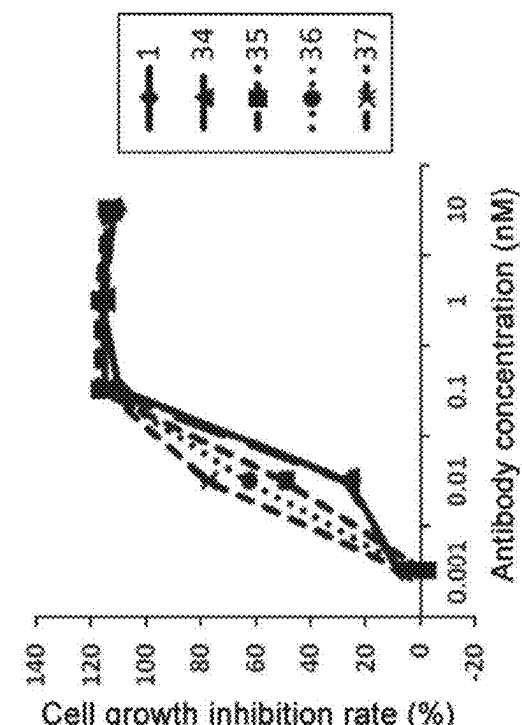


Fig. 9A

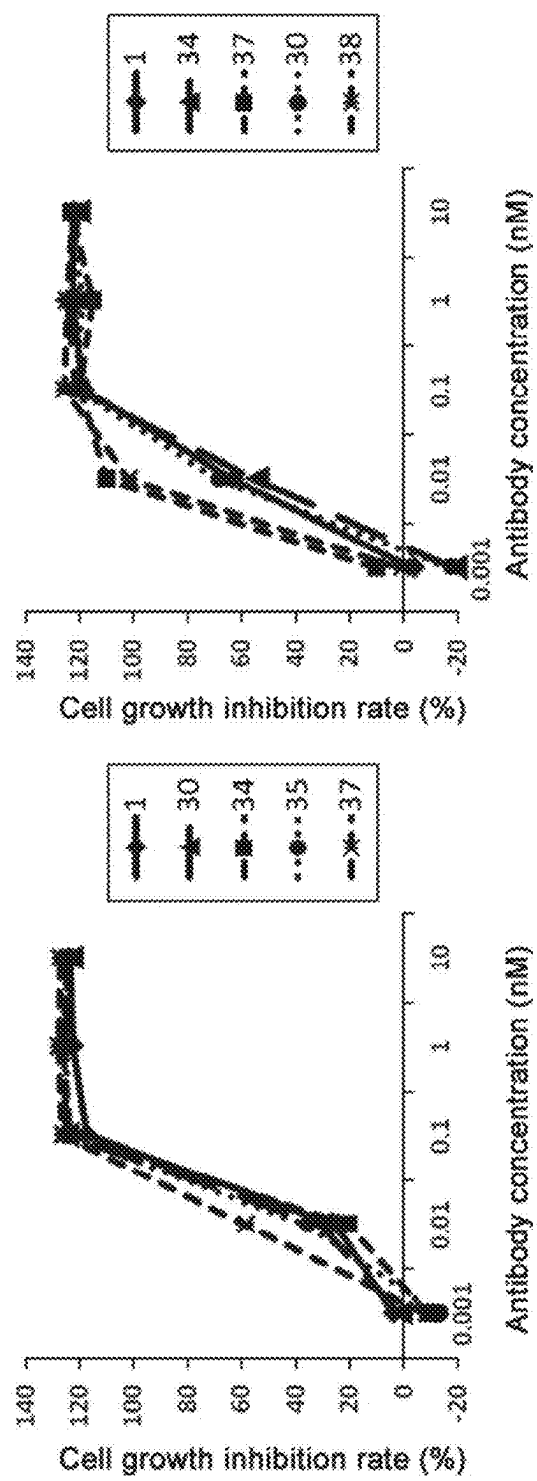


Fig. 9B

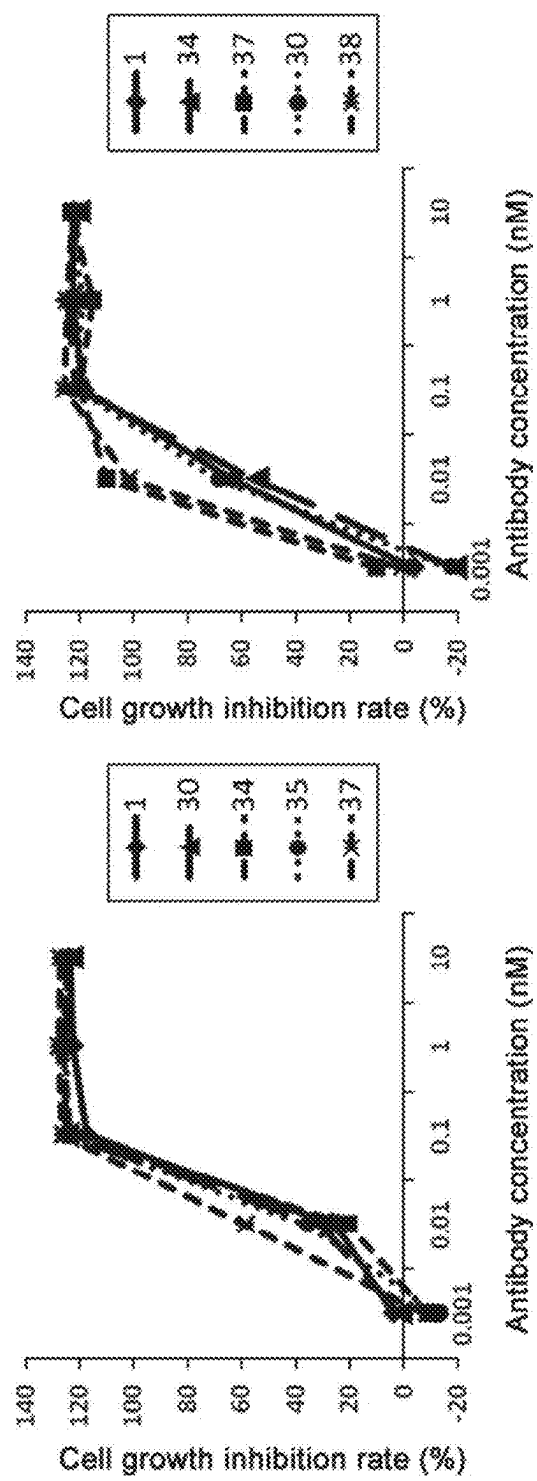


Fig. 10A

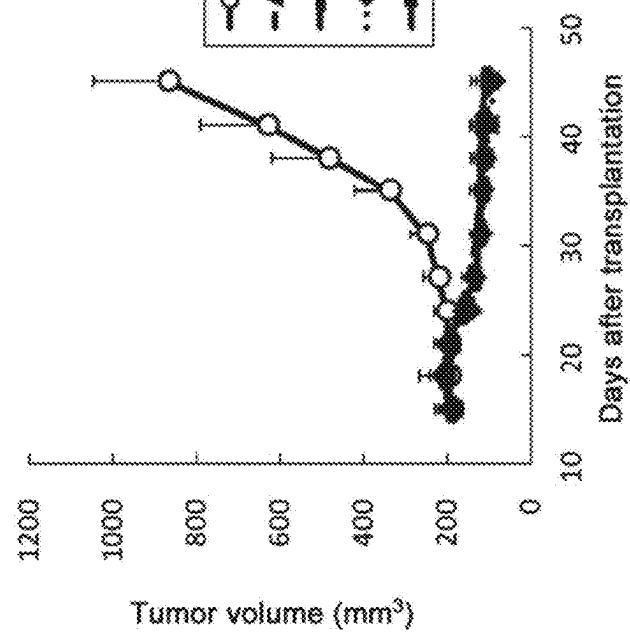
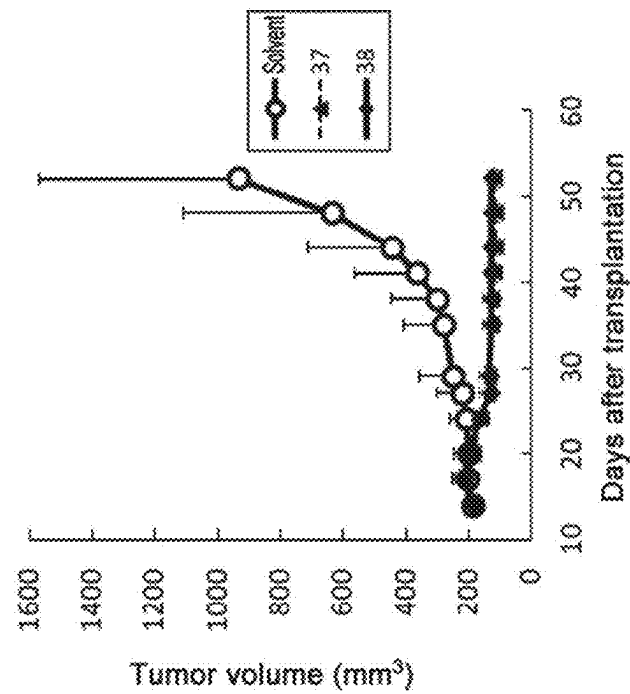
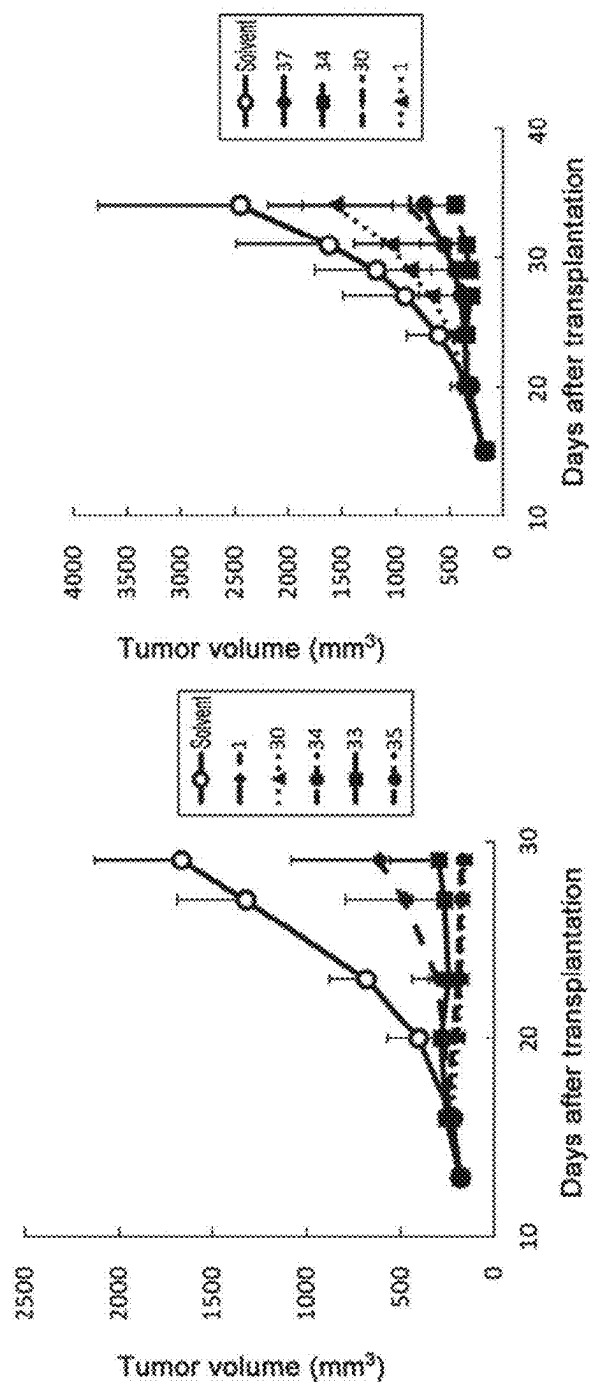


Fig. 10B



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Fig. 12

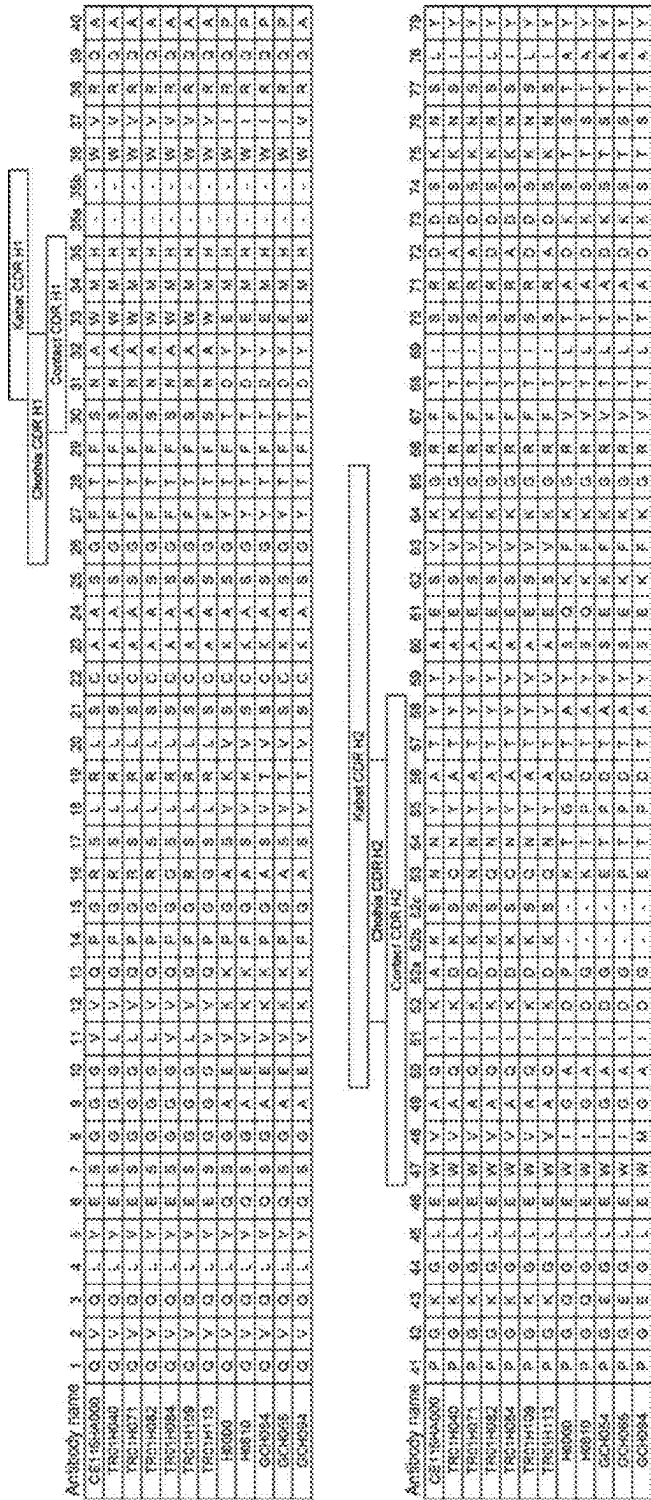


Fig. 13-1

Antibody name	Kabat CDR HS																		
	80	81	82	82a	82b	82c	83	84	85	85	87	88	89	89	89	89	89	89	89
GE11644000	L	G	N	S	L	K	T	E	O	T	A	V	C	R	Y	H	Y	G	T
TRCH1040	L	G	N	S	L	K	T	E	O	T	A	V	C	R	Y	H	Y	G	T
TSR1H071	L	O	N	S	L	K	T	E	D	T	A	V	Y	C	R	Y	H	Y	G
TSR1H092	L	O	N	S	L	K	T	E	D	T	A	V	Y	C	R	Y	H	Y	G
TRCH1084	L	O	N	S	L	K	T	E	D	T	A	V	Y	C	R	Y	H	Y	G
TSR1H108	L	O	N	S	L	K	T	E	D	T	A	V	Y	C	R	Y	H	Y	G
TSR1H113	L	O	N	S	L	K	T	E	D	T	A	V	Y	C	R	Y	H	Y	G
HR090	M	E	L	S	S	L	T	S	E	D	T	A	V	Y	C	T	R	F	Y
HR0910	M	E	L	S	S	L	T	S	E	D	T	A	V	Y	C	T	R	F	Y
GC14054	M	E	L	S	S	L	T	S	E	D	T	A	V	Y	C	T	R	F	Y
GC15285	M	E	L	S	S	L	T	S	E	D	T	A	V	Y	C	T	R	F	Y
GC094	M	E	L	S	S	L	T	S	E	D	T	A	V	Y	C	T	R	F	Y
Cysteine CDR HS																			
Contact CDR HS																			
GE11644000	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H040	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H071	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H092	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TRCH1084	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H108	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TRCH113	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
HR090	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
HR0910	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
GC14054	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
GC15285	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
GC094	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S

Antibody name	Kabat CDR HS																		
	109	110	111	112	113	114	115	116	117	118	119	120	121	122	123	124	125	126	127
GE11644000	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H040	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H071	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H092	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TRCH1084	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TSR1H108	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
TRCH113	T	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
HR090	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
HR0910	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
GC14054	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
GC15285	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S
GC094	L	V	T	V	S	S	S	S	S	S	S	S	S	S	S	S	S	S	S

Fig. 13-2

Antibody name	Kabat CDR L1		Contact CDR L1		Contact CDR L3	
	C	R	C	R	C	R
1.0090	O	-	V	-	S	-
1.0091	D	-	V	-	S	-
1.0091:	D	-	V	-	S	-
1.0201	D	-	V	-	S	-
1.0203	D	-	V	-	S	-
1.0204	D	-	V	-	S	-
1.0206	D	-	V	-	S	-
1.0208	D	-	V	-	S	-
1.0209	D	-	V	-	S	-
1.0211:	D	-	V	-	S	-
1.0212	D	-	V	-	S	-
1.0222	D	-	V	-	S	-
1.0272	D	-	V	-	S	-
Antibody name	Kabat CDR L2		Contact CDR L2		Contact CDR L4	
	C	R	C	R	C	R
1.0090	G	O	K	E	G	A
1.0091:	G	O	K	P	G	A
1.0201	G	O	K	P	G	A
1.0203	G	O	K	P	G	A
1.0204	G	O	K	P	G	A
1.0206	G	O	K	P	G	A
1.0208	G	O	K	P	G	A
1.0209	G	O	K	P	G	A
1.0211:	G	O	K	P	G	A
1.0212	G	O	K	P	G	A
1.0222	G	O	K	P	G	A
1.0272	G	O	K	P	G	A
Antibody name	Kabat CDR L3		Contact CDR L3		Contact CDR L5	
	C	R	C	R	C	R
1.0090	E	A	E	D	V	G
1.0091:	E	A	E	D	V	G
1.0201	E	A	E	D	V	G
1.0203	E	A	E	D	V	G
1.0204	E	A	E	D	V	G
1.0206	E	A	E	D	V	G
1.0208	E	A	E	D	V	G
1.0209	E	A	E	D	V	G
1.0211:	E	A	E	D	V	G
1.0212	E	A	E	D	V	G
1.0222	E	A	E	D	V	G
1.0272	E	A	E	D	V	G

Fig. 14

CYTOTOXICITY-INDUCING THERAPEUTIC AGENT

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application is a continuation of U.S. application Ser. No. 17/225,273, filed on Apr. 8, 2021, which is a divisional of U.S. application Ser. No. 15/963,221, filed on Apr. 26, 2018, now U.S. Pat. No. 11,001,643 B2, issued on May 11, 2021, which is a divisional of U.S. application Ser. No. 15/467,654, filed on Mar. 23, 2017, now U.S. Pat. No. 9,975,966 B2, issued on May 22, 2018, which is a continuation of International Application No. PCT/JP2015/077024, filed on Sep. 25, 2015, which claims the benefit of Japanese Application No. 2014-197315, filed on Sep. 26, 2014, the contents of which are hereby incorporated by reference in their entireties.

REFERENCE TO SEQUENCE LISTING SUBMITTED ELECTRONICALLY

[0002] The content of the electronically submitted sequence listing (Name: 6663_0311_Sequence_Listing.xml; Size: 594 kilobytes; and Date of Creation: Aug. 16, 2024) filed with the application is incorporated herein by reference in its entirety.

TECHNICAL FIELD

[0003] The present invention relates to multispecific antigen-binding molecules, uses thereof, and such.

BACKGROUND ART

[0004] Antibodies are drawing attention as pharmaceuticals because of their high stability in plasma and few adverse reactions (Non-patent Documents 1 and 2). Antibodies are known to induce not only an antigen-binding action, an agonistic action, and an antagonistic action, but also effector-mediated cytotoxic activities (also called effector functions) such as antibody-dependent cellular cytotoxicity (ADCC), antibody dependent cell phagocytosis (ADCP), and complement-dependent cytotoxicity (CDC), and exhibit antitumor effects against cancer cells (Non-Patent Document 3). ADCC is a cytotoxicity exhibited by effector cells against antibody-bound target cancer cells via binding of the antibody Fc region to an Fc receptor present on effector cells such as NK cells and macrophages. A complement complex binds to the complement-binding site present in an antibody structure. CDC is cell injury that results from cell destruction where an influx of water and ions into cells is promoted by pore formation on the cell membrane of the antibody-bound cells by complement components present in the complex. A number of therapeutic antibodies showing excellent anti-tumor effects have been developed as pharmaceuticals for cancer treatment (Non-patent Document 4); and while existing therapeutic antibodies have shown excellent actions, the therapeutic outcome achieved by administration of these antibodies is still not satisfactory.

[0005] For an antibody to express ADCC, ADCP, and CDC, it is necessary for the antibody Fc region, the antibody receptor (FcγR) present on effector cells such as NK cells and macrophages, and various complement components to bind. In humans, isoforms of FcγRIa, FcγRIIa, FcγRIIb, FcγRIIIa, and FcγRIIIb have been reported as the FcγR protein family, and the respective allotypes have been

reported as well (Non-patent Document 5). Among these isoforms, FcγRIa, FcγRIIa, and FcγRIIIa carry a domain called the Immunoreceptor Tyrosine-based Activation Motif (ITAM) in the intracellular domain, and transmit activation signals. On the other hand, only FcγRIIb carries a domain called the Immunoreceptor Tyrosine-based Inhibitory Motif (ITIM) in the intracellular domain, and transmits inhibitory signals. Every one of the FcγRs is known to transmit signals via crosslinking by immune complexes and such (Non-patent Document 6). When antibodies actually exert an effector function on cancer cells, FcγRs on the effector cell membrane form clusters at the Fc regions of several antibodies bound on the cancer cell membrane, and activation signals are transmitted by effector cells. A cytoidal effect is exerted as a result, but since FcγRs are crosslinked only in effector cells present near cancer cells this time, activation of immunity is shown to occur locally in cancer cells (Non-patent Document 7).

[0006] Naturally-occurring immunoglobulins bind to antigens at their variable regions, and bind to receptors such as FcγR, FcRn, FcαR, and FcεR, and complements at their constant regions. FcRn is one of the binding molecules that interact at the IgG Fc region, and since each of the antibody heavy chains binds one molecule of FcRn, two molecules of FcRn have been reported to bind one IgG-type antibody molecule. However, unlike FcRn and such, FcγR interacts at the antibody hinge region and CH2 domain, and only one molecule of FcγR binds to one molecule of IgG-type antibody (Non-patent Document 8). Furthermore, a common naturally-occurring IgG-type antibody recognizes and binds a single epitope via its variable region (Fab); therefore, it can bind to only one antigen. On the other hand, many types of proteins are known to be involved in cancer and inflammation, and there may be crosstalk among the proteins. For example, several inflammatory cytokines (TNF, IL1, and IL6) are known to be involved in immunological diseases (Non-patent Document 9). Furthermore, activation of other receptors is known as one of the mechanisms of cancer in acquiring drug resistance (Non-patent Document 10). In such cases, common antibodies that recognize a single epitope would be unable to inhibit multiple proteins.

[0007] Antibodies (bispecific antibodies) that bind to two or more types of antigens with one molecule are being studied as molecules that inhibit multiple targets. It is possible to confer binding activities to two different antigens (a first antigen and a second antigen) by modifying naturally-occurring IgG-type antibodies (Non-patent Document 11). Accordingly, there will not only be neutralization of two or more types of antigens by a single molecule, but also enhancement of antitumor activity due to crosslinks between cells having cytotoxic activity and cancer cells. As molecular forms of a bispecific antibody, a molecule comprising an antigen-binding site added to the N or C terminus of an antibody (DVD-Ig and scFv-IgG), a molecule having different sequences for the two Fab regions of an antibody (common L-chain bispecific antibody and hybrid hybridoma), a molecule in which one Fab region recognizes two antigens (two-in-one IgG), and a molecule having a CH3 region loop site as a new antigen-binding site (Fcab) have been reported so far (Non-patent Documents 12 and 13). Since all bispecific antibodies interact at their Fc regions with FcγR, antibody effector functions are preserved. Thus, the bispecific antibody binds to any antigen

that it recognizes and at the same time binds to Fc γ R, and exhibits ADCC activity against cells expressing the antigen.

[0008] If all the antigens recognized by the bispecific antibody are antigens specifically expressed in cancer, the bispecific antibody exhibits cytotoxic activity to cancer cells when it binds to any of the antigens. Therefore, in comparison to a conventional antibody pharmaceutical that recognizes one antigen, a more efficient antitumor effect can be expected from such an antibody. However, in the case where any one of the antigens recognized by the bispecific antibody is expressed in normal tissues or cells expressed on immunocytes, damage on normal tissues or release of cytokines occurs due to crosslinking with Fc γ R (Non-patent Document 14). As a result, strong adverse reactions are induced.

[0009] A T-cell redirecting antibody that employs cytotoxicity mobilizing T cells as effector cells as the mechanism for its antitumor effect has been known from the 1980s as a bispecific antibody (Non-patent Documents 15, 16, and 17). Unlike antibodies that employ ADCC mobilizing NK cells or macrophages as effector cells as the mechanism for their antitumor effects, a T-cell redirecting antibody is an antibody against any one of the subunits constituting the T-cell receptor (TCR) complex on T cells, and is specifically a bi-specific antibody comprising an antibody that binds to the CD3 epsilon chain and an antibody that binds to an antigen on the target cancer cell. T cells come close to cancer cells via simultaneous binding of the CD3 epsilon chain and a cancer antigen by a T-cell redirecting antibody. As a result, antitumor effects against cancer cells are considered to be exerted through the cytotoxic activity possessed by T cells.

[0010] Catumaxomab, which is known as a T-cell redirecting antibody, binds at two Fab each to a cancer antigen (EpCAM) and to a CD3 ϵ (CD3 epsilon) chain expressed on T cells. Catumaxomab induces T cell-mediated cytotoxic activity by binding to the cancer antigen and the CD3 ϵ at the same time, and induces cytotoxic activity mediated by antigen-presenting cells such as NK cells and macrophages, by binding to the cancer antigen and Fc γ R at the same time. By use of these two cytotoxic activities, catumaxomab exhibits a high therapeutic effect on malignant ascites by intraperitoneal administration and has thus been approved in Europe (Non-patent Document 18). In addition, there are cases where the administration of catumaxomab reportedly yields cancer cell-reactive antibodies, which clearly shows that acquired immunity is induced (Non-patent Document 19). From this result, antibodies having both T cell-mediated cytotoxic activity and the Fc γ R-mediated actions by cells such as NK cells or macrophages (these antibodies are particularly referred to as trifunctional antibodies) have received attention because a strong antitumor effect and induction of acquired immunity can be expected.

[0011] The trifunctional antibodies, however, bind to CD3 ϵ and Fc γ R at the same time even in the absence of a cancer antigen and therefore crosslink CD3 ϵ -expressing T cells with Fc γ R-expressing cells even in a cancer cell-absent environment, leading to production of various cytokines in large amounts. Such cancer antigen-independent induction of production of various cytokines restricts the current administration of the trifunctional antibodies to an intraperitoneal route (Non-patent Document 20). The trifunctional antibodies are very difficult to administer systemically due to severe cytokine storm-like adverse reactions. In fact, in the Phase I clinical trial of administering catumaxomab

systemically to non-small-cell lung cancer patients, a very low dose of 5 μ g/body is the maximum permissible dose, and administration of a larger dose has been reported to cause various serious adverse reactions (Non-patent Document 21).

[0012] As such, bispecific antibodies by conventional techniques may bind to both antigens, the first antigen being the cancer antigen (EpCAM) and the second antigen being CD3 ϵ , at the same time when they bind to Fc γ R; and therefore, in view of their molecular structure it is impossible to avoid adverse reactions caused by the simultaneous binding to Fc γ R and the second antigen CD3 ϵ .

[0013] Meanwhile, unlike catumaxomab, BiTE has no Fc γ receptor-binding site, and therefore it does not cross-link the receptors expressed on T cells and cells such as NK cells and macrophages in a cancer antigen-dependent manner. Thus, it has been demonstrated that BiTE does not cause cancer antigen-independent cytokine induction which is observed when catumaxomab is administered. However, since BiTE is a modified low-molecular-weight antibody molecule without an Fc region, the problem is that its blood half-life after administration to a patient is significantly shorter than IgG-type antibodies conventionally used as therapeutic antibodies. In fact, the blood half-life of BiTE administered in vivo has been reported to be about several hours (Non-patent Documents 22 and 23). In the clinical trials of blinatumomab, it is administered by continuous intravenous infusion using a minipump. This administration method is not only extremely inconvenient for patients but also has the potential risk of medical accidents due to device malfunction or the like. Thus, it cannot be said that such an administration method is desirable.

[0014] In recent years, use of an Fc region with reduced Fc γ R-binding activity has enabled maintenance of the strong antitumor activity possessed by BiTE and the excellent safety property of not inducing a cytokine storm in a cancer antigen-dependent manner, and has provided novel polypeptide assemblies that have long half-lives in blood (Patent Document 1).

[0015] On the other hand, when expressing a bispecific antibody by conventional techniques, since two types of H chains and two types of L chains are expressed, ten combinations are conceivable. Among them, only one of the produced combinations has the binding specificity of interest. Therefore, to obtain the bispecific antibody of interest, the single antibody of interest must be purified from the ten types of antibodies, which is very inefficient and difficult.

[0016] A method of preferentially secreting IgGs with a heterodimeric combination of H chains, for example, a combination of an H chain against antigen A and an H chain against antigen B, by introducing amino acid substitutions into the IgG H-chain CH3 region has been reported as a method for solving this problem (Patent Documents 2, 3, 4, 5, 6, 7, and Non-patent Documents 24 and 25). A method that utilizes physical disturbance, i.e., "knob" and "hole", and a method that utilizes electric charge repulsion have been reported as such methods.

[0017] To obtain the molecule of interest with better efficiency, methods using L chains that can bind to two different antigens even though the L chains have the same amino acid sequence have been reported (Patent Documents 8 and 9). However, the antigen affinity may decrease greatly with the use of common L chains, and it is difficult to find common L chains that maintain antigen affinity.

CITATION LIST

- [0018] [Patent Documents]: [Patent Document 1] WO 2012/073985; [Patent Document 2] WO 1996/27011; [Patent Document 3] WO 2006/106905; [Patent Document 4] WO 2007/147901; [Patent Document 5] WO 2009/089004; [Patent Document 6] WO 2010/129304; [Patent Document 7] WO 2013/065708; [Patent Document 8] WO 1998/050431; [Patent Document 9] WO 2006/109592
- [0019] [Non-patent Documents]: [Non-patent Document 1] Nat. Biotechnol. (2005) 23, 1073-1078; [Non-patent Document 2] Eur J Pharm Biopharm. (2005) 59 (3), 389-396; [Non-patent Document 3] Drug Des Devel Ther (2009) 3, 7-16; [Non-patent Document 4] Clin Cancer Res. (2010) 16 (1), 11-20; [Non-patent Document 5] Immunol. Lett. (2002) 82, 57-65; [Non-patent Document 6] Nat. Rev. Immunol. (2008) 8, 34-47; [Non-patent Document 7] Ann. Rev. Immunol. (1988). 6. 251-81; [Non-patent Document 8] J. Bio. Chem., (2000) 276, 16469-16477; [Non-patent Document 9] Nat. Biotech., (2011) 28, 502-10; [Non-patent Document 10] Endocr Relat Cancer (2006) 13, 45-51; [Non-patent Document 11] MAbs. (2012) March 1, 4 (2); [Non-patent Document 12] Nat. Rev. (2010) 10, 301-316; [Non-patent Document 13] Peds (2010), 23 (4), 289-297; [Non-patent Document 14] J. Immunol. (1999) August 1, 163 (3), 1246-52; [Non-patent Document 15] Nature (1985) 314 (6012), 628-31; [Non-patent Document 16] Int J Cancer (1988) 41 (4), 609-15; [Non-patent Document 17] Proc Natl Acad Sci USA (1986) 83 (5), 1453-7; [Non-patent Document 18] Cancer Treat Rev. (2010) October 36 (6), 458-67; [Non-patent Document 19] Future Oncol. (2012) January 8 (1), 73-85; [Non-patent Document 20] Cancer Immunol Immunother. (2007) 56 (9), 1397-406; [Non-patent Document 21] Cancer Immunol Immunother. (2007) 56 (10), 1637-44; [Non-patent Document 22] Cancer Immunol Immunother. (2006) 55 (5), 503-14; [Non-patent Document 23] Cancer Immunol Immunother. (2009) 58 (1), 95-109; [Non-patent Document 24] Protein Engineering. (1996) vol. 9, p. 617-621; [Non-patent Document 25] Nature Biotechnology. (1998) vol. 16, p. 677-681

SUMMARY OF THE INVENTION

Problems to be Solved by the Invention

[0020] The present invention was achieved in view of the above circumstances. An objective of the present invention is to provide multispecific antigen-binding molecules that bring T cells close to the target cancer cells, and can treat cancer through the cytotoxic activity of T cells against target cancer tissues containing glycan 3-expressing cells, and are molecular forms that can be produced with high efficiency; methods for producing the antigen-binding molecules; and pharmaceutical compositions comprising the antigen-binding molecules as active ingredient.

Means for Solving the Problems

[0021] The present inventors discovered an L chain common to a domain comprising a glycan 3-binding antibody variable region, and a domain comprising a T-cell receptor complex-binding antibody variable region, where the common L chain is capable of improving affinity to both anti-

gens. This allows preparation of molecular forms that can be produced with high efficiency, and further discovery of novel multispecific antigen-binding molecules that maintain the strong antitumor activity possessed by T-cell redirecting antibodies such as BiTE and the excellent safety property of not inducing a cytokine storm in a cancer antigen-dependent manner, and also have long half-lives in blood. Furthermore, the present inventors discovered that the multispecific antigen-binding molecules comprising common L chains target glycan 3-expressing cancer cells and cause cell injury. Based on this discovery, the present inventors elucidated that the multispecific antigen-binding molecules of the present invention cause injury to cancer tissues containing glycan 3-expressing cancer cells.

[0022] More specifically, the present invention provides the following:

[0023] [1] A multispecific antigen-binding molecule that comprises:

[0024] (1) a domain comprising an antibody variable region having glycan 3-binding activity,

[0025] (2) a domain comprising an antibody variable region having T-cell receptor complex-binding activity, and

[0026] (3) a domain comprising an Fc region with reduced binding activity towards an Fc γ receptor, wherein the L chain variable regions comprised in the variable region of (1) and the variable region of (2) have a common amino acid sequence; wherein the multispecific antigen-binding molecule has a cytotoxic activity equivalent to or greater than that of the bispecific antibody GPC3_ERY22_rCE115 comprising a glycan 3-binding domain comprising SEQ ID NOs: 47 and 48, and a T-cell receptor complex-binding domain comprising SEQ ID NOs: 49 and 50.

[0027] [2] The multispecific antigen-binding molecule of [1], wherein the cytotoxic activity is T-cell-dependent cytotoxic activity.

[0028] [3] The multispecific antigen-binding molecule of [1] or [2], wherein the T-cell receptor complex-binding activity is binding activity towards a T-cell receptor.

[0029] [4] The multispecific antigen-binding molecule of any one of [1] to [3], wherein the T-cell receptor complex-binding activity is binding activity towards a CD38 chain.

[0030] [5] The multispecific antigen-binding molecule of any one of [1] to [4], wherein the antibody variable region of (1) in [1] is an antibody variable region that comprises any one of the combinations of H-chain CDR1, CDR2, and CDR3 selected from (a1) to (a5) below, or an antibody variable region functionally equivalent thereto:

[0031] (a1) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:40;

[0032] (a2) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:197;

[0033] (a3) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:206;

[0034] (a4) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:211; and

[0035] (a5) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:215.

[0036] [6] The multispecific antigen-binding molecule of any one of [1] to [4], wherein the antibody variable region of (2) in [1] is an antibody variable region that comprises any one of the combinations of H-chain CDR1, CDR2, and CDR3 amino acid sequences selected from (b1) to (b15) below, or an antibody variable region functionally equivalent thereto:

[0037] (b1) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:52;

[0038] (b2) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:103;

[0039] (b3) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 122;

[0040] (b4) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:128;

[0041] (b5) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 129;

[0042] (b6) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:132; (b7) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:142;

[0043] (b8) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 144;

[0044] (b9) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:164;

[0045] (b10) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:168;

[0046] (b11) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:421; (b12) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:424;

[0047] (b13) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:426;

[0048] (b14) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:429; and (b15) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:430.

[0049] [7] The multispecific antigen-binding molecule of any one of [1] to [4], wherein the antibody variable regions of (1) and (2) in [1] are antibody variable regions comprising any one of the combinations of H-chain CDR1, CDR2, and CDR3 selected from the following (c1) to (c19), or antibody variable regions functionally equivalent thereto:

[0050] (c1) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:40; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino

acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:52;

[0051] (c2) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:40; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:421;

[0052] (c3) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:40; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:426;

[0053] (c4) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:40; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:429;

[0054] (c5) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:40; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:430;

[0055] (c6) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 197; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:128;

[0056] (c7) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:206; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 142;

[0057] (c8) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:206; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 144;

[0058] (c9) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:206; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino

acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 164;

[0059] (c10) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:206; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:168;

[0060] (c11) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:211; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 142;

[0061] (c12) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:211; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:144;

[0062] (c13) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:211; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:164;

[0063] (c14) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:211; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:168;

[0064] (c15) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:215; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 103;

[0065] (c16) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:215; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 122;

[0066] (c17) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:215; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino

acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:129;

[0067] (c18) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:215; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 132; and

[0068] (c19) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:215; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:424.

[0069] [8] The multispecific antigen-binding molecule of any one of [5] to [7], wherein CDR1, CDR2, and CDR3 are CDR1, CDR2, and CDR3 regions based on Kabat numbering. [9] The multispecific antigen-binding molecule of any one of [1] to [4], wherein the antibody variable region of (1) in [1] is an antibody variable region comprising any one of the H-chain variable regions selected from (a1) to (a5) below, or an antibody variable region functionally equivalent thereto:

[0070] (a1) an H-chain variable region having the amino acid sequence of SEQ ID NO:40;

[0071] (a2) an H-chain variable region having the amino acid sequence of SEQ ID NO: 197; (a3) an H-chain variable region having the amino acid sequence of SEQ ID NO:206; (a4) an H-chain variable region having the amino acid sequence of SEQ ID NO:211; and

[0072] (a5) an H-chain variable region having the amino acid sequence of SEQ ID NO:215.

[0073] [10] The multispecific antigen-binding molecule of any one of [1] to [4], wherein the antibody variable region of (2) in [1] is an antibody variable region comprising any one of the H-chain variable regions selected from (b1) to (b15) below, or an antibody variable region functionally equivalent thereto:

[0074] (b1) an H-chain variable region having the amino acid sequence of SEQ ID NO:52;

[0075] (b2) an H-chain variable region having the amino acid sequence of SEQ ID NO:103;

[0076] (b3) an H-chain variable region having the amino acid sequence of SEQ ID NO:122;

[0077] (b4) an H-chain variable region having the amino acid sequence of SEQ ID NO:128;

[0078] (b5) an H-chain variable region having the amino acid sequence of SEQ ID NO:129;

[0079] (b6) an H-chain variable region having the amino acid sequence of SEQ ID NO:132;

[0080] (b7) an H-chain variable region having the amino acid sequence of SEQ ID NO:142;

[0081] (b8) an H-chain variable region having the amino acid sequence of SEQ ID NO:144;

[0082] (b9) an H-chain variable region having the amino acid sequence of SEQ ID NO:164;

[0083] (b10) an H-chain variable region having the amino acid sequence of SEQ ID NO:168;

variable region of (2) in [1] which has the amino acid sequence of SEQ ID NO:424.

[0109] [12] The multispecific antigen-binding molecule of any one of [1] to [11], wherein the common L chain of [1] is a common L chain comprising any one of the combinations of CDR1, CDR2, and CDR3 selected from (d1) to (d11) below, or a common L chain functionally equivalent thereto:

[0110] (d1) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:53;

[0111] (d2) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:223;

[0112] (d3) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:299;

[0113] (d4) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:301;

[0114] (d5) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:302;

[0115] (d6) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:304;

[0116] (d7) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:306;

[0117] (d8) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:307;

[0118] (d9) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:309;

[0119] (d10) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:310; and (d11) CDR1, CDR2, and CDR3 identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:319.

[0120] [13] The multispecific antigen-binding molecule of any one of [1] to [11], wherein the L chain variable region of [1] is a variable region of any one of the L chain amino acid sequences selected from (d1) to (d11) below:

[0121] (d1) an L chain comprising the amino acid sequence of SEQ ID NO:53;

[0122] (d2) an L chain comprising the amino acid sequence of SEQ ID NO:223;

[0123] (d3) an L chain comprising the amino acid sequence of SEQ ID NO:299;

[0124] (d4) an L chain comprising the amino acid sequence of SEQ ID NO:301;

[0125] (d5) an L chain comprising the amino acid sequence of SEQ ID NO:302;

[0126] (d6) an L chain comprising the amino acid sequence of SEQ ID NO:304;

[0127] (d7) an L chain comprising the amino acid sequence of SEQ ID NO:306;

[0128] (d8) an L chain comprising the amino acid sequence of SEQ ID NO:307;

[0129] (d9) an L chain comprising the amino acid sequence of SEQ ID NO:309;

[0130] (d10) an L chain comprising the amino acid sequence of SEQ ID NO:310; and

[0131] (d11) an L chain comprising the amino acid sequence of SEQ ID NO:319.

[0132] [14] The multispecific antigen-binding molecule of any one of [1] to [4], wherein the antibody variable regions of (1) and (2) of [1] and the common L chain variable region are antibody variable regions comprising any one of the combinations of H-chain CDR1, CDR2, and CDR3 and L-chain CDR1, CDR2, and CDR3 selected from (e1) to (e25) below, or antibody variable regions functionally equivalent thereto:

[0133] (e1) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 197; CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 128; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of the common L chain and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 53;

[0134] (e2) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 197; CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 128; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of the common L chain and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 299;

[0135] (e3) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 197; CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 128; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of the common L chain and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 310;

[0136] (e4) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 197; CDR1, CDR2, and CDR3 comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 128; and CDR1, CDR2, and CDR3 comprised in the antibody variable region of the common L chain and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 319;

[0137] (e5) CDR1, CDR2, and CDR3 comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequences of the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO:206; CDR1,

[0173] (f15) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:299;

[0174] (f16) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:301;

[0175] (f17) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:302;

[0176] (f18) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:304;

[0177] (f19) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:306;

[0178] (f20) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:307;

[0179] (f21) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 103; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:309;

[0180] (f22) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 122; and an antibody variable region of the common L chain identical to the amino acid sequence of the variable region comprised in SEQ ID NO:53;

[0181] (f23) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 129; and an antibody variable region of the common L chain and identical to the amino acid sequence of the variable region comprised in SEQ ID NO: 53;

[0182] (f24) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO: 132; and an antibody variable region of the common L chain and identical to the amino acid sequence of the variable region comprised in SEQ ID NO: 53;

[0183] (f25) an H-chain variable region comprised in the antibody variable region of (1) in [1] and identical to the amino acid sequence of SEQ ID NO:215; an H-chain variable region comprised in the antibody variable region of (2) in [1] and identical to the amino acid sequence of SEQ ID NO:424; and an antibody variable region of the common L chain and identical to the amino acid sequence of the variable region comprised in SEQ ID NO: 53; and

[0184] (f26) multispecific antigen-binding molecule that binds to an epitope overlapping with each of the epitopes on glycan 3 and T-cell receptor complex bound by the multispecific antigen-binding molecule of any one of (f1) to (f25), and which has a common L chain.

[0185] [16] The multispecific antigen-binding molecule of any one of [1] to [15], wherein the Fc region of (3) in [1] is an Fc region with an amino acid mutation at any of the Fc region-constituting amino acids of SEQ ID NOS: 23 to 26 (IgG1 to IgG4).

[0186] [17] The multispecific antigen-binding molecule of [16], wherein the Fc region of (3) in [1] is an Fc region with mutation of at least one amino acid selected from the following amino acid positions specified by EU numbering:

[0187] position 220, position 226, position 229, position 231, position 232, position 233, position 234, position 235, position 236, position 237, position 238, position 239, position 240, position 264, position 265, position 266, position 267, position 269, position 270, position 295, position 296, position 297, position 298, position 299, position 300, position 325, position 327, position 328, position 329, position 330, position 331, and position 332.

[0188] [18] The multispecific antigen-binding molecule of [16], wherein the Fc region of (3) in [1] is an Fc region comprising at least one amino acid selected from the following amino acids specified by EU numbering: Arg at

amino acid position 234, Ala or Arg at amino acid position 235, Lys at amino acid position 239, and Ala at amino acid position 297.

[0189] [19] The multispecific antigen-binding molecule of any one of to [18], wherein the Fc region of (3) in [1] further comprises an amino acid mutation for promoting formation of a heterodimeric Fc region.

[0190] [20] The multispecific antigen-binding molecule of [19], wherein the heterodimeric Fc region is the amino acid sequence combination of (g1) or (g2) below:

[0191] (g1) a combination of an amino acid sequence identical to the Fc region of a constant region comprising the amino acid sequence of SEQ ID NO:57, and an amino acid sequence identical to the Fc region of a constant region comprising the amino acid sequence of SEQ ID NO:58; and

[0192] (g2) a combination of an amino acid sequence identical to the Fc region of a constant region comprising the amino acid sequence of SEQ ID NO:60 or 62, and an amino acid sequence identical to the Fc region of a constant region comprising the amino acid sequence of SEQ ID NO:61.

[0193] [21] The multispecific antigen-binding molecule of any one of [1] to [20], wherein the multispecific antigen-binding molecule is a bispecific antibody.

[0194] [22] A bispecific antibody of any one of (h1) to (h25) below:

[0195] (h1) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 424 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:53;

[0196] (h2) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:53;

[0197] (h3) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:299;

[0198] (h4) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:301;

[0199] (h5) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:302;

[0200] (h6) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:304;

[0201] (h7) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:306;

[0202] (h8) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 103 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:307;

[0203] (h9) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:215 and a constant region

- [0214] (h20) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:206 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 144 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:223;
- [0215] (h21) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:206 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 142 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:223;
- [0216] (h22) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:206 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 164 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:223;
- [0217] (h23) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:206 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 168 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:223;
- [0218] (h24) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:211 and a constant region having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 164 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:223; and
- [0219] (h25) a bispecific antibody having an antibody H chain having glycan 3-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO:211 and a constant region

having the amino acid sequence of SEQ ID NO:61; an antibody H chain having T-cell receptor complex-binding activity that comprises an antibody H-chain variable region having the amino acid sequence of SEQ ID NO: 168 and a constant region having the amino acid sequence of SEQ ID NO:60 or 62; and a common antibody L chain having the amino acid sequence of SEQ ID NO:223.

[0220] [23] A nucleic acid that encodes the multispecific antigen-binding molecule of any one of [1] to or the bispecific antibody of or [22].

[0221] [24] A vector into which the nucleic acid of is introduced.

[0222] [25] A cell comprising the nucleic acid of or the vector of [24].

[0223] [26] A method for producing the multispecific antigen-binding molecule of any one of [1] to [20] or the bispecific antibody of or by culturing the cell of [25].

[0224] [27] A multispecific antigen-binding molecule or a bispecific antibody produced by the method of [26].

[0225] [28] A pharmaceutical composition comprising the multispecific antigen-binding molecule of any one of [1] to or the bispecific antibody of or [22], and a pharmaceutically acceptable carrier.

[0226] [29] The pharmaceutical composition of [28], which induces cytotoxicity.

[0227] [30] The pharmaceutical composition of [29], wherein the cytotoxicity is T-cell-dependent cytotoxicity.

[0228] [31] The pharmaceutical composition of [28], which is for administration to a patient in need of the multispecific antigen-binding molecule of any one of [1] to or the bispecific antibody of or [22].

[0229] Furthermore, the present invention relates to a kit to be used in a method of the present invention, wherein the kit comprises a multispecific antigen-binding molecule of the present invention, or a multispecific antigen-binding molecule produced by the production method of the present invention. The present invention also relates to use of a multispecific antigen-binding molecule of the present invention, or use of a multispecific antigen-binding molecule produced by the production method of the present invention in the manufacture of a pharmaceutical composition for activating cytotoxic activity. The present invention additionally relates to a multispecific antigen-binding molecule of the present invention, or a multispecific antigen-binding molecule produced by the production method of the present invention to be used in a method of the present invention. Herein, multispecific antigen-binding molecules include bispecific antibodies of the present invention.

[0230] Furthermore, the present invention relates to a multispecific antigen-binding molecule comprising the following domains:

[0231] (1) a domain comprising an antibody variable region having glycan 3-binding activity;

[0232] (2) a domain comprising an antibody variable region having T-cell receptor complex-binding activity;

[0233] wherein the L-chain variable regions contained in the variable regions of (1) and (2) have a commonly shared amino acid sequence. The present invention also relates to the domain of (1), which is more specifically a domain that comprises antibody heavy-chain and/or light-chain variable regions having glycan 3-binding activity, and which is comprised in the multispecific antigen-binding molecule. The present invention addi-

tionally relates to the domain of (2), which is more specifically a domain that comprises an antibody variable region having T-cell receptor complex-binding activity, and which is comprised in the multispecific antigen-binding molecule. Details of the domains of (1) and (2) may include those described in [1] to [22] mentioned above. The multispecific antigen-binding molecule may be a bispecific antibody. Furthermore, the multispecific antigen-binding molecule may further comprise a domain comprising an Fc region, and the Fc region may have a reduced Fcγ receptor-binding activity. Details of the domain comprising an Fc region may include those described in [1] to [22] mentioned above. The present invention relates to a nucleic acid encoding the multispecific antigen-binding molecule or the domains, a vector introduced with the nucleic acid, a cell comprising the nucleic acid or the vector, a method for producing the multispecific antigen-binding molecule by culturing the cells, and a multispecific antigen-binding molecule or domains comprising an antibody variable region having glycan 3-binding activity or T-cell receptor complex-binding activity produced by the method. Furthermore, the present invention relates to a pharmaceutical composition comprising the multispecific antigen-binding molecule and a pharmaceutically acceptable carrier. The pharmaceutical composition may induce cell injury, the cell injury may be T-cell-dependent cellular cytotoxicity, and the composition may be for administration to a patient in need of the multispecific antigen-binding molecule.

[0234] The present invention also provides a multispecific antigen-binding molecule that binds to epitopes overlapping and/or competing with epitopes on each of glycan 3 and T-cell receptor complex bound by the multispecific antigen-binding molecule of any one of (e1) to (e25) of [14] mentioned above, and a multispecific antigen-binding molecule that binds to epitopes overlapping and/or competing with epitopes on each of glycan 3 and T-cell receptor complex bound by the multispecific antigen-binding molecule of any one of (f1) to (f25) of [15].

[0235] Regarding (g1) and (g2) of mentioned above, of the two Fc regions, the former Fc region may be included in the antibody H chain having glycan 3-binding activity and the latter Fc region may be included in the antibody H chain having T-cell receptor complex-binding activity; or the former Fc region may be included in the antibody H chain having T-cell receptor complex-binding activity and the latter Fc region may be included in the antibody H chain having glycan 3-binding activity.

Effects of the Invention

[0236] The present invention provides novel multispecific antigen-binding molecules with molecular forms that can be produced with high efficiency, which maintain the strong antitumor activity possessed by BiTE and the excellent safety property of not causing cancer antigen-independent induction of a cytokine storm and such, and have long half-lives in blood. Pharmaceutical compositions that activate cytotoxic activity, which comprise a multispecific antigen-binding molecule of the present invention as active ingredient, target cancer tissues containing glycan 3-expressing cancer cells to cause cell injury, and can treat or prevent various cancers. The invention enables desirable

treatment which has not only a high level of safety but also reduced physical burden, and is highly convenient for patients.

BRIEF DESCRIPTION OF THE DRAWINGS

[0237] FIGS. 1A and 1B show schematic diagrams of ERY22 (FIG. 1A) and ERY27 (FIG. 1B).

[0238] FIG. 2 is a graph showing the cytotoxic activities of GPC3_ERY22_rCE115 and GPC3_ERY27_hCE115 when NCI-H446 is used as the target cell. The filled diamond (◆) and the filled triangle (▲) indicate the cytotoxic activity of GPC3_ERY22_rCE115 and GPC3_ERY27_hCE115, respectively.

[0239] FIG. 3 is a graph showing the cytotoxic activities of GPC3_ERY22_rCE115 and GPC3_ERY27_hCE115 when PC-10 is used as the target cell. The filled diamond (◆) and the filled triangle (▲) indicate the cytotoxic activity of GPC3_ERY22_rCE115 and GPC3_ERY27_hCE115, respectively.

[0240] FIGS. 4A and 4B is a graph showing the cytotoxic activities of the optimized antibodies when NCI-H446 is used as the target cell.

[0241] FIGS. 5A and 5B is a graph showing the cytotoxic activities of the optimized antibodies when NCI-H446 is used as the target cell.

[0242] FIGS. 6A and 6B is a graph showing the cytotoxic activities of the optimized antibodies when NCI-H446 is used as the target cell.

[0243] FIGS. 7A and 7B is a graph showing the cytotoxic activities of the optimized antibodies when NCI-H446 is used as the target cell.

[0244] FIGS. 8A and 8B is a graph showing the cytotoxic activities of the optimized antibodies when NCI-H446 is used as the target cell.

[0245] FIGS. 9A and 9B is a graph showing the cytotoxic activities of the optimized antibodies when NCI-H446 is used as the target cell.

[0246] FIGS. 10A and 10B show the in vivo antitumor effects of the optimized antibodies when PC-10 is used as the target cell.

[0247] FIGS. 11A and 11B show the in vivo antitumor effects of the optimized antibodies when NCI-H446 is used as the target cell.

[0248] FIG. 12 shows the relationship between the amino acid residues constituting the Fc regions of IgG1, IgG2, IgG3, and IgG4, and the Kabat EU numbering system (herein, also referred to as EU INDEX).

[0249] FIG. 13-1 shows the heavy-chain variable region sequences of amino acid residues 1-79 of CE115HA000 (SEQ ID NO:52), TR01H040 (SEQ ID NO:103), TR01H071 (SEQ ID NO: 132), TR01H082 (SEQ ID NO:142), TR01H084 (SEQ ID NO:144), TR01H109 (SEQ ID NO:164), TR01H113 (SEQ ID NO:168), H0000 (SEQ ID NO:40), H0610 (SEQ ID NO: 215), GCH054 (SEQ ID NO:197), GCH065 (SEQ ID NO:206), and GCH094 (SEQ ID NO:211) according to Kabat et al.

[0250] FIG. 13-2 shows the heavy-chain variable region sequences of amino acid residues 80-113 of CE115HA000 (SEQ ID NO:52), TR01H040 (SEQ ID NO:103), TR01H071 (SEQ ID NO:132), TR01H082 (SEQ ID NO:142), TR01H084 (SEQ ID NO:144), TR01H109 (SEQ ID NO:164), TR01H113 (SEQ ID NO: 168), H0000 (SEQ ID NO:40), H0610 (SEQ ID NO:215), GCH054 (SEQ ID

NO:197), GCH065 (SEQ ID NO:206), and GCH094 (SEQ ID NO:211) according to Kabat et al.

[0251] FIG. 14 shows the light-chain variable region sequences of L0000 (SEQ ID NO: 53), L0011 (SEQ ID NO:223), L0201 (SEQ ID NO:299), L0203 (SEQ ID NO:301), L0204 (SEQ ID NO:302), L0206 (SEQ ID NO:304), L0208 (SEQ ID NO:306), L0209 (SEQ ID NO:307), L0211 (SEQ ID NO:309), L0212 (SEQ ID NO:310), L0222 (SEQ ID NO: 319), and L0272 (SEQ ID NO:359), according to Kabat et al.

MODE FOR CARRYING OUT THE INVENTION

[0252] The definitions below are provided to help understanding of the present invention illustrated herein.

Antibody

[0253] Herein, "antibody" refers to a natural immunoglobulin or an immunoglobulin produced by partial or complete synthesis. Antibodies can be isolated from natural sources such as naturally-occurring plasma and serum, or culture supernatants of antibody-producing hybridomas. Alternatively, antibodies can be partially or completely synthesized using techniques such as genetic recombination. Preferred antibodies include, for example, antibodies of an immunoglobulin isotype or subclass belonging thereto. Known human immunoglobulins include antibodies of the following nine classes (isotypes): IgG1, IgG2, IgG3, IgG4, IgA1, IgA2, IgD, IgE, and IgM. Of these isotypes, antibodies of the present invention include IgG1, IgG2, IgG3, and IgG4.

[0254] Methods for producing an antibody with desired binding activity are known to those skilled in the art. Below is an example that describes a method for producing an antibody (anti-GPC3 antibody) that binds to Glycan-3 (hereinafter, also referred to as GPC3), which belongs to the GPI-anchored receptor family (Int J Cancer. (2003) 103 (4), 455-65). Antibodies that bind to a T-cell receptor complex can also be produced according to the example described below.

[0255] Anti-GPC3 antibodies can be obtained as polyclonal or monoclonal antibodies using known methods. The anti-GPC3 antibodies preferably produced are monoclonal antibodies derived from mammals. Such mammal-derived monoclonal antibodies include antibodies produced by hybridomas or host cells transformed with an expression vector carrying an antibody gene by genetic engineering techniques.

[0256] Monoclonal antibody-producing hybridomas can be produced using known techniques, for example, as described below. Specifically, mammals are immunized by conventional immunization methods using a GPC3 protein as a sensitizing antigen. Resulting immune cells are fused with known parental cells by conventional cell fusion methods. Then, hybridomas producing an anti-GPC3 antibody can be selected by screening for monoclonal antibody-producing cells using conventional screening methods.

[0257] Specifically, monoclonal antibodies are prepared as mentioned below. First, the GPC3 gene whose nucleotide sequence is disclosed in RefSeq accession number NM_001164617.1 (SEQ ID NO:1) can be expressed to produce a GPC3 protein shown in RefSeq accession number NP_001158089.1 (SEQ ID NO:2), which will be used as a sensitizing antigen for antibody preparation. That is, a gene

sequence encoding GPC3 is inserted into a known expression vector, and appropriate host cells are transformed with this vector. The desired human GPC3 protein is purified from the host cells or their culture supernatants by known methods. For example, to prepare soluble GPC3 from culture supernatants, amino acids at positions 564 to 580 that form the hydrophobic region corresponding to the GPI-anchor sequence used to anchor GPC3 on the cell membrane are deleted from the GPC3 polypeptide sequence of SEQ ID NO:2, and then the resulting protein is expressed instead of the GPC3 protein of SEQ ID NO:2. Alternatively, it is possible to use a purified natural GPC3 protein as a sensitizing antigen.

[0258] The purified GPC3 protein can be used as a sensitizing antigen for use in immunization of mammals. Partial peptides of GPC3 can also be used as sensitizing antigens. In this case, the partial peptides may also be obtained by chemical synthesis from the human GPC3 amino acid sequence. Furthermore, they may also be obtained by incorporating a portion of the GPC3 gene into an expression vector and expressing it. Moreover, they may also be obtained by degrading the GPC3 protein using proteases, but the region and size of the GPC3 peptide used as the partial peptide are not particularly limited to a special embodiment. As the preferred region, any sequence from the amino acid sequence corresponding to the amino acids at positions 524 to 563, or more preferably any sequence from the amino acid sequence corresponding to the amino acids at positions 537 to 563 in the amino acid sequence of SEQ ID NO:2 may be selected. Preferably, any sequence may be selected from the amino acid sequence of the region not containing the amino acid sequence corresponding to amino acids at positions 550 to 663 in the amino acid sequence of SEQ ID NO: 2. Preferably, any sequence may be selected from the amino acid sequence corresponding to positions 544 to 553, and more preferably, any sequence may be selected from the amino acid sequence corresponding to positions 546 to 551 in the amino acid sequence of SEQ ID NO:2. The number of amino acids constituting a peptide to be used as the sensitizing antigen is at least five or more, or preferably for example, six or more, or seven or more. More specifically, peptides consisting of 8 to 50 residues or preferably 10 to 30 residues may be used as the sensitizing antigen.

[0259] For sensitizing antigen, alternatively it is possible to use a fusion protein prepared by fusing a desired partial polypeptide or peptide of the GPC3 protein with a different polypeptide. For example, antibody Fc fragments and peptide tags are preferably used to produce fusion proteins to be used as sensitizing antigens. Vectors for expression of such fusion proteins can be constructed by fusing in frame genes encoding two or more desired polypeptide fragments and inserting the fusion gene into an expression vector as described above. Methods for producing fusion proteins are described in Molecular Cloning 2nd ed. (Sambrook, J et al., Molecular Cloning 2nd ed., 9.47-9.58 (1989) Cold Spring Harbor Lab. Press). Methods for preparing GPC3 to be used as a sensitizing antigen, and immunization methods using GPC3 are specifically described in WO 2003/000883, WO 2004/022754, and WO 2006/006693.

[0260] There is no particular limitation on the mammals to be immunized with the sensitizing antigen. However, it is preferable to select the mammals by considering their compatibility with the parent cells to be used for cell fusion. In

general, rodents such as mice, rats, and hamsters, rabbits, and monkeys are preferably used.

[0261] The above animals are immunized with a sensitizing antigen by known methods. Generally performed immunization methods include, for example, intraperitoneal or subcutaneous injection of a sensitizing antigen into mammals. Specifically, a sensitizing antigen is appropriately diluted with PBS (Phosphate-Buffered Saline), physiological saline, or the like. If desired, a conventional adjuvant such as Freund's complete adjuvant is mixed with the antigen, and the mixture is emulsified. Then, the sensitizing antigen is administered to a mammal several times at 4- to 21-day intervals. Appropriate carriers may be used in immunization with the sensitizing antigen. In particular, when a low-molecular-weight partial peptide is used as the sensitizing antigen, it is sometimes desirable to couple the sensitizing antigen peptide to a carrier protein such as albumin or keyhole limpet hemocyanin for immunization.

[0262] Alternatively, hybridomas producing a desired antibody can be prepared using DNA immunization as mentioned below. DNA immunization is an immunization method that confers immunostimulation by expressing a sensitizing antigen in an animal immunized as a result of administering a vector DNA constructed to allow expression of an antigen protein-encoding gene in the animal. As compared to conventional immunization methods in which a protein antigen is administered to animals to be immunized, DNA immunization is expected to be superior in that:

[0263] immunostimulation can be provided while retaining the structure of a membrane protein such as GPC3; and

[0264] there is no need to purify the antigen for immunization.

[0265] In order to prepare a monoclonal antibody of the present invention using DNA immunization, first, a DNA expressing a GPC3 protein is administered to an animal to be immunized. The GPC3-encoding DNA can be synthesized by known methods such as PCR. The obtained DNA is inserted into an appropriate expression vector, and then this is administered to an animal to be immunized. Preferably used expression vectors include, for example, commercially-available expression vectors such as pcDNA3.1. Vectors can be administered to an organism using conventional methods. For example, DNA immunization is performed by using a gene gun to introduce expression vector-coated gold particles into cells in the body of an animal to be immunized. Antibodies that recognized GPC3 can also be produced by the methods described in WO 2003/104453.

[0266] After immunizing a mammal as described above, an increase in the titer of a GPC3-binding antibody is confirmed in the serum. Then, immune cells are collected from the mammal, and then subjected to cell fusion. In particular, splenocytes are preferably used as immune cells.

[0267] A mammalian myeloma cell is used as a cell to be fused with the above-mentioned immunocyte. The myeloma cells preferably comprise a suitable selection marker for screening. A selection marker confers characteristics to cells for their survival (or death) under a specific culture condition. Hypoxanthine-guanine phosphoribosyltransferase deficiency (hereinafter abbreviated as HGPRT deficiency) and thymidine kinase deficiency (hereinafter abbreviated as TK deficiency) are known as selection markers. Cells with HGPRT or TK deficiency have hypoxanthine-aminopterin-thymidine sensitivity (hereinafter abbreviated as HAT sen-

sitivity). HAT-sensitive cells cannot synthesize DNA in a HAT selection medium, and are thus killed. However, when the cells are fused with normal cells, they can continue DNA synthesis using the salvage pathway of the normal cells, and therefore they can grow even in the HAT selection medium. [0268] HGPRT-deficient and TK-deficient cells can be selected in a medium containing 6-thioguanine, 8-azaguanine (hereinafter abbreviated as 8AG), or 5'-bromodeoxyuridine, respectively. Normal cells are killed because they incorporate these pyrimidine analogs into their DNA. Meanwhile, cells that are deficient in these enzymes can survive in the selection medium, since they cannot incorporate these pyrimidine analogs. In addition, a selection marker referred to as G418 resistance provided by the neomycin-resistant gene confers resistance to 2-deoxystreptamine antibiotics (gentamycin analogs). Various types of myeloma cells that are suitable for cell fusion are known.

[0269] For example, myeloma cells including the following cells can be preferably used: P3 (P3x63Ag8.653) (J. Immunol. (1979) 123 (4), 1548-1550); P3x63Ag8U.1 (Current Topics in Microbiology and Immunology (1978) 81, 1-7); NS-1 (C. Eur. J. Immunol. (1976) 6 (7), 511-519); MPC-11 (Cell (1976) 8 (3), 405-415); SP2/0 (Nature (1978) 276 (5685), 269-270); FO (J. Immunol. Methods (1980) 35 (1-2), 1-21); S194/5.XXO.BU.1 (J. Exp. Med. (1978) 148 (1), 313-323); R210 (Nature (1979) 277 (5692), 131-133), etc.

[0270] Cell fusions between the immunocytes and myeloma cells are essentially carried out using known methods, for example, a method by Kohler and Milstein et al. (Methods Enzymol. (1981) 73:3-46).

[0271] More specifically, cell fusion can be carried out, for example, in a conventional culture medium in the presence of a cell fusion-promoting agent. The fusion-promoting agents include, for example, polyethylene glycol (PEG) and Sendai virus (HVJ). If required, an auxiliary substance such as dimethyl sulfoxide is also added to improve fusion efficiency.

[0272] The ratio of immunocytes to myeloma cells may be determined at one's own discretion, preferably, for example, one myeloma cell for every one to ten immunocytes. Culture media to be used for cell fusions include, for example, media that are suitable for the growth of myeloma cell lines, such as RPMI1640 medium and MEM medium, and other conventional culture medium used for this type of cell culture. In addition, serum supplements such as fetal calf serum (FCS) may be preferably added to the culture medium.

[0273] For cell fusion, predetermined amounts of the above immune cells and myeloma cells are mixed well in the above culture medium. Then, a PEG solution (for example, the average molecular weight is about 1,000 to 6,000) prewarmed to about 37° C. is added thereto at a concentration of generally 30% to 60% (w/v). This is gently mixed to produce desired fusion cells (hybridomas). Then, an appropriate culture medium mentioned above is gradually added to the cells, and this is repeatedly centrifuged to remove the supernatant. Thus, cell fusion agents and such which are unfavorable to hybridoma growth can be removed.

[0274] The hybridomas thus obtained can be selected by culture using a conventional selective medium, for example, HAT medium (a culture medium containing hypoxanthine, aminopterin, and thymidine). Cells other than the desired hybridomas (non-fused cells) can be killed by continuing culture in the above HAT medium for a sufficient period of

time. Typically, the period is several days to several weeks. Then, hybridomas producing the desired antibody are screened and singly cloned by conventional limiting dilution methods.

[0275] The hybridomas thus obtained can be selected using a selection medium based on the selection marker possessed by the myeloma used for cell fusion. For example, HGPRT- or TK-deficient cells can be selected by culture using the HAT medium (a culture medium containing hypoxanthine, aminopterin, and thymidine). Specifically, when HAT-sensitive myeloma cells are used for cell fusion, cells successfully fused with normal cells can selectively proliferate in the HAT medium. Cells other than the desired hybridomas (non-fused cells) can be killed by continuing culture in the above HAT medium for a sufficient period of time. Specifically, desired hybridomas can be selected by culture for generally several days to several weeks. Then, hybridomas producing the desired antibody are screened and singly cloned by conventional limiting dilution methods.

[0276] Desired antibodies can be preferably selected and singly cloned by screening methods based on known antigen/antibody reaction. For example, a GPC3-binding monoclonal antibody can bind to GPC3 expressed on the cell surface. Such a monoclonal antibody can be screened by fluorescence activated cell sorting (FACS). FACS is a system that assesses the binding of an antibody to cell surface by analyzing cells contacted with a fluorescent antibody using laser beam, and measuring the fluorescence emitted from individual cells.

[0277] To screen for hybridomas that produce a monoclonal antibody of the present invention by FACS, GPC3-expressing cells are first prepared. Cells preferably used for screening are mammalian cells in which GPC3 is forcedly expressed. As control, the activity of an antibody to bind to cell-surface GPC3 can be selectively detected using non-transformed mammalian cells as host cells. Specifically, hybridomas producing an anti-GPC3 monoclonal antibody can be isolated by selecting hybridomas that produce an antibody which binds to cells forced to express GPC3, but not to host cells.

[0278] Alternatively, the activity of an antibody to bind to immobilized GPC3-expressing cells can be assessed based on the principle of ELISA. For example, GPC3-expressing cells are immobilized to the wells of an ELISA plate. Culture supernatants of hybridomas are contacted with the immobilized cells in the wells, and antibodies that bind to the immobilized cells are detected. When the monoclonal antibodies are derived from mouse, antibodies bound to the cells can be detected using an anti-mouse immunoglobulin antibody. Hybridomas producing a desired antibody having the antigen-binding ability are selected by the above screening, and they can be cloned by a limiting dilution method or the like.

[0279] Monoclonal antibody-producing hybridomas thus prepared can be passaged in a conventional culture medium, and stored in liquid nitrogen for a long period.

[0280] The above hybridomas are cultured by a conventional method, and desired monoclonal antibodies can be prepared from the culture supernatants. Alternatively, the hybridomas are administered to and grown in compatible mammals, and monoclonal antibodies are prepared from the ascites. The former method is suitable for preparing antibodies with high purity.

[0281] Antibodies encoded by antibody genes that are cloned from antibody-producing cells such as the above hybridomas can also be preferably used. A cloned antibody gene is inserted into an appropriate vector, and this is introduced into a host to express the antibody encoded by the gene. Methods for isolating antibody genes, inserting the genes into vectors, and transforming host cells have already been established, for example, by Vandamme et al. (*Eur. J. Biochem.* (1990) 192 (3), 767-775). Methods for producing recombinant antibodies are also known as described below.

[0282] For example, a cDNA encoding the variable region (V region) of an anti-GPC3 antibody is prepared from hybridoma cells expressing the anti-GPC3 antibody. For this purpose, total RNA is first extracted from hybridomas. Methods used for extracting mRNAs from cells include, for example:

[0283] the guanidine ultracentrifugation method (*Biochemistry* (1979) 18 (24), 5294-5299), and

[0284] the AGPC method (*Anal. Biochem.* (1987) 162 (1), 156-159)

[0285] Extracted mRNAs can be purified using the mRNA Purification Kit (GE Healthcare Bioscience) or such. Alternatively, kits for extracting total mRNA directly from cells, such as the QuickPrep mRNA Purification Kit (GE Healthcare Bioscience), are also commercially available. mRNAs can be prepared from hybridomas using such kits. cDNAs encoding the antibody V region can be synthesized from the prepared mRNAs using a reverse transcriptase. cDNAs can be synthesized using the AMV Reverse Transcriptase First-strand cDNA Synthesis Kit (Seikagaku Co.) or such. Furthermore, the SMART RACE cDNA amplification kit (Clontech) and the PCR-based 5'-RACE method (*Proc. Natl. Acad. Sci. USA* (1988) 85 (23), 8998-9002; *Nucleic Acids Res.* (1989) 17 (8), 2919-2932) can be appropriately used to synthesize and amplify cDNAs. In such a cDNA synthesis process, appropriate restriction enzyme sites described below may be introduced into both ends of a cDNA.

[0286] The cDNA fragment of interest is purified from the resulting PCR product, and then this is ligated to a vector DNA. A recombinant vector is thus constructed, and introduced into *E. coli* or such. After colony selection, the desired recombinant vector can be prepared from the colony-forming *E. coli*. Then, whether the recombinant vector has the cDNA nucleotide sequence of interest is tested by a known method such as the dideoxy nucleotide chain termination method.

[0287] The 5'-RACE method which uses primers to amplify the variable region gene is conveniently used for isolating the gene encoding the variable region. First, a 5'-RACE cDNA library is constructed by cDNA synthesis using RNAs extracted from hybridoma cells as a template. A commercially available kit such as the SMART RACE cDNA amplification kit is appropriately used to synthesize the 5'-RACE cDNA library.

[0288] The antibody gene is amplified by PCR using the prepared 5'-RACE cDNA library as a template. Primers for amplifying the mouse antibody gene can be designed based on known antibody gene sequences. The nucleotide sequences of the primers vary depending on the immunoglobulin subclass. Therefore, it is preferable that the subclass is determined in advance using a commercially available kit such as the Iso Strip mouse monoclonal antibody isotyping kit (Roche Diagnostics).

[0289] Specifically, for example, primers that allow amplification of genes encoding $\gamma 1$, $\gamma 2a$, $\gamma 2b$, and $\gamma 3$ heavy chains and k and A light chains are used to isolate mouse IgG-encoding genes. In general, a primer that anneals to a constant region site close to the variable region is used as a 3'-side primer to amplify an IgG variable region gene. Meanwhile, a primer attached to a 5' RACE cDNA library construction kit is used as a 5'-side primer.

[0290] PCR products thus amplified are used to reshape immunoglobulins composed of a combination of heavy and light chains. A desired antibody can be selected using the GPC3-binding activity of a reshaped immunoglobulin as an indicator. For example, when the objective is to isolate an antibody against GPC3, it is more preferred that the binding of the antibody to GPC3 is specific. A GPC3-binding antibody can be screened, for example, by the following steps:

[0291] (1) contacting a GPC3-expressing cell with an antibody comprising the V region encoded by a cDNA isolated from a hybridoma;

[0292] (2) detecting the binding of the antibody to the GPC3-expressing cell; and

[0293] (3) selecting an antibody that binds to the GPC3-expressing cell.

[0294] Methods for detecting the binding of an antibody to GPC3-expressing cells are known. Specifically, the binding of an antibody to GPC3-expressing cells can be detected by the above-described techniques such as FACS. Immobilized samples of GPC3-expressing cells are appropriately used to assess the binding activity of an antibody.

[0295] Preferred antibody screening methods that use the binding activity as an indicator also include panning methods using phage vectors. Screening methods using phage vectors are advantageous when the antibody genes are isolated from heavy-chain and light-chain subclass libraries from a polyclonal antibody-expressing cell population. Genes encoding the heavy-chain and light-chain variable regions can be linked by an appropriate linker sequence to form a single-chain Fv (scFv). Phages presenting scFv on their surface can be produced by inserting a gene encoding scFv into a phage vector. The phages are contacted with an antigen of interest. Then, a DNA encoding scFv having the binding activity of interest can be isolated by collecting phages bound to the antigen. This process can be repeated as necessary to enrich scFv having a desired binding activity.

[0296] After isolation of the cDNA encoding the V region of the anti-GPC3 antibody of interest, the cDNA is digested with restriction enzymes that recognize the restriction sites introduced into both ends of the cDNA. Preferred restriction enzymes recognize and cleave a nucleotide sequence that occurs in the nucleotide sequence of the antibody gene at a low frequency. Furthermore, a restriction site for an enzyme that produces a sticky end is preferably introduced into a vector to insert a single-copy digested fragment in the correct orientation. The cDNA encoding the V region of the anti-GPC3 antibody is digested as described above, and this is inserted into an appropriate expression vector to construct an antibody expression vector. In this case, if a gene encoding the antibody constant region (C region) and a gene encoding the above V region are fused in-frame, a chimeric antibody is obtained. Herein, "chimeric antibody" means that the origin of the constant region is different from that of the variable region. Thus, in addition to mouse/human heterochimeric antibodies, human/human allochimeric antibodies are included in the chimeric antibodies of the present

invention. A chimeric antibody expression vector can be constructed by inserting the above V region gene into an expression vector that already has the constant region. Specifically, for example, a recognition sequence for a restriction enzyme that excises the above V region gene can be appropriately placed on the 5' side of an expression vector carrying a DNA encoding a desired antibody constant region (C region). A chimeric antibody expression vector is constructed by fusing in frame the two genes digested with the same combination of restriction enzymes.

[0297] To produce an anti-GPC3 monoclonal antibody, antibody genes are inserted into an expression vector so that the genes are expressed under the control of an expression regulatory region. The expression regulatory region for antibody expression includes, for example, enhancers and promoters. Furthermore, an appropriate signal sequence may be attached to the amino terminus so that the expressed antibody is secreted to the outside of cells. In the Examples described below, a peptide having the amino acid sequence MGWSCLILFLVATATGVHS (SEQ ID NO:3) is used as a signal sequence. Meanwhile, other appropriate signal sequences may be attached. The expressed polypeptide is cleaved at the carboxyl terminus of the above sequence, and the resulting polypeptide is secreted to the outside of cells as a mature polypeptide. Then, appropriate host cells are transformed with the expression vector, and recombinant cells expressing the anti-GPC3 antibody-encoding DNA are obtained.

[0298] DNAs encoding the antibody heavy chain (H chain) and light chain (L chain) are separately inserted into different expression vectors to express the antibody gene. An antibody molecule having the H and L chains can be expressed by co-transfected the same host cell with vectors into which the H-chain and L-chain genes are respectively inserted. Alternatively, host cells can be transformed with a single expression vector into which DNAs encoding the H and L chains are inserted (see WO 94/11523).

[0299] There are various known host cell/expression vector combinations for antibody preparation by introducing isolated antibody genes into appropriate hosts. All of these expression systems are applicable to isolation of domains including antibody variable regions of the present invention. Appropriate eukaryotic cells used as host cells include animal cells, plant cells, and fungal cells. Specifically, the animal cells include, for example, the following cells: (1) mammalian cells: CHO, COS, myeloma, baby hamster kidney (BHK), HeLa, Vero, or such; (2) amphibian cells: *Xenopus* oocytes, or such; and

[0300] (3) insect cells: sf9, sf21, Tn5, or such.

[0301] In addition, as a plant cell, an antibody gene expression system using cells derived from the *Nicotiana* genus such as *Nicotiana tabacum* is known. Callus cultured cells can be appropriately used to transform plant cells.

[0302] Furthermore, the following cells can be used as fungal cells: yeasts: the *Saccharomyces* genus such as *Saccharomyces cerevisiae*, and the *Pichia* genus such as *Pichia pastoris*; and filamentous fungi: the *Aspergillus* genus such as *Aspergillus niger*.

[0303] Furthermore, antibody gene expression systems that utilize prokaryotic cells are also known. For example, when using bacterial cells, *E. coli* cells, *Bacillus subtilis* cells, and such can suitably be utilized in the present invention. Expression vectors carrying the antibody genes of interest are introduced into these cells by transfection. The

transfected cells are cultured in vitro, and the desired antibody can be prepared from the culture of transformed cells.

[0304] In addition to the above-described host cells, transgenic animals can also be used to produce a recombinant antibody. That is, the antibody can be obtained from an animal into which the gene encoding the antibody of interest is introduced. For example, the antibody gene can be constructed as a fusion gene by inserting in frame into a gene that encodes a protein produced specifically in milk. Goat β-casein or such can be used, for example, as the protein secreted in milk. DNA fragments containing the fused gene inserted with the antibody gene is injected into a goat embryo, and then this embryo is introduced into a female goat. Desired antibodies can be obtained as a protein fused with the milk protein from milk produced by the transgenic goat born from the embryo-recipient goat (or progeny thereof). In addition, to increase the volume of milk containing the desired antibody produced by the transgenic goat, hormones can be administered to the transgenic goat as necessary (Ebert, K. M. et al., Bio/Technology (1994) 12 (7), 699-702).

[0305] When an antigen-binding molecule described herein is administered to human, a domain derived from a genetically recombinant antibody that has been artificially modified to reduce the heterologous antigenicity against human and such, can be appropriately used as the domain of the antigen-binding molecule including an antibody variable region. Such genetically recombinant antibodies include, for example, humanized antibodies. These modified antibodies are appropriately produced by known methods.

[0306] An antibody variable region used to produce a domain of an antigen-binding molecule including an antibody variable region described herein is generally formed by three complementarity-determining regions (CDRs) that are separated by four framework regions (FRs). CDR is a region that substantially determines the binding specificity of an antibody. The amino acid sequences of CDRs are highly diverse. On the other hand, the FR-forming amino acid sequences often have high identity even among antibodies with different binding specificities. Therefore, generally, the binding specificity of a certain antibody can be introduced to another antibody by CDR grafting.

[0307] A humanized antibody is also called a reshaped human antibody. Specifically, humanized antibodies prepared by grafting the CDR of a non-human animal antibody such as a mouse antibody to a human antibody and such are known. Common genetic engineering techniques for obtaining humanized antibodies are also known. Specifically, for example, overlap extension PCR is known as a method for grafting a mouse antibody CDR to a human FR. In overlap extension PCR, a nucleotide sequence encoding a mouse antibody CDR to be grafted is added to primers for synthesizing a human antibody FR. Primers are prepared for each of the four FRs. It is generally considered that when grafting a mouse CDR to a human FR, selecting a human FR that has high identity to a mouse FR is advantageous for maintaining the CDR function. That is, it is generally preferable to use a human FR comprising an amino acid sequence which has high identity to the amino acid sequence of the FR adjacent to the mouse CDR to be grafted.

[0308] Nucleotide sequences to be ligated are designed so that they will be connected to each other in frame. Human FRs are individually synthesized using the respective primers. As a result, products in which the mouse CDR-encoding

DNA is attached to the individual FR-encoding DNAs are obtained. Nucleotide sequences encoding the mouse CDR of each product are designed so that they overlap with each other. Then, complementary strand synthesis reaction is conducted to anneal the overlapping CDR regions of the products synthesized using a human antibody gene as template. Human FRs are ligated via the mouse CDR sequences by this reaction.

[0309] The full length V region gene, in which three CDRs and four FRs are ultimately ligated, is amplified using primers that anneal to its 5'- or 3'-end, which are added with suitable restriction enzyme recognition sequences. An expression vector for humanized antibody can be produced by inserting the DNA obtained as described above and a DNA that encodes a human antibody C region into an expression vector so that they will ligate in frame. After the recombinant vector is transfected into a host to establish recombinant cells, the recombinant cells are cultured, and the DNA encoding the humanized antibody is expressed to produce the humanized antibody in the cell culture (see, European Patent Publication No. EP 239400 and International Patent Publication No. WO 1996/002576).

[0310] By qualitatively or quantitatively measuring and evaluating the antigen-binding activity of the humanized antibody produced as described above, one can suitably select human antibody FRs that allow CDRs to form a favorable antigen-binding site when ligated through the CDRs. Amino acid residues in FRs may be substituted as necessary, so that the CDRs of a reshaped human antibody form an appropriate antigen-binding site. For example, amino acid sequence mutations can be introduced into FRs by applying the PCR method used for grafting a mouse CDR into a human FR. More specifically, partial nucleotide sequence mutations can be introduced into primers that anneal to the FR. Nucleotide sequence mutations are introduced into the FRs synthesized by using such primers. Mutant FR sequences having the desired characteristics can be selected by measuring and evaluating the activity of the amino acid-substituted mutant antibody to bind to the antigen by the above-mentioned method (Sato, K. et al., Cancer Res. (1993) 53:851-856).

[0311] Alternatively, desired human antibodies can be obtained by immunizing transgenic animals having the entire repertoire of human antibody genes (see WO 1993/012227; WO 1992/003918; WO 1994/002602; WO 1994/025585; WO 1996/034096; WO 1996/033735) by DNA immunization.

[0312] Furthermore, techniques for preparing human antibodies by panning using human antibody libraries are also known. For example, the V region of a human antibody is expressed as a single-chain antibody (scFv) on phage surface by the phage display method. Phages expressing an scFv that binds to the antigen can be selected. The DNA sequence encoding the human antibody V region that binds to the antigen can be determined by analyzing the genes of selected phages. The DNA sequence of the scFv that binds to the antigen is determined. An expression vector is prepared by fusing the V region sequence in frame with the C region sequence of a desired human antibody, and inserting this into an appropriate expression vector. The expression vector is introduced into cells appropriate for expression such as those described above. The human antibody can be produced by expressing the human antibody-encoding gene in the cells. These methods are already known (see WO

1992/001047; WO 1992/020791; WO 1993/006213; WO 1993/011236; WO 1993/019172; WO 1995/001438; WO 1995/015388).

A domain comprising an antibody variable region having glycan 3 (GPC3)-binding activity

[0313] Herein, the phrase “a domain comprising an antibody variable region having glycan 3 (GPC3)-binding activity” refers to an antibody portion that comprises a region that specifically binds to the above-mentioned GPC3 protein, or to all or a portion of a partial peptide of the GPC3 protein, and is also complementary thereto. Domains comprising an antibody variable region may be provided from variable domains of one or a plurality of antibodies. Preferably, domains comprising an antibody variable region comprise antibody light-chain and heavy-chain variable regions (VL and VH). Suitable examples of such domains comprising antibody variable regions include “single chain Fv (scFv)”, “single chain antibody”, “Fv”, “single chain Fv 2 (scFv2)”, “Fab”, “F(ab')2”, etc.

A domain comprising an antibody variable region having T-cell receptor complex-binding activity

[0314] Herein, the phrase “a domain comprising an antibody variable region having T-cell receptor complex-binding activity” refers to a T-cell receptor complex-binding antibody portion that comprises a region that specifically binds to all or a portion of a T-cell receptor complex and is also complementary thereto. The T-cell receptor complex may be a T-cell receptor itself, or an adaptor molecule constituting a T-cell receptor complex along with a T-cell receptor. CD3 is suitable as an adaptor molecule.

A Domain Comprising an Antibody Variable Region that has T-Cell Receptor-Binding Activity

[0315] Herein, the phrase “a domain comprising an antibody variable region having T-cell receptor-binding activity” refers to a T-cell receptor-binding antibody portion produced by including a region that specifically binds to all or a portion of a T-cell receptor and is also complementary thereto.

[0316] The portion of a T cell receptor to which the domain of the present invention binds may be a variable region or a constant region, but an epitope present in the constant region is preferred. Examples of the constant region sequence include the T cell receptor α chain of RefSeq Accession No. CAA26636.1 (SEQ ID NO:4), the T cell receptor β chain of RefSeq Accession No. C25777 (SEQ ID NO:5), the T cell receptor $\gamma 1$ chain of RefSeq Accession No. A26659 (SEQ ID NO:6), the T cell receptor $\gamma 2$ chain of RefSeq Accession No. AAB63312.1 (SEQ ID NO:7), and the T cell receptor δ chain of RefSeq Accession No. AAA61033.1 (SEQ ID NO:8).

A Domain Comprising an Antibody Variable Region that has CD3-Binding Activity

[0317] Herein, the phrase “a domain comprising an antibody variable region that has CD3-binding activity” refers to a CD3-binding antibody portion produced by including a region that specifically binds to all or a portion of CD3 and is also complementary thereto. Preferably, the domain comprises the light-chain and heavy-chain variable regions (VL and VH) of an anti-CD3 antibody. Suitable examples of such a domain include “single chain Fv (scFv)”, “single chain antibody”, “Fv”, “single chain Fv 2 (scFv2)”, “Fab”, “F(ab')2”, etc.

[0318] The domain comprising an antibody variable region that has CD3-binding activity of the present invention

may be any epitope-binding domain as long as the epitope exists in the γ -chain, δ -chain, or ϵ -chain sequence that constitutes human CD3. In the present invention, preferably, a domain comprising an anti-CD3 antibody light-chain variable region (VL) and an anti-CD3 antibody heavy-chain variable region (VH) that bind to an epitope present in the extracellular region of the γ chain of the human CD3 complex is suitably used. Besides the anti-CD3 antibody light chain variable region (VL) and anti-CD3 antibody heavy chain variable region (VH) described in the Examples, various known CD3-binding domains containing a CD3-binding antibody light chain variable region (VL) and a CD3-binding antibody heavy chain variable region (VH), and those of the OKT3 antibody (Proc. Natl. Acad. Sci. USA (1980) 77, 4914-4917) are suitably used as such domains. One may appropriately use an antibody variable region-containing domain derived from the anti-CD3 antibody having desired properties, which is obtained by immunizing a desired animal by the above-mentioned method using the γ -chain, δ -chain, or ϵ -chain constituting the human CD3. Human antibodies and properly humanized antibodies as described above may be appropriately used as the anti-CD3 antibody to give rise to the domain containing the antibody variable region having CD3-binding activity. Regarding the structure of the γ -chain, δ -chain, or ϵ -chain constituting CD3, their polynucleotide sequences are shown in SEQ ID NOS: 9 (NM_000073.2), 10 (NM_000732.4), and 11 (NM_000733.3), and their polypeptide sequences are shown in SEQ ID NOS: 12 (NP_000064.1), 13 (NP_000723.1), and 14 (NP_000724.1) (the RefSeq accession number is shown in parentheses).

[0319] Antibody variable region-containing domains in antigen binding molecules of the present invention may bind to the same epitope. Herein, the same epitope may be present in a protein comprising the amino acid sequence of SEQ ID NO:2 or 14. Alternatively, antibody variable region-containing domains in antigen binding molecules of the present invention may bind to different epitopes, respectively. Herein, the different epitopes may be present in a protein comprising the amino acid sequence of SEQ ID NO:2 or 14.

Specific

[0320] The term “specific” means that one of molecules involved in specific binding does not show any significant binding to molecules other than a single or a number of binding partner molecules. Furthermore, the term is also used when a domain containing an antibody variable region is specific to a particular epitope among multiple epitopes in an antigen. When an epitope bound by a domain containing an antibody variable region is included in a number of different antigens, antigen-binding molecules comprising the antibody variable region-containing domain can bind to various antigens that have the epitope.

Epitope

[0321] “Epitope” means an antigenic determinant in an antigen, and refers to an antigen site to which a domain of an antigen-binding molecule including an antibody variable region disclosed herein binds. Thus, for example, the epitope can be defined according to its structure. Alternatively, the epitope may be defined according to the antigen-binding activity of an antigen-binding molecule that recognizes the epitope. When the antigen is a peptide or polypeptide, the

epitope can be specified by the amino acid residues forming the epitope. Alternatively, when the epitope is a sugar chain, the epitope can be specified by its specific sugar chain structure.

[0322] A linear epitope is an epitope that contains an epitope whose primary amino acid sequence is recognized. Such a linear epitope typically contains at least three and most commonly at least five, for example, about 8 to 10 or 6 to 20 amino acids in its specific sequence.

[0323] In contrast to the linear epitope, "conformational epitope" is an epitope in which the primary amino acid sequence containing the epitope is not the only determinant of the recognized epitope (for example, the primary amino acid sequence of a conformational epitope is not necessarily recognized by an epitope-defining antibody). Conformational epitopes may contain a greater number of amino acids compared to linear epitopes. A conformational epitope-recognizing antibody recognizes the three-dimensional structure of a peptide or protein. For example, when a protein molecule folds and forms a three-dimensional structure, amino acids and/or polypeptide main chains that form a conformational epitope become aligned, and the epitope is made recognizable by the antibody. Methods for determining epitope conformations include, for example, X ray crystallography, two-dimensional nuclear magnetic resonance, site-specific spin labeling, and electron paramagnetic resonance, but are not limited thereto. See, for example, *Epitope Mapping Protocols in Methods in Molecular Biology* (1996), Vol. 66, Morris (ed.).

[0324] A method for confirming binding to an epitope by a test antigen-binding molecule comprising a domain that contains an antibody variable region having GPC3-binding activity is exemplified below, and a method for confirming binding to an epitope by a test antigen-binding molecule comprising a domain that contains an antibody variable region having T-cell receptor complex-binding activity may also be performed suitably according to the examples below.

[0325] For example, recognition of a linear epitope present in the GPC3 molecule by a test antigen-binding molecule comprising a domain that contains an antibody variable region having GPC3-binding activity can be confirmed below. A linear peptide comprising the amino acid sequence constituting the extracellular domain of GPC3 is synthesized for the above-mentioned objective. The peptide may be synthesized chemically. Alternatively, it can be obtained by genetic engineering methods using a region in the cDNA of GPC3 that encodes an amino acid sequence corresponding to the extracellular domain. Next, the binding activity between a linear peptide comprising the amino acid sequence constituting the extracellular domain and the test antigen-binding molecule comprising a domain that contains an antibody variable region having GPC3-binding activity is evaluated. For example, ELISA which uses an immobilized linear peptide as the antigen may enable evaluation of the binding activity of the antigen-binding molecule towards the peptide. Alternatively, binding activity towards the linear peptide may be elucidated based on the level of inhibition caused by the linear peptide in the binding of the antigen-binding molecule to GPC3-expressing cells. These tests may elucidate the binding activity of the antigen-binding molecules toward the linear peptide.

[0326] Furthermore, recognition of the three-dimensional structure of the epitope by a test antigen-binding molecule comprising a domain that contains an antibody variable

region having GPC3-binding activity can be confirmed below. GPC3-expressing cells are prepared for the above-mentioned objective. For example, when the test antigen-binding molecule comprising a domain that contains an antibody variable region having GPC3-binding activity contacts GPC3-expressing cells, it binds strongly to the cells, but on the other hand, there are cases when the antigen-binding molecule does not substantially bind to the immobilized linear peptide comprising the amino acid sequence constituting the extracellular domain of GPC3. In these cases, "does not substantially bind" refers to a binding activity of 80% or less, generally 50% or less, preferably 30% or less, and particularly preferably 15% or less relative to the binding activity towards human GPC3-expressing cells.

[0327] Methods for assaying the binding activity of a test antigen-binding molecule containing a GPC3 antigen-binding domain towards GPC3-expressing cells include, for example, the methods described in *Antibodies: A Laboratory Manual* (Ed Harlow, David Lane, Cold Spring Harbor Laboratory (1988) 359-420). Specifically, the assessment can be performed based on the principle of ELISA or fluorescence activated cell sorting (FACS) using GPC3-expressing cells as antigen.

[0328] In the ELISA format, the binding activity of a test antigen-binding molecule containing a GPC3 antigen-binding domain towards GPC3-expressing cells can be assessed quantitatively by comparing the levels of signal generated by enzymatic reaction. Specifically, a test antigen-binding molecule is added to an ELISA plate onto which GPC3-expressing cells are immobilized. Then, the test antigen-binding molecule bound to the cells is detected using an enzyme-labeled antibody that recognizes the test antigen-binding molecule. Alternatively, when FACS is used, a dilution series of a test antigen-binding molecule is prepared, and the antibody binding titer for GPC3-expressing cells can be determined to compare the binding activity of the test antigen-binding molecule towards GPC3-expressing cells.

[0329] The binding of a test antigen-binding molecule towards an antigen expressed on the surface of cells suspended in buffer or the like can be detected using a flow cytometer. Known flow cytometers include, for example, the following devices: FACSCanto™ II; FACSAria™; FACSArray™; FACS Vantage™ SE; FACSCalibur™ (all are trade names of BD Biosciences); EPICS ALTRA HyPer-Sort; Cytomics FC 500; EPICS XL-MCL ADC EPICS XL ADC; Cell Lab Quanta/Cell Lab Quanta SC (all are trade names of Beckman Coulter)

[0330] Preferable methods for assaying the binding activity of a test antigen-binding molecule containing a GPC3 antigen-binding domain towards an antigen include, for example, the following method. First, GPC3-expressing cells are reacted with a test antigen-binding molecule, and then this is stained with an FITC-labeled secondary antibody that recognizes the polypeptide complex. The test antigen-binding molecule is appropriately diluted with a suitable buffer to prepare the complex at a desired concentration. For example, the complex can be used at a concentration within the range of 10 µg/ml to 10 ng/ml. Then, the fluorescence intensity and cell count are determined using FACSCalibur (BD). The fluorescence intensity obtained by analysis using the CELL QUEST Software (BD), i.e., the Geometric Mean value, reflects the quantity of antibody bound to cells. That

is, the binding activity of a test antigen-binding molecule, which is represented by the quantity of the test antigen-binding molecule bound, can be determined by measuring the Geometric Mean value.

[0331] Whether a test antigen-binding molecule containing a GPC3 antigen-binding domain shares a common epitope with another antigen-binding molecule can be assessed based on the competition between the two complexes for the same epitope. The competition between antigen-binding molecules can be detected by cross-blocking assay or the like. For example, the competitive ELISA assay is a preferred cross-blocking assay.

[0332] Specifically, in cross-blocking assay, the GPC3 protein immobilized to the wells of a microtiter plate is pre-incubated in the presence or absence of a candidate competitor antigen-binding molecule, and then a test antigen-binding molecule is added thereto. The quantity of test antigen-binding molecule bound to the GPC3 protein in the wells is indirectly correlated with the binding ability of a candidate competitor antigen-binding molecule that competes for the binding to the same epitope. That is, the greater the affinity of the competitor antigen-binding molecule for the same epitope, the lower the binding activity of the test antigen-binding molecule towards the GPC3 protein-coated wells.

[0333] The quantity of the test antigen-binding molecule bound to the wells via the GPC3 protein can be readily determined by labeling the antigen-binding molecule in advance. For example, a biotin-labeled antigen-binding molecule is measured using an avidin/peroxidase conjugate and appropriate substrate. In particular, cross-blocking assay that uses enzyme labels such as peroxidase is called "competitive ELISA assay". The antigen-binding molecule can also be labeled with other labeling substances that enable detection or measurement. Specifically, radiolabels, fluorescent labels, and such are known.

[0334] When the candidate competitor antigen-binding molecule can block the binding by a test antigen-binding molecule containing a GPC3 antigen-binding domain by at least 20%, preferably at least 20 to 50%, and more preferably at least 50% compared to the binding activity in a control experiment conducted in the absence of the competitor antigen-binding molecule, the test antigen-binding molecule is determined to substantially bind to the same epitope bound by the competitor antigen-binding molecule, or compete for the binding to the same epitope.

[0335] When the structure of an epitope bound by a test antigen-binding molecule containing a GPC3 antigen-binding domain has already been identified, whether the test and control antigen-binding molecules share a common epitope can be assessed by comparing the binding activities of the two antigen-binding molecules towards a peptide prepared by introducing amino acid mutations into the peptide forming the epitope.

[0336] To measure the above binding activities, for example, the binding activities of test and control antigen-binding molecules towards a linear peptide into which a mutation is introduced are compared in the above ELISA format. Besides the ELISA methods, the binding activity towards the mutant peptide bound to a column can be determined by flowing test and control antigen-binding molecules in the column, and then quantifying the antigen-binding molecule eluted in the elution solution. Methods for

adsorbing a mutant peptide to a column, for example, in the form of a GST fusion peptide, are known.

[0337] Alternatively, when the identified epitope is a conformational epitope, whether test and control antigen-binding molecules share a common epitope can be assessed by the following method. First, GPC3-expressing cells and cells expressing GPC3 with a mutation introduced into the epitope are prepared. The test and control antigen-binding molecules are added to a cell suspension prepared by suspending these cells in an appropriate buffer such as PBS. Then, the cell suspensions are appropriately washed with a buffer, and an FITC-labeled antibody that recognizes the test and control antigen-binding molecules is added thereto. The fluorescence intensity and number of cells stained with the labeled antibody are determined using FACSCalibur (BD). The test and control polypeptide complexes are appropriately diluted using a suitable buffer, and used at desired concentrations. For example, they may be used at a concentration within the range of 10 µg/ml to 10 ng/ml. The fluorescence intensity determined by analysis using the CELL QUEST Software (BD), i.e., the Geometric Mean value, reflects the quantity of labeled antibody bound to cells. That is, the binding activities of the test and control antigen-binding molecules, which are represented by the quantity of labeled antibody bound, can be determined by measuring the Geometric Mean value.

[0338] In the above method, whether an antigen-binding molecule does "not substantially bind to cells expressing mutant GPC3" can be assessed, for example, by the following method. First, the test and control antigen-binding molecules bound to cells expressing mutant GPC3 are stained with a labeled antibody. Then, the fluorescence intensity of the cells is determined. When FACSCalibur is used for fluorescence detection by flow cytometry, the determined fluorescence intensity can be analyzed using the CELL QUEST Software. From the Geometric Mean values in the presence and absence of the antigen-binding molecule, the comparison value (Δ Geo-Mean) can be calculated according to the following formula to determine the ratio of increase in fluorescence intensity as a result of the binding by the antigen-binding molecule.

$$\Delta\text{Geo-Mean} = \text{Geo-Mean} (\text{in the presence of the antigen-binding molecule}) / \text{Geo-Mean} (\text{in the absence of the antigen-binding molecule})$$

[0339] The Geometric Mean comparison value (Δ Geo-Mean value for the mutant GPC3 molecule) determined by the above analysis, which reflects the quantity of a test antigen-binding molecule bound to cells expressing mutant GPC3, is compared to the Δ Geo-Mean comparison value that reflects the quantity of the test antigen-binding molecule bound to GPC3-expressing cells. In this case, the concentrations of the test antigen-binding molecule used to determine the Δ Geo-Mean comparison values for GPC3-expressing cells and cells expressing mutant GPC3 are particularly preferably adjusted to be equal or substantially equal. An antigen-binding molecule that has been confirmed to recognize an epitope in GPC3 is used as a control antigen-binding molecule.

[0340] If the Δ Geo-Mean comparison value of a test antigen-binding molecule for cells expressing mutant GPC3 is smaller than the Δ Geo-Mean comparison value of the test antigen-binding molecule for GPC3-expressing cells by at least 80%, preferably 50%, more preferably 30%, and particularly preferably 15%, then the test polypeptide complex

"does not substantially bind to cells expressing mutant GPC3". The formula for determining the Geo-Mean (Geometric Mean) value is described in the CELL QUEST Software User's Guide (BD biosciences). When the comparison shows that the comparison values are substantially equivalent, the epitope for the test and control antigen-binding molecules can be determined to be the same.

Variable Fragment (Fv)

[0341] Herein, the term "variable fragment (Fv)" refers to the minimum unit of an antibody-derived antigen-binding domain that is composed of a pair of the antibody light chain variable region (VL) and antibody heavy chain variable region (VH). In 1988, Skerra and Pluckthun found that homogeneous and active antibodies can be prepared from the *E. coli* periplasm fraction by inserting an antibody gene downstream of a bacterial signal sequence and inducing expression of the gene in *E. coli* (Science (1988) 240 (4855), 1038-1041). In the Fv prepared from the periplasm fraction, VH associates with VL in a manner so as to bind to an antigen.

[0342] Herein, Fv preferably includes, for example, a pair of Fv which is an antigen-binding molecule or such comprising:

[0343] (1) a bivalent antigen-binding domain which is a bivalent scFv, wherein one monovalent scFv of the bivalent scFv is linked to one polypeptide forming an Fc domain by a heavy-chain Fv fragment forming a CD3-binding domain, and the other monovalent scFv is linked to the other polypeptide forming an Fc domain by a light-chain Fv fragment forming a CD3-binding domain;

[0344] (2) a domain comprising an Fc domain that has no Fcy receptor-binding activity, and which is derived from amino acids forming the Fc domain of IgG1, IgG2a, IgG3, or IgG4; and

[0345] (3) at least a monovalent CD3-binding domain,

[0346] wherein the light-chain and heavy-chain Fv fragments associate to form a CD3-binding domain such that it can bind to the CD3 antigen.

scFv, Single-Chain Antibody, and Sc(Fv)2

[0347] Herein, the terms "scFv", "single-chain antibody", and "sc(Fv)2" all refer to an antibody fragment of a single polypeptide chain that contains variable regions derived from the heavy and light chains, but not the constant region. In general, a single-chain antibody also contains a polypeptide linker between the VH and VL domains, which enables formation of a desired structure that is thought to allow antigen binding. The single-chain antibody is discussed in detail by Pluckthun in "The Pharmacology of Monoclonal Antibodies, Vol. 113, Rosenberg and Moore, eds., Springer-Verlag, New York, 269-315 (1994)". See also International Patent Publication WO 1988/001649; U.S. Pat. Nos. 4,946,778 and 5,260,203. In a particular embodiment, the single-chain antibody can be bispecific and/or humanized.

[0348] scFv is an antigen-binding domain in which VH and VL forming Fv are linked together by a peptide linker (Proc. Natl. Acad. Sci. U.S.A. (1988) 85 (16), 5879-5883). VH and VL can be retained in close proximity by the peptide linker.

[0349] sc(Fv)2 is a single-chain antibody in which four variable regions of two VL and two VH are linked by linkers such as peptide linkers to form a single chain (J Immunol. Methods (1999) 231 (1-2), 177-189). The two VH and two

VL may be derived from different monoclonal antibodies. Such sc(Fv)2 preferably includes, for example, a bispecific sc(Fv)2 that recognizes two epitopes present in a single antigen as disclosed in the Journal of Immunology (1994) 152 (11), 5368-5374. sc(Fv)2 can be produced by methods known to those skilled in the art. For example, sc(Fv)2 can be produced by linking scFv by a linker such as a peptide linker.

[0350] Herein, the form of an antigen-binding domain forming an sc(Fv)2 include an antibody in which the two VH units and two VL units are arranged in the order of VH, VL, VH, and VL ([VH]-linker-[VL]-linker-[VH]-linker-[VL]) beginning from the N terminus of a single-chain polypeptide. The order of the two VH units and two VL units is not limited to the above form, and they may be arranged in any order. Example order of the form is listed below.

[0351] [VL]-linker-[VH]-linker-[VH]-linker-[VL]

[0352] [VH]-linker-[VL]-linker-[VL]-linker-[VH]

[0353] [VH]-linker-[VH]-linker-[VL]-linker-[VL]

[0354] [VL]-linker-[VL]-linker-[VH]-linker-[VH]

[0355] [VL]-linker-[VH]-linker-[VL]-linker-[VH]

[0356] The molecular form of sc(Fv)2 is also described in detail in WO 2006/132352. According to these descriptions, those skilled in the art can appropriately prepare desired sc(Fv)2 to produce the antigen-binding molecules disclosed herein.

[0357] Furthermore, the antigen-binding molecules of the present invention may be conjugated with a carrier polymer such as PEG or an organic compound such as an anticancer agent. Alternatively, a sugar chain addition sequence is preferably inserted into the polypeptide complexes such that the sugar chain produces a desired effect.

[0358] The linkers to be used for linking the variable regions of an antibody comprise arbitrary peptide linkers that can be introduced by genetic engineering, synthetic linkers, and linkers disclosed in, for example, Protein Engineering, 9 (3), 299-305, 1996. However, peptide linkers are preferred in the present invention. The length of the peptide linkers is not particularly limited, and can be suitably selected by those skilled in the art according to the purpose. The length is preferably five amino acids or more (without particular limitation, the upper limit is generally 30 amino acids or less, preferably 20 amino acids or less), and particularly preferably 15 amino acids. When sc(Fv)2 contains three peptide linkers, their length may be all the same or different.

[0359] For example, such peptide linkers include:

Ser

Gly•Ser

Gly•Gly•Ser

Ser•Gly•Gly

Gly•Gly•Gly•Ser

(SEQ ID NO: 15)

Ser•Gly•Gly•Gly

(SEQ ID NO: 16)

Gly•Gly•Gly•Gly•Ser

(SEQ ID NO: 17)

-continued	
Ser•Gly•Gly•Gly•Gly	(SEQ ID NO: 18)
Gly•Gly•Gly•Gly•Gly•Ser	(SEQ ID NO: 19)
Ser•Gly•Gly•Gly•Gly•Gly	(SEQ ID NO: 20)
Gly•Gly•Gly•Gly•Gly•Gly•Ser	(SEQ ID NO: 21)
Ser•Gly•Gly•Gly•Gly•Gly	(SEQ ID NO: 22)
(Gly•Gly•Gly•Gly•Ser (SEQ ID NO: 17)) ⁿ	
(Ser•Gly•Gly•Gly•Gly (SEQ ID NO: 18)) ⁿ	

[0360] where n is an integer of 1 or larger. The length or sequences of peptide linkers can be selected accordingly by those skilled in the art depending on the purpose.

[0361] Synthetic linkers (chemical crosslinking agents) is routinely used to crosslink peptides, and for example:

- [0362] N-hydroxy succinimide (NHS),
- [0363] disuccinimidyl suberate (DSS),
- [0364] bis(sulfosuccinimidyl) suberate (BS³),
- [0365] dithiobis(succinimidyl propionate) (DSP),
- [0366] dithiobis(sulfosuccinimidyl propionate) (DTSSP),
- [0367] ethylene glycol bis(succinimidyl succinate) (EGS),
- [0368] ethylene glycol bis(sulfosuccinimidyl succinate) (sulfo-EGS),
- [0369] disuccinimidyl tartrate (DST), disulfosuccinimidyl tartrate (sulfo-DST),
- [0370] bis [2-(succinimidoxycarbonyloxy)ethyl] sulfone (BSOCOES),
- [0371] and bis [2-(sulfosuccinimidoxycarbonyloxy)ethyl] sulfone (sulfo-BSOCOES). These
- [0372] crosslinking agents are commercially available.
- [0373] In general, three linkers are required to link four antibody variable regions together. The linkers to be used may be of the same type or different types.

Fab, F(ab')2, and Fab'

[0374] "Fab" consists of a single light chain, and a CH1 domain and variable region from a single heavy chain. The heavy chain of Fab molecule cannot form disulfide bonds with another heavy chain molecule.

[0375] "F(ab')2" or "Fab" is produced by treating an immunoglobulin (monoclonal antibody) with a protease such as pepsin and papain, and refers to an antibody fragment generated by digesting an immunoglobulin (monoclonal antibody) at near the disulfide bonds present between the hinge regions in each of the two H chains. For example, papain cleaves IgG upstream of the disulfide bonds present between the hinge regions in each of the two H chains to generate two homologous antibody fragments, in which an L chain comprising VL (L-chain variable region) and CL (L-chain constant region) is linked to an H-chain fragment comprising VH (H-chain variable region) and CH γ 1 (γ 1 region in an H-chain constant region) via a disulfide bond at their C-terminal regions. Each of these two homologous antibody fragments is called Fab'.

[0376] "F(ab')2" consists of two light chains and two heavy chains comprising the constant region of a CH1 domain and a portion of CH2 domains so that disulfide bonds are formed between the two heavy chains. The F(ab')2 forming an antigen-binding molecule disclosed herein can be preferably produced as follows. A whole monoclonal antibody or such comprising a desired antigen-binding domain is partially digested with a protease such as pepsin; and Fc fragments are removed by adsorption onto a Protein A column. The protease is not particularly limited, as long as it can cleave the whole antibody in a selective manner to produce F(ab')2 under an appropriate setup enzyme reaction condition such as pH. Such proteases include, for example, pepsin and ficin.

Fc Domain

[0377] An Fc domain that forms an antigen-binding molecule disclosed herein can be preferably produced in the following manner. An antibody such as a monoclonal antibody is partially digested with a protease such as pepsin. Then, the resulting fragment is adsorbed onto a Protein A or Protein G column, and eluted with an appropriate elution buffer. The protease is not particularly limited, as long as it can cleave antibodies such as monoclonal antibodies under an appropriate setup enzyme reaction condition such as pH. Such proteases include, for example, pepsin and ficin.

[0378] The antigen-binding molecules described herein comprise an Fc domain with reduced Fc γ receptor-binding activity, which includes amino acids forming the Fc domain of IgG1, IgG2, IgG3, or IgG4.

[0379] Antibody isotype is determined according to the structure of the constant region. Constant regions of the isotypes IgG1, IgG2, IgG3, and IgG4 are called C γ 1, C γ 2, C γ 3, and C γ 4, respectively. The amino acid sequences of Fc domain polypeptides forming human C γ 1, C γ 2, C γ 3, and C γ 4 are exemplified in SEQ ID NO:23, 24, 25, and 26, respectively. The relationship between amino acid residues forming each amino acid sequence and Kabat's EU numbering (herein also referred to as EU INDEX) are shown in FIG. 12.

[0380] The Fc domain refers to the region besides F(ab')2 which comprises two light chains and two heavy chains comprising a portion of the constant region that comprises a CH1 domain and a region between the CH1 and CH2 domains so that disulfide bonds are formed between the two heavy chains. The Fc domain forming an antigen-binding molecule disclosed herein can be preferably produced as follows. A monoclonal IgG1, IgG2, IgG3, or IgG4 antibody or the like is partially digested with a protease such as pepsin, followed by elution of the fraction adsorbed onto a Protein A column. The protease is not particularly limited, as long as it can cleave the whole antibody in a selective manner to produce F(ab')2 in an appropriate setup enzyme reaction condition such as pH. Such proteases include, for example, pepsin and ficin.

Fc γ Receptor

[0381] Fc γ receptor refers to a receptor capable of binding to the Fc domain of monoclonal IgG1, IgG2, IgG3, or IgG4 antibodies, and includes all members belonging to the family of proteins substantially encoded by an Fc γ receptor gene. In human, the family includes Fc γ RI (CD64) including isoforms Fc γ RIa, Fc γ RIb and Fc γ RIc; Fc γ RII (CD32)

including isoforms Fc γ RIIa (including allotype H131 and R131), Fc γ RIIb (including Fc γ RIIb-1 and Fc γ RIIb-2), and Fc γ RIIC; and Fc γ RIII (CD16) including isoform Fc γ RIIIa (including allotype V158 and F158) and Fc γ RIIIb (including allotype Fc γ RIIIb-NA1 and Fc γ RIIIb-NA2); as well as all unidentified human Fc γ Rs, Fc γ R isoforms, and allotypes thereof. However, Fc γ receptor is not limited to these examples. Without being limited thereto, Fc γ R includes those derived from humans, mice, rats, rabbits, and monkeys. Fc γ R may be derived from any organisms. Mouse Fc γ R includes, without being limited to, Fc γ RI (CD64), Fc γ RII (CD32), Fc γ RIII (CD16), and Fc γ RIII-2 (CD16-2), as well as all unidentified mouse Fc γ Rs, Fc γ R isoforms, and allotypes thereof. Such preferred Fc γ receptors include, for example, human Fc γ RI (CD64), Fc γ RIIA (CD32), Fc γ RIIB (CD32), Fc γ RIIIA (CD16), and/or Fc γ RIIIB (CD16). The polynucleotide sequence and amino acid sequence of Fc γ RI are shown in SEQ ID NOs: 27 (NM_000566.3) and 28 (NP_000557.1), respectively; the polynucleotide sequence and amino acid sequence of Fc γ RIIA are shown in SEQ ID NOs: 29 (BC020823.1) and 30 (AAH20823.1), respectively; the polynucleotide sequence and amino acid sequence of Fc γ RIIB are shown in SEQ ID NOs: 31 (BC146678.1) and 32 (AAI46679.1), respectively; the polynucleotide sequence and amino acid sequence of Fc γ RIIIA are shown in SEQ ID NOs: 33 (BC033678.1) and 34 (AAH33678.1), respectively; and the polynucleotide sequence and amino acid sequence of Fc γ RIIIB are shown in SEQ ID NOs: 35 (BC128562.1) and 36 (AAI28563.1), respectively (RefSeq accession number is shown in each parentheses). Whether an Fc γ receptor has binding activity to the Fc domain of a monoclonal IgG1, IgG2, IgG3, or IgG4 antibody can be assessed by ALPHA screen (Amplified Luminescent Proximity Homogeneous Assay), surface plasmon resonance (SPR)-based BIACORE method, and others (Proc. Natl. Acad. Sci. USA (2006) 103 (11), 4005-4010), in addition to the above-described FACS and ELISA formats.

[0382] Meanwhile, "Fc ligand" or "effector ligand" refers to a molecule and preferably a polypeptide that binds to an antibody Fc domain, forming an Fc/Fc ligand complex. The molecule may be derived from any organisms. The binding of an Fc ligand to Fc preferably induces one or more effector functions. Such Fc ligands include, but are not limited to, Fc receptors, Fc γ R, Fc α R, Fc ϵ R, FcRn, C1q, and C3, mannabinding lectin, mannose receptor, *Staphylococcus* Protein A, *Staphylococcus* Protein G, and viral Fc γ Rs. The Fc ligands also include Fc receptor homologs (FcRH) (Davis et al., (2002) Immunological Reviews 190, 123-136), which are a family of Fc receptors homologous to Fc γ R. The Fc ligands also include unidentified molecules that bind to Fc.

Fc γ Receptor-Binding Activity

[0383] The impaired binding activity of Fc domain to any of the Fc γ receptors Fc γ I, Fc γ IIA, Fc γ IIB, Fc γ IIIa, and/or Fc γ IIIb can be assessed by using the above-described FACS and ELISA formats as well as ALPHA screen (Amplified Luminescent Proximity Homogeneous Assay) and surface plasmon resonance (SPR)-based BIACORE method (Proc. Natl. Acad. Sci. USA (2006) 103 (11), 4005-4010).

[0384] ALPHA screen is performed by the ALPHA technology based on the principle described below using two types of beads: donor and acceptor beads. A luminescent signal is detected only when molecules linked to the donor beads interact biologically with molecules linked to the

acceptor beads and when the two beads are located in close proximity. Excited by laser beam, the photosensitizer in a donor bead converts oxygen around the bead into excited singlet oxygen. When the singlet oxygen diffuses around the donor beads and reaches the acceptor beads located in close proximity, a chemiluminescent reaction within the acceptor beads is induced. This reaction ultimately results in light emission. If molecules linked to the donor beads do not interact with molecules linked to the acceptor beads, the singlet oxygen produced by donor beads do not reach the acceptor beads and chemiluminescent reaction does not occur.

[0385] For example, a biotin-labeled antigen-binding molecule is immobilized to the donor beads and glutathione S-transferase (GST)-tagged Fc receptor is immobilized to the acceptor beads. In the absence of an antigen-binding molecule comprising a competitive mutant Fc domain, Fc γ receptor interacts with an antigen-binding molecule comprising a wild-type Fc domain, inducing a signal of 520 to 620 nm as a result. The antigen-binding molecule having a non-tagged mutant Fc domain competes with the antigen-binding molecule comprising a wild-type Fc domain for the interaction with Fc γ receptor. The relative binding affinity can be determined by quantifying the reduction of fluorescence as a result of competition. Methods for biotinyling antigen-binding molecules such as antibodies using Sulfo-NHS-biotin or the like are known. Appropriate methods for adding the GST tag to an Fc γ receptor include methods that involve fusing polypeptides encoding Fc γ and GST in-frame, expressing the fused gene using cells introduced with a vector carrying the gene, and then purifying using a glutathione column. The induced signal can be preferably analyzed, for example, by fitting to a one-site competition model based on nonlinear regression analysis using software such as GRAPHPAD PRISM (GraphPad; San Diego).

[0386] One of the substances for observing their interaction is immobilized as a ligand onto the gold thin layer of a sensor chip. When light is shed on the rear surface of the sensor chip so that total reflection occurs at the interface between the gold thin layer and glass, the intensity of reflected light is partially reduced at a certain site (SPR signal). The other substance for observing their interaction is injected as an analyte onto the surface of the sensor chip. The mass of immobilized ligand molecule increases when the analyte binds to the ligand. This alters the refraction index of solvent on the surface of the sensor chip. The change in refraction index causes a positional shift of SPR signal (conversely, the dissociation shifts the signal back to the original position). In the Biacore system, the amount of shift described above (i.e., the change of mass on the sensor chip surface) is plotted on the vertical axis, and thus the change of mass over time is shown as measured data (sensogram). Kinetic parameters (association rate constant (ka) and dissociation rate constant (kd)) are determined from the curve of sensogram, and affinity (KD) is determined from the ratio between these two constants. Inhibition assay is preferably used in the BIACORE methods. Examples of such inhibition assay are described in Proc. Natl. Acad. Sci. USA (2006) 103 (11), 4005-4010.

[0387] Herein, "Fc γ receptor-binding activity is reduced" means, for example, that based on the above-described analysis method the competitive activity of a test antigen-binding molecule is 50% or less, preferably 45% or less, 40% or less, 35% or less, 30% or less, 20% or less, or 15%

or less, and particularly preferably 10% or less, 9% or less, 8% or less, 7% or less, 6% or less, 5% or less, 4% or less, 3% or less, 2% or less, or 1% or less than the competitive activity of a control antigen-binding molecule.

[0388] Antigen-binding molecules comprising the Fc domain of a monoclonal IgG1, IgG2, IgG3, or IgG4 antibody can be appropriately used as control antigen-binding molecules. The Fc domain structures are shown in SEQ ID NOs: 37 (A is added to the N terminus of RefSeq accession number AAC82527.1), 38 (A is added to the N terminus of RefSeq accession number AAB59393.1), 25 (A is added to the N terminus of RefSeq accession number CAA27268.1), and 39 (A is added to the N terminus of RefSeq accession number AAB59394.1). Furthermore, when an antigen-binding molecule comprising an Fc domain mutant of an antibody of a particular isotype is used as a test substance, the effect of the mutation of the mutant on the Fcγ receptor-binding activity is assessed using as a control an antigen-binding molecule comprising an Fc domain of the same isotype. As described above, antigen-binding molecules comprising an Fc domain mutant whose Fcγ receptor-binding activity has been judged to be reduced are appropriately prepared.

[0389] Such known mutants include, for example, mutants having a deletion of amino acids 231A-238S (EU numbering) (WO 2009/011941), as well as mutants C226S, C229S, P238S, (C220S) (J. Rheumatol. (2007) 34, 11); C226S and C229S (Hum. Antibod. Hybridomas (1990) 1 (1), 47-54); C226S, C229S, E233P, L234V, and L235A (Blood (2007) 109, 1185-1192).

[0390] Specifically, the preferred antigen-binding molecules include those comprising an Fc domain with a substitution of the amino acid at position 220, 226, 229, 231, 232, 233, 234, 235, 236, 237, 238, 239, 240, 264, 265, 266, 267, 269, 270, 295, 296, 297, 298, 299, 300, 325, 327, 328, 329, 330, 331, or 332 (EU numbering) in the amino acids forming the Fc domain of an antibody of a particular isotype. The isotype of antibody from which the Fc domain originates is not particularly limited, and it is possible to use an appropriate Fc domain derived from a monoclonal IgG1, IgG2, IgG3, or IgG4 antibody. It is preferable to use Fc domains derived from IgG1 antibodies.

[0391] The preferred antigen-binding molecules include, for example, those comprising an Fc domain which has any one of the substitutions shown below, whose positions are specified according to EU numbering (each number represents the position of an amino acid residue in the EU numbering; and the one-letter amino acid symbol before the number represents the amino acid residue before substitution, while the one-letter amino acid symbol after the number represents the amino acid residue before the substitution) in the amino acids forming the Fc domain of IgG1 antibody:

- [0392] (a) L234F, L235E, P331S;
- [0393] (b) C226S, C229S, P238S;
- [0394] (c) C226S, C229S;
- [0395] (d) C226S, C229S, E233P, L234V, L235A;
- [0396] (e) L234A, L235A or L235R, N297A;
- [0397] (f) L235A or L235R, S239K, N297A
- [0398] as well as those having an Fc domain which has a deletion of the amino acid sequence at positions 231 to 238.

[0399] Furthermore, the preferred antigen-binding molecules also include those comprising an Fc domain that has

any one of the substitutions shown below, whose positions are specified according to EU numbering in the amino acids forming the Fc domain of an IgG2 antibody:

- [0400] (g) H268Q, V309L, A330S, and P331S;
- [0401] (h) V234A;
- [0402] (i) G237A;
- [0403] (j) V234A and G237A;
- [0404] (k) A235E and G237A;
- [0405] (l) V234A, A235E, and G237A. Each number represents the position of an amino acid residue in EU numbering; and the one-letter amino acid symbol before the number represents the amino acid residue before substitution, while the one-letter amino acid symbol after the number represents the amino acid residue before the substitution.

[0406] Furthermore, the preferred antigen-binding molecules also include those comprising an Fc domain that has any one of the substitutions shown below, whose positions are specified according to EU numbering in the amino acids forming the Fc domain of an IgG3 antibody:

- [0407] (m) F241A;
- [0408] (n) D265A;
- [0409] (o) V264A. Each number represents the position of an amino acid residue in EU numbering; and the one-letter amino acid symbol before the number represents the amino acid residue before substitution, while the one-letter amino acid symbol after the number represents the amino acid residue before the substitution.

[0410] Furthermore, the preferred antigen-binding molecules also include those comprising an Fc domain that has any one of the substitutions shown below, whose positions are specified according to EU numbering in the amino acids forming the Fc domain of an IgG4 antibody:

- [0411] (p) L235A, G237A, and E318A;
- [0412] (q) L235E;
- [0413] (r) F234A and L235A. Each number represents the position of an amino acid residue in EU numbering; and the one-letter amino acid symbol before the number represents the amino acid residue before substitution, while the one-letter amino acid symbol after the number represents the amino acid residue before the substitution.

[0414] The other preferred antigen-binding molecules include, for example, those comprising an Fc domain in which any amino acid at position 233, 234, 235, 236, 237, 327, 330, or 331 (EU numbering) in the amino acids forming the Fc domain of an IgG1 antibody is substituted with an amino acid of the corresponding position in EU numbering in the corresponding IgG2 or IgG4.

[0415] The preferred antigen-binding molecules also include, for example, those comprising an Fc domain in which any one or more of the amino acids at positions 234, 235, and 297 (EU numbering) in the amino acids forming the Fc domain of an IgG1 antibody is substituted with other amino acids. The type of amino acid after substitution is not particularly limited; however, the antigen-binding molecules comprising an Fc domain in which any one or more of the amino acids at positions 234, 235, and 297 are substituted with alanine are particularly preferred.

[0416] The preferred antigen-binding molecules also include, for example, those comprising an Fc domain in which an amino acid at position 265 (EU numbering) in the amino acids forming the Fc domain of an IgG1 antibody is

substituted with another amino acid. The type of amino acid after substitution is not particularly limited; however, antigen-binding molecules comprising an Fc domain in which an amino acid at position 265 is substituted with alanine are particularly preferred.

Multispecific Antigen-Binding Molecule

[0417] Examples of a preferred embodiment of the “multispecific antigen-binding molecule” of the present invention include multispecific antibodies. When an Fc region with reduced Fc γ receptor-binding activity is used as the multispecific antibody Fc region, an Fc region derived from the multispecific antibody may be used appropriately. Bispecific antibodies are particularly preferred as the multispecific antibodies of the present invention. In this case, a bispecific antibody is an antibody having two different specificities. IgG-type bispecific antibodies can be secreted from a hybrid hybridoma (quadroma) produced by fusing two types of hybridomas that produce IgG antibodies (Milstein et al., Nature (1983) 305, 537-540).

[0418] Furthermore, IgG-type bispecific antibodies are secreted by introducing the genes of L chains and H chains constituting the two types of IgGs of interest, i.e. a total of four genes, into cells, and co-expressing them. However, the number of combinations of H and L chains of IgG that can be produced by these methods is theoretically ten combinations. Accordingly, it is difficult to purify an IgG comprising the desired combination of H and L chains from ten types of IgGs. Furthermore, theoretically the amount of secretion of the IgG having the desired combination will decrease remarkably, and therefore large-scale culturing will be necessary, and production costs will increase further.

[0419] Therefore, techniques for promoting the association among H chains and between L and H chains having the desired combinations can be applied to the multispecific antigen-binding molecules of the present invention.

[0420] For example, techniques for suppressing undesired H-chain association by introducing electrostatic repulsion at the interface of the second constant region or the third constant region of the antibody H chain (CH2 or CH3) can be applied to multispecific antibody association (WO 2006/106905).

[0421] In the technique of suppressing unintended H-chain association by introducing electrostatic repulsion at the interface of CH2 or CH3, examples of amino acid residues in contact at the interface of the other constant region of the H chain include regions corresponding to the residues at EU numbering positions 356, 439, 357, 370, 399, and 409 in the CH3 region.

[0422] More specifically, examples include an antibody comprising two types of H-chain CH3 regions, in which one to three pairs of amino acid residues in the first H-chain CH3 region, selected from the pairs of amino acid residues indicated in (1) to (3) below, carry the same type of charge: (1) amino acid residues comprised in the H chain CH3 region at EU numbering positions 356 and 439; (2) amino acid residues comprised in the H-chain CH3 region at EU numbering positions 357 and 370; and (3) amino acid residues comprised in the H-chain CH3 region at EU numbering positions 399 and 409.

[0423] Furthermore, the antibody may be an antibody in which pairs of the amino acid residues in the second H-chain CH3 region which is different from the first H-chain CH3 region mentioned above, are selected from the aforemen-

tioned pairs of amino acid residues of (1) to (3), wherein the one to three pairs of amino acid residues that correspond to the aforementioned pairs of amino acid residues of (1) to (3) carrying the same type of charges in the first H-chain CH3 region mentioned above carry opposite charges from the corresponding amino acid residues in the first H-chain CH3 region mentioned above.

[0424] Each of the amino acid residues indicated in (1) to (3) above come close to each other during association. Those skilled in the art can find out positions that correspond to the above-mentioned amino acid residues of (1) to (3) in a desired H-chain CH3 region or H-chain constant region by homology modeling and such using commercially available software, and amino acid residues of these positions can be appropriately subjected to modification.

[0425] In the antibodies mentioned above, “charged amino acid residues” are preferably selected, for example, from amino acid residues included in either one of the following groups: (a) glutamic acid (E) and aspartic acid (D); and (b) lysine (K), arginine (R), and histidine (H).

[0426] In the above-mentioned antibodies, the phrase “carrying the same charge” means, for example, that all of the two or more amino acid residues are selected from the amino acid residues included in either one of groups (a) and (b) mentioned above. The phrase “carrying opposite charges” means, for example, that when at least one of the amino acid residues among two or more amino acid residues is selected from the amino acid residues included in either one of groups (a) and (b) mentioned above, the remaining amino acid residues are selected from the amino acid residues included in the other group.

[0427] In a preferred embodiment, the antibodies mentioned above may have their first H-chain CH3 region and second H-chain CH3 region crosslinked by disulfide bonds.

[0428] In the present invention, amino acid residues subjected to modification are not limited to the above-mentioned amino acid residues of the antibody variable regions or the antibody constant regions. Those skilled in the art can identify the amino acid residues that form an interface in mutant polypeptides or heterooligomers by homology modeling and such using commercially available software; and amino acid residues of these positions can then be subjected to modification so as to regulate the association.

[0429] Other known techniques can also be used for the association of multispecific antibodies of the present invention. Fc region-containing polypeptides comprising different amino acids can be efficiently associated with each other by substituting an amino acid side chain present in one of the H-chain Fc regions of the antibody with a larger side chain (knob), and substituting an amino acid side chain present in the corresponding Fc region of the other H chain with a smaller side chain (hole) to allow placement of the knob within the hole (WO 1996/027011; Ridgway J B et al., Protein Engineering (1996) 9, 617-621; Merchant A. M. et al. Nature Biotechnology (1998) 16, 677-681; and US20130336973).

[0430] In addition, other known techniques can also be used for formation of multispecific antibodies of the present invention. Association of polypeptides having different sequences can be induced efficiently by complementary association of CH3 using a strand-exchange engineered domain CH3 produced by changing part of one of the H-chain CH3s of an antibody to a corresponding IgA-derived sequence and introducing a corresponding IgA-

derived sequence into the complementary portion of the other H-chain CH3 (Protein Engineering Design & Selection, 23; 195-202, 2010). This known technique can also be used to efficiently form multispecific antibodies of interest.

[0431] In addition, technologies for antibody production using association of antibody CH1 and CL and association of VH and VL as described in WO 2011/028952, WO 2014/018572, and Nat Biotechnol. 2014 February; 32 (2): 191-8; technologies for producing bispecific antibodies using separately prepared monoclonal antibodies in combination (Fab Arm Exchange) as described in WO 2008/119353 and WO 2011/131746; technologies for regulating association between antibody heavy-chain CH3s as described in WO 2012/058768 and WO 2013/063702; technologies for producing bispecific antibodies composed of two types of light chains and one type of heavy chain as described in WO 2012/023053; technologies for producing bispecific antibodies using two bacterial cell strains that individually express one of the chains of an antibody comprising a single H chain and a single L chain as described by Christoph et al. (Nature Biotechnology Vol. 31, p 753-758 (2013)); and such may be used for the formation of multispecific antibodies.

[0432] Alternatively, even when a multispecific antibody of interest cannot be formed efficiently, a multispecific antibody of the present invention can be obtained by separating and purifying the multispecific antibody of interest from the produced antibodies. For example, a method for enabling purification of two types of homomeric forms and the heteromeric antibody of interest by ion-exchange chromatography by imparting a difference in isoelectric points by introducing amino acid substitutions into the variable regions of the two types of H chains has been reported (WO 2007/114325). To date, as a method for purifying heteromeric antibodies, methods using Protein A to purify a heterodimeric antibody comprising a mouse IgG2a H chain that binds to Protein A and a rat IgG2b H chain that does not bind to Protein A have been reported (WO 1998/050431 and WO 1995/033844). Furthermore, a heterodimeric antibody can be purified efficiently on its own by using H chains comprising substitution of amino acid residues at EU numbering positions 435 and 436, which is the IgG-Protein A binding site, with Tyr, His, or such which are amino acids that yield a different Protein A affinity, or using H chains with a different protein A affinity obtained according to the method of Reference Example 5, to change the interaction of each of the H chains with Protein A, and then using a Protein A column.

[0433] Alternatively, a common L chain that can provide binding ability to a plurality of different H chains can be obtained and used as the common L chain of a multispecific antibody. Efficient expression of a multispecific IgG can be achieved by introducing the genes of such a common L chain and a plurality of different H chains into cells to express the IgG (Nature Biotechnology (1998) 16, 677-681). A method for selecting a common L chain that shows a strong binding ability to any of the different H chains can also be used when selecting the common H chain (WO 2004/065611).

[0434] Furthermore, an Fc region whose Fc region C-terminal heterogeneity has been improved can be appropriately used as an Fc region of the present invention. More specifically, the present invention provides Fc regions produced by deleting glycine at position 446 and lysine at position 447 as

specified by EU numbering from the amino acid sequences of two polypeptides constituting an Fc region derived from IgG1, IgG2, IgG3, or IgG4.

[0435] A plurality, such as two or more, of these technologies can be used in combination. Furthermore, these technologies can be appropriately and separately applied to the two H chains to be associated. Furthermore, these techniques can be used in combination with the above-mentioned Fc region which has reduced binding activity to an Fcγ receptor. Furthermore, an antigen-binding molecule of the present invention may be a molecule produced separately so that it has the same amino acid sequence, based on the antigen-binding molecule subjected to the above-described modifications.

[0436] An appropriate multispecific antigen-binding molecule of the present invention comprises

[0437] (1) a domain comprising an antibody variable region having glycan 3-binding activity;

[0438] (2) a domain comprising an antibody variable region having T-cell receptor complex-binding activity; and

[0439] (3) a domain comprising an Fc region with reduced Fcγ receptor-binding activity mentioned above, without limitation to its structure.

[0440] In the present invention, each of the above-mentioned domains can be linked directly by peptide bonds. For example, when using F(ab')2 as the domain comprising an antibody variable region of (1) and (2), and these Fc regions as the domain comprising an Fc region with reduced Fcγ receptor-binding activity of (3), the polypeptides formed by linking the antibody variable region-containing domains of (1) and (2) and the Fc region-containing domain of (3) by peptide bonds will form an antibody structure. Such antibodies can be produced by purification from the above-mentioned hybridoma culture medium, and also by purifying antibodies from the culture medium of desired host cells that stably carry polynucleotides encoding the polypeptides constituting the antibody.

[0441] Examples of a preferred antibody H-chain variable region of the present invention contained in the antibody variable region having glycan 3-binding activity comprises the antibody H-chain variable regions of Table 1, or antibody H-chain variable regions having CDR sequences whose CDR1, CDR2, and CDR3 amino acid sequences are the same as the CDR1, CDR2, and CDR3 amino acid sequences contained in the H-chain variable regions of Table 1, or antibody H-chain variable regions which are functionally equivalent to the above-mentioned variable regions.

TABLE 1

Sequence Name	SEQ ID NO:
H0000	40
GCH003	170
GCH005	171
GCH006	172
GCH007	173
GCH008	174
GCH010	175
GCH012	176
GCH013	177
GCH014	178
GCH015	179
GCH016	180
GCH019	181
GCH022	182

TABLE 1-continued

Sequence Name	SEQ ID NO:
GCH023	183
GCH025	184
GCH026	185
GCH027	186
GCH029	187
GCH032	188
GCH034	189
GCH035	190
GCH039	191
GCH040	192
GCH042	193
GCH043	194
GCH045	195
GCH053	196
GCH054	197
GCH055	198
GCH056	199
GCH057	200
GCH059	201
GCH060	202
GCH061	203
GCH062	204
GCH064	205
GCH065	206
GCH066	207
GCH067	208
GCH068	209
GCH073	210
GCH094	211
GCH098	212
GCH099	213
GCH100	214
H0610	215

TABLE 2-continued

Sequence Name	SEQ ID NO:
TR01H017	80
TR01H018	81
TR01H019	82
TR01H020	83
TR01H021	84
TR01H022	85
TR01H023	86
TR01H024	87
TR01H025	88
TR01H026	89
TR01H027	90
TR01H028	91
TR01H029	92
TR01H030	93
TR01H031	94
TR01H032	95
TR01H033	96
TR01H034	97
TR01H035	98
TR01H036	99
TR01H037	100
TR01H038	101
TR01H039	102
TR01H040	103
TR01H041	104
TR01H042	105
TR01H043	106
TR01H044	107
TR01H045	108
TR01H046	109
TR01H047	110
TR01H048	111
TR01H049	112
TR01H050	113
TR01H051	114
TR01H052	115
TR01H053	116
TR01H054	117
TR01H055	118
TR01H056	119
TR01H057	120
TR01H058	121
TR01H061	122
TR01H062	123
TR01H063	124
TR01H064	125
TR01H065	126
TR01H066	127
TR01H067	128
TR01H068	129
TR01H069	130
TR01H070	131
TR01H071	132
TR01H072	133
TR01H073	134
TR01H074	135
TR01H075	136
TR01H076	137
TR01H077	138
TR01H079	139
TR01H080	140
TR01H081	141
TR01H082	142
TR01H083	143
TR01H084	144
TR01H090	145
TR01H091	146
TR01H092	147
TR01H093	148
TR01H094	149
TR01H095	150
TR01H096	151
TR01H097	152
TR01H098	153
TR01H099	154

TABLE 2

Sequence Name	SEQ ID NO:
hCE115HA	52
CE115HA177	64
CE115HA178	65
CE115HA179	66
CE115HA180	67
hCE115HAa	68
TR01H006	69
TR01H007	70
TR01H008	71
TR01H009	72
TR01H010	73
TR01H011	74
TR01H012	75
TR01H013	76
TR01H014	77
TR01H015	78
TR01H016	79

TABLE 2-continued

Sequence Name	SEQ ID NO:
TR01H100	155
TR01H101	156
TR01H102	157
TR01H103	158
TR01H104	159
TR01H105	160
TR01H106	161
TR01H107	162
TR01H108	163
TR01H109	164
TR01H110	165
TR01H111	166
TR01H112	167
TR01H113	168
TR01H114	169
TR01H001	420
TR01H002	421
TR01H003	422
TR01H004	423
rCE115H	424
CE115HA121	425
CE115HA122	426
CE115HA124	427
CE115HA192	428
CE115HA236	429
CE115HA251	430
CE115HA252	431

[0443] The relationship between the CDR regions of the amino acid residues constituting the antibody H chain amino acid sequence and Kabat numbering is as shown in FIG. 13-1 and FIG. 13-2.

[0444] For the antibody L-chain variable regions contained in the antibody variable region having glypican 3-binding activity and the antibody variable region having T-cell receptor complex-binding activity of the present invention, it is preferable to obtain a common L chain that may provide a binding activity to the H chain having glypican3-binding activity and a binding activity to the H chain having T-cell receptor complex, and to use this as the common L-chain variable region of the multispecific antigen-binding molecule.

[0445] Examples of the common L-chain variable region to be used in the present invention include the L-chain variable regions of Table 3, antibody L-chain variable regions having CDR sequences whose CDR1, CDR2, and CDR3 amino acid sequences are the same as the CDR1, CDR2, and CDR3 amino acid sequences contained in the antibody L-chain variable regions of Table 3, and antibody L-chain variable regions that are functionally equivalent to the above-mentioned variable regions.

TABLE 3

Sequence Name	SEQ ID NO:
L0000	53
L0002	217
L0003	218
L0006	219
L0007	220
L0008	221
L0009	222
L0011	223
L0012	224
L0013	225
L0014	226
L0015	227

TABLE 3-continued

Sequence Name	SEQ ID NO:
L0016	228
L0032	229
L0038	230
L0039	231
L0041	232
L0042	233
L0043	234
L0044	235
L0045	236
L0046	237
L0047	238
L0062	239
L0063	240
L0064	241
L0065	242
L0066	243
L0069	244
L0075	245
L0079	246
L0082	247
L0085	248
L0089	249
L0090	250
L0091	251
L0093	252
L0104	253
L0106	254
L0107	255
L0109	256
L0113	257
L0115	258
L0117	259
L0120	260
L0122	261
L0123	262
L0124	263
L0125	264
L0126	265
L0127	266
L0129	267
L0132	268
L0134	269
L0136	270
L0137	271
L0138	272
L0139	273
L0140	274
L0141	275
L0143	276
L0144	277
L0145	278
L0147	279
L0148	280
L0149	281
L0151	282
L0152	283
L0154	284
L0155	285
L0157	286
L0160	287
L0161	288
L0163	289
L0167	290
L0168	291
L0173	292
L0175	293
L0180	294
L0181	295
L0186	296
L0187	297
L0200	298
L0201	299
L0202	300
L0203	301
L0204	302

TABLE 3-continued

Sequence Name	SEQ ID NO:
L0205	303
L0206	304
L0207	305
L0208	306
L0209	307
L0210	308
L0211	309
L0212	310
L0213	311
L0214	312
L0215	313
L0216	314
L0217	315
L0218	316
L0219	317
L0220	318
L0222	319
L0223	320
L0224	321
L0226	322
L0227	323
L0228	324
L0229	325
L0230	326
L0231	327
L0232	328
L0233	329
L0234	330
L0235	331
L0236	332
L0237	333
L0238	334
L0239	335
L0240	336
L0241	337
L0242	338
L0243	339
L0246	340
L0247	341
L0248	342
L0249	343
L0250	344
L0258	345
L0259	346
L0260	347
L0261	348
L0262	349
L0263	350
L0264	351
L0265	352
L0266	353
L0267	354
L0268	355
L0269	356
L0270	357
L0271	358
L0272	359

[0446] The relationship between the CDR regions of the amino acid residues constituting the antibody L-chain amino acid sequence and Kabat numbering is as shown in FIG. 14.

[0447] In the present invention, the phrase “functionally equivalent” means that the binding affinities for an antigen are equivalent, or alternatively, it means that the cytotoxic activities against glypcan 3-expressing cells or tissues containing these cells are equivalent when it is used as a multispecific antigen-binding molecule. The binding affinity and cytotoxic activity can be measured based on the description herein. The cells used for measurement of cytotoxic activity may be the desired GPC3-expressing cells or a desired tissue containing these cells, and for example, PC-10 or NCI-H446 which are GPC3-expressing human cancer cell

lines can be used. Regarding the antibody constant regions, the phrase may mean that the decreases in Fcγ receptor-binding activity are equivalent.

[0448] For example, an antibody H-chain variable region functionally equivalent to the antibody H chain variable region described herein (i.e., the original H chain variable region) means that this region has the same binding affinity when it is combined with the antibody L-chain variable region described herein which forms a pair with the original H chain, or alternatively that the region has the same cytotoxic activity towards glypcan 3-expressing cells or a tissue containing these cells when used for a multispecific antigen-binding molecule. Furthermore, an antibody L-chain variable region functionally equivalent to the antibody L-chain variable region described herein (i.e., the original L-chain variable region) means that this region has the same binding affinity when it is combined with the antibody H-chain variable region described herein which forms a pair with the original L chain, or alternatively that the region has the same cytotoxic activity towards glypcan 3-expressing cells or a tissue containing these cells when used for a multispecific antigen-binding molecule.

[0449] The term “equivalent” does not necessarily have to mean the same degree of activity, and the activity may be enhanced. Specifically, for antigen-binding affinity, examples include the case where the value (KD value/parent KD value) obtained by comparison to the binding affinity of the antibody variable region serving as the control (parent KD value) is 1.5 or less. The value of KD value/parent KD value is preferably 1.3 or less, more preferably 1.2 or less, 1.1 or less, 1.0 or less, 0.9 or less, 0.8 or less, 0.7 or less, 0.6 or less, or 0.5 or less. While there is no lower limit, examples include 10^{-1} , 10^{-2} , 10^{-3} , 10^{-4} , 10^{-5} , or 10^{-6} . More specifically, in the present invention, the value of KD value/parent KD value is preferably 10^{-6} to 1.5×10^{-0} , more preferably 10^{-6} to 10^{-1} , even more preferably 10^{-6} to 10^{-2} , and yet even more preferably 10^{-6} to 10^{-3} . For cytotoxic activity, examples include the case where the value (cell growth inhibition rate/parent cell growth inhibition rate) obtained by comparison to the cell growth inhibition rate of the multispecific antigen-binding molecule serving as the control (parent cell growth inhibition rate) is 0.7 or more. The concentration of the added multispecific antigen-binding molecule can be determined appropriately, but is preferably, for example, 0.01 nM, 0.05 nM, 0.1 nM, 0.5 nM, or 1 nM; and preferably, measurements are taken at 0.05 nM or 0.1 nM. The value for cell growth inhibition rate/parent cell growth inhibition rate is preferably 0.8 or higher, more preferably 0.9 or higher, 1.0 or higher, 1.2 or higher, 1.5 or higher, 2 or higher, 3 or higher, 5 or higher, 10 or higher, or 20 or higher. While there is no upper limit, the value may be 10 , 10^2 , 10^3 , 10^4 , 10^5 , or 10^6 .

[0450] Furthermore, for cytotoxic activity, examples include the case where the value (concentration for 50% inhibition of cell growth/parent concentration for 50% inhibition of cell growth) obtained by comparison to the concentration of the original multispecific antigen-binding molecule for 50% inhibition of cell growth (parent concentration for 50% inhibition of cell growth) is 1.5 or less. Concentration for 50% growth inhibition refers to the concentration of the multispecific antigen-binding molecule necessary for reducing the cell proliferation rate to one half compared to when the multispecific antigen-binding molecule is not added. The value of “concentration for 50% inhibition of

cell growth/parent concentration for 50% inhibition of cell growth" is preferably 1.3 or less, more preferably 1.2 or less, 1.1 or less, 1.0 or less, 0.9 or less, 0.8 or less, 0.7 or less, 0.6 or less, or 0.5 or less. While there is no lower limit, the value may be, for example, 10^{-1} , 10^{-2} , 10^{-3} , 10^{-4} , 10^{-5} , or 10^{-6} . Specifically, the value is preferably 10^{-6} to 1.5×10^{-6} , more preferably 10^{-6} to 10^{-1} , even more preferably 10^{-6} to 10^{-2} , and yet even more preferably 10^{-6} to 10^{-3} .

[0451] Regarding the domain comprising an antibody variable region having GPC3-binding activity, the KD value towards GPC3 (for example, human GPC3) may be, for example, 5×10^{-9} M or less, preferably 4×10^{-9} M or less, such as 3×10^{-9} M or less, 2×10^{-9} M or less, 1×10^{-9} M or less, 8×10^{-10} M or less, 5×10^{-10} M or less, 4×10^{-10} M or less, 3×10^{-10} M or less, 2×10^{-10} M or less, 1×10^{-10} M or less, 8×10^{-11} M or less, 5×10^{-11} M or less, 4×10^{-11} M or less, 3×10^{-11} M or less, 2×10^{-11} M or less, 1×10^{-11} M or less, 8×10^{-12} M or less, 5×10^{-12} M or less, 4×10^{-12} M or less, 3×10^{-12} M or less, 2×10^{-12} M or less, 1×10^{-12} M or less, 8×10^{-13} M or less, 5×10^{-13} M or less, 4×10^{-13} M or less, 3×10^{-13} M or less, 2×10^{-13} M or less, or 1×10^{-13} M or less.

[0452] Regarding the domain comprising an antibody variable region having T-cell receptor complex-binding activity, the KD value towards a human T-cell receptor complex such as a human T cell receptor, or more specifically for example human CD38 may be, for example, 2×10^{-7} M or less, preferably 1.5×10^{-7} M or less, such as 1.4×10^{-7} M or less, 1.3×10^{-7} M or less, 1.2×10^{-7} M or less, 1×10^{-7} M or less, 3×10^{-8} M or less, 2×10^{-8} M or less, 1×10^{-8} M or less, 8×10^{-9} M or less, 5×10^{-9} M or less, 4×10^{-9} M or less, 3×10^{-9} M or less, 2×10^{-9} M or less, 1×10^{-9} M or less, 8×10^{-10} M or less, 5×10^{-10} M or less, 4×10^{-10} M or less, 3×10^{-10} M or less, 2×10^{-10} M or less, 1×10^{-10} M or less, 8×10^{-11} M or less, 5×10^{-11} M or less, 4×10^{-11} M or less, 3×10^{-11} M or less, 2×10^{-11} M or less, 1×10^{-11} M or less, 8×10^{-12} M or less, 5×10^{-12} M or less, 4×10^{-12} M or less, 3×10^{-12} M or less, 2×10^{-12} M or less, or 1×10^{-12} M or less.

[0453] The multispecific antigen-binding molecules of the present invention preferably have KD values toward human GPC3 and human T-cell receptor complex (for example, human CD3& chain) that are 5×10^{-9} M or less and 2×10^{-7} M or less, respectively, and more preferably 1×10^{-9} M or less and 5×10^{-8} M or less, respectively.

[0454] In the present invention, antibody variable regions that are "functionally equivalent" are not particularly limited as long as they are antibody H-chain and/or antibody L-chain variable regions that satisfy the above-described conditions. Examples of such antibody variable regions include regions produced by introducing substitution, deletion, addition, and/or insertion of one or more amino acids (for example, 1, 2, 3, 4, 5, or 10 amino acids) into the amino acid sequences of the variable regions of Tables 1 to 3 mentioned above. A method well known to those skilled in the art for introducing one or more amino-acid substitutions, deletions, additions, and/or insertions into an amino acid sequence is a method of introducing mutations into proteins. For example, those skilled in the art can prepare variable regions that are functionally equivalent to the antibody variable regions having the above-mentioned functions by appropriately introducing mutations into amino acid sequences using methods such as site-directed mutagenesis (Hashimoto-Gotoh, T., Mizuno, T., Ogasahara, Y., and Nakagawa, M. (1995) An oligodeoxyribonucleotide-directed

dual amber method for site-directed mutagenesis. Gene 152, 271-275; Zoller, M. J., and Smith, M. (1983) Oligonucleotide-directed mutagenesis of DNA fragments cloned into M13 vectors. Methods Enzymol. 100, 468-500; Kramer, W., Drutsa, V., Jansen, H. W., Kramer, B., Pflugfelder, M., and Fritz, H. J. (1984). The gapped duplex DNA approach to oligonucleotide-directed mutation construction. Nucleic Acids Res. 12, 9441-9456; Kramer, W., and Fritz, H. J. (1987) Oligonucleotide-directed construction of mutations via gapped duplex DNA Methods. Enzymol. 154, 350-367; and Kunkel, T. A. (1985) Rapid and efficient site-specific mutagenesis without phenotypic selection. Proc Natl Acad. Sci. USA. 82, 488-492).

[0455] When an amino acid residue is altered, the amino acid is preferably mutated into a different amino acid(s) that conserves the properties of the amino acid side-chain. Examples of amino-acid side chain properties are: hydrophobic amino acids (A, I, L, M, F, P, W, Y, and V), hydrophilic amino acids (R, D, N, C, E, Q, G, H, K, S, and T), amino acids containing aliphatic side chains (G, A, V, L, I, and P), amino acids containing hydroxyl group-containing side chains (S, T, and Y), amino acids containing sulfur atom-containing side chains (C and M), amino acids containing carboxylic acid- and amide-containing side chains (D, N, E, and Q), amino acids containing basic side chains (R, K, and H), and amino acids containing aromatic side chains (H, F, Y, and W) (amino acids are represented by one-letter codes in parentheses). Amino acid substitutions within each of these groups are called conservative substitutions. It is already known that a polypeptide containing a modified amino acid sequence in which one or more amino acid residues in a given amino acid sequence are deleted, added, and/or substituted with other amino acids can retain the original biological activity (Mark, D. F. et al., Proc. Natl. Acad. Sci. USA; (1984) 81:5662-6; Zoller, M. J. and Smith, M., Nucleic Acids Res. (1982) 10:6487-500; Wang, A. et al., Science (1984) 224:1431-3; Dalbadie-McFarland, G. et al., Proc. Natl. Acad. Sci. USA (1982) 79:6409-13). Variable regions of the present invention containing such amino acid modifications have an amino acid sequence identity of at least 70%, more preferably at least 75%, even more preferably at least 80%, still more preferably at least 85%, yet more preferably at least 90%, and most preferably at least 95%, with the amino acid sequence of the CDR sequences, FR sequences, or whole variable regions of the variable region prior to modification. Herein, sequence identity is defined as the percentage of residues identical to those in the original amino acid sequence of the H-chain variable region or L-chain variable region determined after the sequences are aligned, and gaps are appropriately introduced to maximize the sequence identity as necessary. The identity of amino acid sequences can be determined by the method described below.

[0456] Furthermore, a "functionally equivalent antibody variable region" can be obtained, for example, from nucleic acids that hybridize under stringent conditions with nucleic acids comprising a nucleotide sequence encoding the amino acid sequence of a variable region in Tables 1 to 3 mentioned above. Stringent hybridization conditions for isolating a nucleic acid that hybridizes under stringent conditions with a nucleic acid comprising a nucleotide sequence encoding the amino acid sequence of a variable region include, for example, the conditions of 6 M urea, 0.4% SDS, 0.5×SSC, and 37° C., or hybridization conditions with a stringency

equivalent thereto. Isolation of nucleic acids with a much higher homology can be expected with more stringent conditions, for example, the conditions of 6 M urea, 0.4% SDS, 0.1×SSC, and 42° C. The washing conditions following the hybridization are, for example, washing using 0.5× SSC (1×SSC is 0.15 M NaCl and 0.015 M sodium citrate at pH7.0) and 0.1% SDS at 60° C., more preferably washing using 0.2×SSC and 0.1% SDS at 60° C., even more preferably washing using 0.2×SSC and 0.1% SDS at 62° C., yet even more preferably washing using 0.2×SSC and 0.1% SDS at 65° C., and still more preferably washing using 0.1×SSC and 0.1% SDS at 65° C. The sequences of the isolated nucleic acids can be determined by the known methods described below. The overall nucleotide sequence homology of the isolated nucleic acid is at least 50% or higher, preferably 70% or higher, and more preferably 90% or higher (for example, 95%, 96%, 97%, 98%, 99%, or higher) sequence identity.

[0457] Nucleic acids that hybridize under stringent conditions to a nucleic acid comprising a nucleotide sequence encoding the amino acid sequence of a variable region can also be isolated by using, instead of the above-described methods using hybridization techniques, gene amplification methods such as polymerase chain reaction (PCR) that uses primers synthesized based on information of the nucleotide sequence encoding the variable-region amino acid sequence.

[0458] The identity of one nucleotide sequence or amino acid sequence to another can be determined using the algorithm BLAST, by Karlin and Altschul (Proc. Natl. Acad. Sci. USA (1993) 90:5873-7). Programs called BLASTN and BLASTX were developed based on this algorithm (Altschul et al., J. Mol. Biol. (1990) 215:403-10). To analyze nucleotide sequences according to BLASTN based on BLAST, the parameters are set, for example, as score=100 and wordlength=12. On the other hand, parameters used for the analysis of amino acid sequences by BLASTX based on BLAST include, for example, score=50 and wordlength=3. Default parameters for each program are used when using the BLAST and Gapped BLAST programs. Specific techniques for such analyses are known in the art (see the website of the National Center for Biotechnology Information (NCBI), Basic Local Alignment Search Tool (BLAST); www.ncbi.nlm.nih.gov).

[0459] The combination of the antibody variable region having glycan 3-binding activity and the antibody variable region having T-cell receptor complex binding activity as comprised in the multispecific antigen-binding molecule of the present invention is not particularly limited as long as it has the above-described activities. However, in the present invention, the cytotoxic activity of the multispecific antigen-binding molecule is preferably equivalent to or greater than that of the bispecific antibody GPC3_ERY22_rCE115 described in Example 3. Here, the term "equivalent" does not necessarily have to mean the same degree of activity as described above, and the activity may be enhanced. Being equivalent to GPC3_ERY22_rCE115 is, for example, when the value of (cell growth inhibition rate/cell growth inhibition rate (GPC3_ERY22_rCE115)) relative to the cell growth inhibition rate of GPC3_ERY22_rCE115 (cell growth inhibition rate (GPC3_ERY22_rCE115)) is 0.7 or greater, preferably 0.8 or greater, 0.9 or greater, 1.0 or greater, 1.2 or greater, 1.5 or greater, 2 or greater, 3 or greater, 5 or greater, 10 or greater, or 20 or greater. While there is no upper limit, the value may be, for example, 10,

10², 10³, 10⁴, 10⁵, or 10⁶. The concentration of the multispecific antigen-binding molecule to be added can be determined appropriately, but is preferably, for example, 0.01 nM, 0.05 nM, 0.1 nM, 0.5 nM, or 1 nM; and preferably, measurements are taken at 0.05 nM or 0.1 nM.

[0460] Furthermore, examples include the case where the value (concentration for 50% inhibition of cell growth/concentration for 50% inhibition of cell growth (GPC3_ERY22_rCE115)) obtained by comparison to the concentration for 50% inhibition of growth of GPC3_ERY22_rCE115 cells (concentration for 50% inhibition of cell growth (GPC3_ERY22_rCE115)) is 1.5 or less. The value for "concentration for 50% inhibition of cell growth/concentration for 50% inhibition of cell growth (GPC3_ERY22_rCE115)" is preferably 1.3 or less, more preferably 1.2 or less, 1.1 or less, 1.0 or less, 0.9 or less, 0.8 or less, 0.7 or less, 0.6 or less, or 0.5 or less. While there is no lower limit, the value may be for example, 10⁻¹, 10⁻², 10⁻³, 10⁻⁴, 10⁻⁵, or 10⁻⁶. Specifically, the value is preferably 10⁶ to 1.5×10⁶, more preferably 10⁻⁶ to 10⁻¹, even more preferably 10⁶ to 10⁻², and yet even more preferably 10⁻⁶ to 10⁻³.

[0461] The preferred specific KD values for human GPC3 and human T cell receptor complex (for example, human CD38 chain) are also as indicated above. Desired cells showing GPC3 expression or desired tissues containing these cells may be used for the cells, and for example, PC-10 or NCI-H446 which are GPC3-expressing human cancer cell lines can be used.

[0462] Examples of such a combination of the antibody variable region having glycan 3-binding activity and the antibody variable region having T-cell receptor complex binding activity, include the combinations of antibody H-chain variable regions shown in Table 4, combinations of antibody H-chain variable regions having CDR sequences whose CDR1, CDR2, and CDR3 amino acid sequences are the same as the CDR1, CDR2, and CDR3 amino acid sequences carried by the antibody H-chain variable regions of Table 4, and combinations of antibody H-chain variable regions functionally equivalent to these variable regions. Here, "functionally equivalent" has the same meaning described above.

TABLE 4

GPC3 side/T cell receptor complex side	SEQ ID NO:
H0000/hCE115HA	40/52
H0000/CE115HA251	40/500
H0000/CE115HA236	40/429
H0000/TR01H002	40/421
H0000/CE115HA122	40/426
H0610/rCE115H	215/424
H0610/TR01H040	215/103
H0610/TR01H061	215/122
H0610/TR01H068	215/129
H0610/TR01H071	215/132
GCH054/TR01H067	197/128
GCH094/TR01H082	211/142
GCH094/TR01H084	211/144
GCH065/TR01H084	206/144
GCH065/TR01H082	206/142
GCH094/TR01H109	211/164
GCH065/TR01H109	206/164
GCH094/TR01H113	211/168
GCH065/TR01H113	206/168

[0463] A preferred common L chain for such combinations of an antibody variable region having glycan 3-binding activity and an antibody variable region having T-cell receptor complex binding activity includes, for example, L0000, L0011, L0201, L0203, L0204, L0206, L0208, L0209, L0211, L0212, L0222, and a common L chain having CDR sequences (CDR1, CDR2, and CDR3 amino acid sequences) identical to the CDR1, CDR2, and CDR3 amino acid sequences as in the above common L chain. Specific combinations include, for example, the combinations of antibody H-chain variable regions and a common L chain shown in Table 5, combinations of antibody variable regions having CDR sequences (CDR1, CDR2, and CDR3 amino acid sequences) identical to the amino acid sequences of CDR1, CDR2, and CDR3 carried by the antibody variable regions and a common L chain of Table 5, and combinations of antibody H-chain variable regions and a common L chain functionally equivalent to these variable regions. Here, “functionally equivalent” has the same meaning as described above.

TABLE 5

GPC3 side/T cell receptor complex side/common L chain	SEQ ID NO:
H0610/rCE115H/L0000	215/424/53
H0610/TR01H040/L0000	215/103/53
H0610/TR01H040/L0201	215/103/299
H0610/TR01H040/L0203	215/103/301
H0610/TR01H040/L0204	215/103/302
H0610/TR01H040/L0206	215/103/304
H0610/TR01H040/L0208	215/103/306
H0610/TR01H040/L0209	215/103/307
H0610/TR01H040/L0211	215/103/309
H0610/TR01H061/L0000	215/122/53
H0610/TR01H068/L0000	215/129/53
H0610/TR01H071/L0000	215/132/53
GCH054/TR01H067/L0201	197/128/299
GCH054/TR01H067/L0212	197/128/310
GCH054/TR01H067/L0222	197/128/319
GCH054/TR01H067/L0000	197/128/53
GCH094/TR01H082/L0201	211/142/299
GCH094/TR01H082/L0011	211/142/223
GCH094/TR01H084/L0011	211/144/223
GCH065/TR01H084/L0011	206/144/223
GCH065/TR01H082/L0011	206/142/223
GCH094/TR01H109/L0011	211/164/223
GCH065/TR01H109/L0011	206/164/223
GCH094/TR01H113/L0011	211/168/223
GCH065/TR01H113/L0011	206/168/223

[0464] The Fc region comprised in the multispecific antigen-binding molecule of the present invention is not particularly limited as long as it is an Fc region having reduced Fc γ receptor-binding activity, but examples of a preferred Fc region of the present invention include a combination of the Fc-region portion of E22Hh and the Fc-region portion of E22Hk, a combination of the Fc-region portion of E2702GsKsc and the Fc-region portion of E2704sEpsc, and a combination of the Fc-region portion of E2702sKsc and the Fc-region portion of E2704sEpsc.

[0465] Examples of a preferred multispecific antigen-binding molecule of the present invention include bispecific antibodies comprising an antibody variable region having glycan 3-binding activity and an antibody variable region having CD38-binding activity. More preferably, the cytotoxic activity is the same or greater than that of the GPC3₃-ERY22_rCE115 bispecific antibody. Examples of such bispecific antibodies include bispecific antibodies comprising

H and L chains described in Table 13, and bispecific antibodies that bind to an epitope overlapping with an epitope bound by the above antibodies, and which contain an Fc region with reduced Fc γ receptor-binding activity.

[0466] Whether an antibody recognizes an epitope that overlaps with an epitope recognized by another antibody can be confirmed by the competition between the two antibodies against the epitope. Competition between the antibodies can be evaluated by competitive binding assays using means such as enzyme-linked immunosorbent assay (ELISA), fluorescence energy transfer method (FRET), and fluorometric microvolume assay technology (FMAT (Registered trademark)). The amount of an antibody bound to an antigen indirectly correlates with the binding ability of a candidate competitor antibody (a test antibody) that competitively binds to the overlapping epitope. In other words, as the amount or affinity of a test antibody against the overlapping epitope increases, the amount of the antibody bound to the antigen decreases, and the amount of the antigen-bound test antibody increases. Specifically, the appropriately labeled antibody and antibody to be evaluated are simultaneously added to the antigen, and the antibody bound as a result are detected using the label. The amount of the antigen-bound antibody can be easily determined by labeling the antibody beforehand. This label is not particularly limited, and the labeling method is selected according to the assay technique used. Specifically, the labeling method includes fluorescent labeling, radiolabeling, enzymatic labeling, and such.

[0467] For example, the fluorescently labeled antibody and the unlabeled antibody or test antibody are simultaneously added to beads immobilized with GPC3 or CD38, and the labeled antibody is detected by fluorometric microvolume assay technology.

[0468] Herein, the “antibody that binds to the overlapping epitope” refers to a test antibody that can reduce the amount of the bound labeled antibody by at least 50% at a concentration that is usually 100 times higher, preferably 80 times higher, more preferably 50 times higher, even more preferably 30 times higher, and still more preferably 10 times higher than the concentration at which the non-labeled antibody reduces 50% of the amount of the labeled antibody bound (IC50).

[0469] Multispecific antigen-binding molecules, which have the antigen-binding sites of antibodies that bind to epitopes overlapping with epitopes bound by the above-mentioned antibodies, can yield excellent cytotoxic activity.

[0470] The multispecific antigen-binding molecules of the present invention are produced by the same technique as the method for producing recombinant antibodies mentioned above.

[0471] The present invention also relates to polynucleotides encoding the antigen-binding molecules of the present invention, and they can be inserted into discretionary expression vectors. Suitable hosts can be transformed with the expression vectors to produce cells that express the antigen-binding molecules. Antigen-binding molecules encoded by the polynucleotides can be obtained by culturing the cells that express the antigen-binding molecules, and collecting the expression products from culture supernatants. That is, the present invention relates to vectors comprising a polynucleotide encoding an antigen-binding molecule of the present invention, cells carrying such a vector, and methods for producing antigen-binding molecules, which comprise culturing the cells and collecting antigen-

binding molecules from culture supernatants. These can be obtained by techniques similar to those for recombinant antibodies mentioned above.

Pharmaceutical Compositions

[0472] From another viewpoint, the present invention provides pharmaceutical compositions comprising as the active ingredient a multispecific antigen-binding molecule that comprises: (1) a domain comprising an antibody variable region having glypican 3-binding activity, (2) a domain comprising an antibody variable region having T-cell receptor complex-binding activity, and (3) a domain comprising an Fc region with reduced binding activity towards an Fcγ receptor. Furthermore, the present invention relates to pharmaceutical compositions that induce cell injury, which comprise the antigen-binding molecule as an active ingredient. Pharmaceutical compositions of the present invention which induce the described cell injury, particularly T-cell-dependent cellular cytotoxicity, are preferably administered to a subject suffering from a disease for which the activities are needed for prevention or treatment, or a subject in which the disease is possible to relapse.

[0473] Furthermore, in the present invention, cytotoxicity-inducing agents and cell growth-inhibiting agents comprising as the active ingredient a multispecific antigen-binding molecule that comprises:

[0474] (1) a domain comprising an antibody variable region having glypican 3-binding activity,

[0475] (2) a domain comprising an antibody variable region having T-cell receptor complex-binding activity, and

[0476] (3) a domain comprising an Fc region with reduced binding activity towards an Fcγ receptor

[0477] can be presented as a method for inducing cell injury which comprises the step of administering the antigen-binding molecule to a subject, or it can be presented as use of the antigen-binding molecule in the manufacture of a cytotoxicity-inducing agent and a cell growth-inhibiting agent.

[0478] In the present invention “comprising as the active ingredient a multispecific antigen-binding molecule that comprises (1) a domain comprising an antibody variable region having glypican 3-binding activity, (2) a domain comprising an antibody variable region having T-cell receptor complex-binding activity, and (3) a domain comprising an Fc region with reduced binding activity towards an Fcγ receptor” means comprising the antigen-binding molecule as a major active component, without limitation to the content ratio of the antigen-binding molecule.

[0479] If necessary, multispecific antigen-binding molecules of the present invention may be encapsulated in microcapsules (e.g., those made of hydroxymethylcellulose, gelatin, and poly(methylmethacrylate)), or incorporated as components of a colloidal drug delivery system (e.g., liposomes, albumin microspheres, microemulsion, nanoparticles, and nanocapsules) (see, for example, “Remington’s Pharmaceutical Science 16th edition”, Oslo Ed. (1980)). Methods for preparing the pharmaceutical agents as controlled-release pharmaceutical agents are also well known, and such methods may be applied to the multispecific antigen-binding molecules of the present invention (J. Biomed. Mater. Res. (1981) 15:267-277; Chemtech. (1982)

12:98-105; U.S. Pat. No. 3,773,719; European Patent Application Publication Nos. EP 58,481 and EP 133,988; Biopolymers (1983) 22:547-556).

[0480] The pharmaceutical compositions of the present invention or cytotoxicity-inducing agents and cell growth-inhibiting agents may be administered to patients by oral or parenteral administration, and parenteral administration is preferred. Specific examples of the administration method include administration by injection, transnasal administration, transpulmonary administration, and transdermal administration. Examples of administration by injection include intravenous injection, intramuscular injection, intra-peritoneal injection, and subcutaneous injection. A pharmaceutical composition of the present invention or a cytotoxicity-inducing agent and a cell growth-inhibiting agent can be administered systemically or locally, for example, through administration by injection. The method of administration can be selected appropriately according to the age and symptoms of the patient. The dose can be selected from the range of 0.0001 mg to 1000 mg per kilogram body weight for a single administration. Alternatively, for example, the dose may be selected from the range of 0.001 mg/body to 100000 mg/body per patient. However, the pharmaceutical compositions of the present invention or a cytotoxicity-inducing agent and cell growth-inhibiting agent are not limited to these doses.

[0481] The pharmaceutical compositions of the present invention or cytotoxicity-inducing agents and cell growth-inhibiting agents can be formulated according to conventional methods (for example, Remington’s Pharmaceutical Science, latest edition, Mark Publishing Company, Easton, U.S.A.) and may also contain pharmaceutically acceptable carriers and additives. Examples include, but are not limited to surfactants, excipients, coloring agents, perfumes, preservatives, stabilizers, buffers, suspending agents, isotonicization agents, binders, disintegrants, lubricants, fluidity promoting agents, and flavoring agents; and other commonly used carriers can be suitably used. Specific examples of the carriers include light anhydrous silicic acid, lactose, crystalline cellulose, mannitol, starch, carmellose calcium, carmellose sodium, hydroxypropyl cellulose, hydroxypropyl methyl cellulose, polyvinylacetal diethylaminoacetate, polyvinylpyrrolidone, gelatin, medium chain fatty acid triglyceride, polyoxyethylene hardened castor oil 60, saccharose, carboxymethyl cellulose, corn starch, inorganic salt, and such.

[0482] The present invention also provides methods for damaging glypican 3 antigen-expressing cells or tumor tissues containing the antigen-expressing cells, or methods for suppressing growth of these cells or tumor tissues by contacting the glypican 3 antigen-expressing cells with a multispecific antigen-binding molecule of the present invention that binds to the antigen. The multispecific antigen-binding molecule that binds to the antigen is as described above for an antigen-binding molecule of the present invention that binds to the antigen, which is comprised in the cytotoxicity-inducing agents and cell growth-inhibiting agents of the present invention. The cells bound by a multispecific antigen-binding molecule of the present invention that binds to the antigen are not particularly limited as long as they are cells expressing the antigen.

[0483] In the present invention, “contact” is carried out, for example, by adding a multispecific antigen-binding molecule of the present invention which binds to the antigen

to the culture medium of GPC3 antigen-expressing cells cultured in vitro. In this case, a liquid or a solid obtained by freeze-drying or such may be suitably used as the form of the added antigen-binding molecule. When added as an aqueous solution, it may be an aqueous solution that simply contains only the multispecific antigen-binding molecule of the present invention, or it may be a solution containing also, for example, the above-mentioned surfactants, excipients, coloring agents, perfumes, preservatives, stabilizers, buffers, suspending agents, isotonization agents, binders, disintegrants, lubricants, fluidity promoting agents, and flavoring agents. The concentration at which the addition is performed is not particularly limited, but a suitable final concentration in the culture solution is preferably in the range of 1 µg/ml to 1 g/ml, more preferably 1 ng/ml to 1 mg/ml, and even more preferably 1 µg/mL to 1 mg/mL.

[0484] Furthermore, in another embodiment, "contact" of the present invention is also carried out by administering an antigen-binding molecule of the present invention to non-human animals with cells expressing the GPC3 antigen transplanted into their bodies, and to animals carrying cells that intrinsically express the antigen. The method of administration may be oral or parenteral, and parenteral administration is particularly preferred. Specific examples of the administration method include administration by injection, transnasal administration, transpulmonary administration, and transdermal administration. Examples of administration by injection include intravenous injection, intramuscular injection, intraperitoneal injection, and subcutaneous injection. A pharmaceutical composition of the present invention or a cytotoxicity-inducing agent and a cell growth-inhibiting agent can be administered systemically or locally, for example, through administration by injection. The method of administration can be selected appropriately according to the age and symptoms of the test animal. When administered as an aqueous solution, an aqueous solution containing simply only a multispecific antigen-binding molecule of the present invention may be used, or a solution containing also the above-mentioned surfactants, excipients, coloring agents, perfumes, preservatives, stabilizers, buffers, suspending agents, isotonization agents, binders, disintegrants, lubricants, fluidity promoting agents, flavoring agents, and such may be used. The dose can be selected from the range of 0.0001 mg to 1000 mg per kilogram body weight for a single administration.

[0485] Alternatively, for example, the dose may be selected from the range of 0.001 mg/body to 100000 mg/body per patient. However, the amount of the multispecific antigen-binding molecule of the present invention administered is not limited to these doses.

[0486] The following method is suitably used as a method for evaluating or measuring cell injury induced in cells expressing the glycan 3 antigen which is bound by a domain carrying an antibody variable region having glycan 3-binding activity that constitutes the antigen-binding molecule as a result of contacting the cells with a multispecific antigen-binding molecule of the present invention. Examples of a method for evaluating or measuring the cytotoxic activity in vitro include methods for measuring cytotoxic T cell activity and such. Whether or not a multispecific antigen-binding molecule of the present invention has T cellular cytotoxicity can be measured by known methods (for example, Current protocols in Immunology, Chapter 7. Immunologic studies in humans, Editor, John E.

Coligan et al., John Wiley & Sons, Inc., (1993) and the like). For activity measurements, an antigen-binding molecule that binds to an antigen different from glycan 3, which is an antigen not expressed in the cells used for the examination, can be used as a control in the same manner as a multispecific antigen-binding molecule of the present invention, and the activity can be determined to be present when the multispecific antigen-binding molecule of the present invention shows a stronger cytotoxic activity than when the antigen-binding molecule is used as a control.

[0487] To evaluate or measure cytotoxic activity in vivo, for example, cells expressing a glycan 3 antigen are intradermally or subcutaneously transplanted to a non-human test animal, and then a test antigen-binding molecule is intravenously or intraperitoneally administered daily or with an interval of few days, starting from the day of transplantation or the following day. Cytotoxic activity can be determined by daily measurement of tumor size and by observing difference in the change of tumor size. In a similar manner to the in vitro evaluation, the cytotoxic activity of an antigen-binding molecule of the present invention can be determined to be present when administration of a control antigen-binding molecule shows that the tumor size in the group subjected to administration of an antigen-binding molecule of the present invention is significantly smaller than the tumor size in the group subjected to administration of the control antigen-binding molecule.

[0488] As a method for evaluating or measuring the suppressive effect on proliferation of cells expressing a glycan 3 antigen, a method of measuring the uptake of isotope-labeled thymidine into cells or the MTT method may be suitably used. As a method for evaluating or measuring the cell proliferation-suppressing activity in vivo, the same method described above for evaluating or measuring cytotoxic activity in vivo may be suitably used.

[0489] The present invention also provides kits for use in the methods of the present invention, which comprise a multispecific antigen-binding molecule of the present invention or a multispecific antigen-binding molecule produced by a production method of the present invention. Additionally, the kit may include in its package, a pharmaceutically acceptable carrier, solvent, and instructions describing the method of use.

[0490] The present invention also relates to a multispecific antigen-binding molecule of the present invention or a multispecific antigen-binding molecule produced by a production method of the present invention for use in a method of the present invention.

[0491] The present invention also relates to molecules having GPC3-binding activity, which contain a domain comprising an antibody variable region having GPC3-binding activity of the multispecific antigen-binding molecule of the present invention. Furthermore, the present invention relates to a molecule having GPC3-binding activity, which comprises the antibody variable regions of H and L chains respectively comprising the three CDRs of the H and L chains (total of six CDRs) contained in the molecule. The present invention also relates to molecules having T-cell receptor complex-binding activity, which contain a domain comprising an antibody variable region having T-cell receptor complex-binding activity of the multispecific antigen-binding molecule of the present invention. Furthermore, the present invention relates to a molecule having T-cell receptor complex-binding activity that comprises the antibody

variable regions of the H and L chains respectively comprising the three CDRs of the H and L chains (total of six CDRs) contained in the molecule. Such molecules may be antibodies or polypeptides comprising antigen-binding fragments of an antibody. The present invention also relates to antibodies that bind to epitopes overlapping or competing with these molecules or polypeptides containing the antigen-binding fragments thereof. Suitable examples of such polypeptides comprising antigen-binding fragments of an antibody include scFv, single chain antibody, Fv, single chain Fv 2 (scFv2), Fab, and F(ab')2. Furthermore, these molecules do not have to be multispecific (bispecific), and may bind only to either GPC3 or a T cell receptor complex (for example, the CD38 chain).

[0492] These molecules include a molecule comprising a domain that comprises an antibody variable region having GPC-binding activity of the multispecific antigen-binding molecule exemplified in detail in the Examples herein (which comprises the H-chain variable regions having GPC3-binding activity and the common L-chain variable region), a molecule comprising a domain that comprises an antibody variable region having T cell receptor complex-binding activity of the multispecific antigen-binding molecule exemplified in the Examples herein (which comprises the H-chain variable regions having T cell receptor complex-binding activity and the common L-chain variable region), and also a molecule having an activity to bind to the same antigenic protein (GPC3 or T-cell receptor complex), which comprises the three CDRs of each of the H and L chains (total of six CDRs) contained in the above molecule.

[0493] These molecules have CDRs that are in common with those of a multispecific antigen-binding molecule of the present invention; and therefore, they are expected to bind to an epitope overlapping with an epitope for the multispecific antigen-binding molecule of the present invention. Therefore, these molecules can compete with multispecific antigen-binding molecules of the present invention when they coexist with the multispecific antigen-binding molecules of the present invention. Accordingly, these molecules can be used, for example, as regulatory agents for suppressing activities (such as antigen-binding activity, cytotoxic activity, and antitumor activity) of the multispecific antigen-binding molecules of the present invention. Furthermore, such a molecule can be bound to a target protein (GPC3 or T cell receptor complex) in advance, and when a multispecific antigen-binding molecule of the present invention is added, the molecules that dissociate through competition can be detected. This way, the molecule is useful as an agent for detecting binding of a multispecific antigen-binding molecule of the present invention to a target protein. Here, such molecules may be labeled appropriately with fluorescent substances or such. Alternatively, these molecules are useful for screening novel antibodies that bind to epitopes overlapping with the epitopes bound by the multispecific antigen-binding molecules of the present invention. As described above, such a molecule can be bound to a target protein (GPC3 or T cell receptor complex) in advance, and when a test antibody is added, if the bound molecules dissociate, then the test antibody is a candidate for an antibody that binds to an epitope overlapping with the epitope bound by the multispecific antigen-binding molecule of the present invention. This will enable efficient screening of novel multispecific antigen-binding molecules.

[0494] The combinations presented as examples herein as combinations of each CDR of the multispecific antigen-binding molecules of the present invention can be directly used as specific combinations of CDRs of the H-chain and L-chain variable regions in these molecules. The antigen affinity of these molecules (KD values) is preferably a value exemplified herein as the KD value of a multispecific antigen-binding molecule of the present invention, but is not limited thereto.

[0495] The present invention also relates to nucleic acids encoding these molecules, vectors comprising the nucleic acids, cells comprising the nucleic acids or the vectors, methods for producing the molecules by culturing the cells, and molecules produced by these methods.

[0496] All prior art references cited herein are incorporated by reference into this description.

EXAMPLES

[0497] Herein below, the present invention will be specifically described with reference to the Examples, but it is not to be construed as being limited thereto.

[Example 1] Production of GPC3_ERY22_rCE115 and Measurement of Cytotoxic Activity

(1-1) Production of GPC3_ERY22_rCE115

[0498] A molecule in which one of the Fabs has been replaced with a CD3 epsilon-binding domain was produced using IgG against a cancer antigen (GPC3) as the basic structure. In this case, the IgG Fc used as the basic structure was a silent Fc with attenuated affinity for FcgR (an Fcy (Fc gamma) receptor). An anti-GPC3 antibody, H0000 (SEQ ID NO: 40)/GL4 (SEQ ID NO:41), was used as the GPC3-binding domain. An anti-CD3 antibody, rCE115H/rCE115L (SEQ ID NO:42/SEQ ID NO:43), was used as the CD3-binding domain.

[0499] G1d produced by removing Gly and Lys at the C terminus of IgG1 was used as the antibody H-chain constant region, and this was used in combination with H0000/GL4 and rCE115H/rCE115L. When the antibody H-chain constant region was named H1, the sequence corresponding to the H chain of the antibody carrying H0000 in the variable region was shown as H0000-H1. Here, an amino acid alteration was shown, for example, as D356K. The first alphabet (corresponding to D in D356K) is the one-letter code representation for the amino acid residue before modification, the number that follows (corresponding to 356 of D356K) is the position of modification indicated by EU numbering, and the final alphabet (corresponding to K of D356K) is the one-letter code representation for the amino acid residue after modification. Gldh (SEQ ID NO:44) produced by removing Gly and Lys at the C terminus of IgG1, ERY22_Hk (SEQ ID NO:45) produced by introducing the L234A/L235A/Y349C/T366W mutations into Gldh, and ERY22_Hh (SEQ ID NO:46) produced by introducing the L234A/L235A/D356C/T366S/L368A/Y407V mutations into G1dh were prepared according to the method of Reference Example 1. The L234A and L235A mutations were introduced into the respective H chains to attenuate affinity for FcgR (an Fcy receptor), and the Y349C/T366W and D356C/T366S/L368A/Y407V mutations were introduced to efficiently form heteromers of each H chain when producing heterodimeric antibodies comprising two types of H chains.

[0500] The heterodimeric antibody, GPC3_ERY22_rCE115, produced by substitution with the VH and VL domains of Fab against GPC3 was prepared according to Reference Example 1 (FIG. 1A).

[0501] A series of expression vectors inserted with a polynucleotide encoding each of GL4-ERY22_Hk (SEQ ID NO:47), H0000-ERY22_L (SEQ ID NO:48), rCE115H-ERY22_Hh (SEQ ID NO:49), and rCE115L-k0 (SEQ ID NO:50) were produced by methods well-known to those skilled in the art, such as PCR methods using primers added with an appropriate sequence similar to those in the above-described method.

[0502] The following combination of expression vectors were introduced into FreeStyle 293-F cells for transient expression of each target molecule.

[0503] Target molecule: GPC3_ERY22_rCE115

[0504] Polypeptides encoded by the polynucleotides inserted into the expression vectors: GL4-ERY22_Hk, H0000-ERY22_L, rCE115H-ERY22_Hh, rCE115L-k0

(1-2) Purification of GPC3_ERY22_rCE115

[0505] The obtained culture supernatant was added to an anti-FLAG M2 column (Sigma), and then the column was washed, followed by elution using 0.1 mg/mL of a FLAG peptide (Sigma). The fractions containing the molecule of interest were added to a HisTrap HP column (GE Healthcare), and then the column was washed, followed by elution using an imidazole concentration gradient. Fractions containing the molecule of interest were concentrated using an ultrafiltration membrane, then the fractions were added to a Superdex 200 column (GE Healthcare), and each of the purified molecules of interest was obtained by collecting only the monomeric fractions from the eluted solution.

(1-3) Measurement of the Cytotoxic Activity of GPC3_ERY22_rCE115 Using Human Peripheral Blood Mononuclear Cells

[0506] The in vitro cytotoxic activity of GPC3_ERY22_rCE115 was assessed.

(1-3-1) Preparation of a Human Peripheral Blood Mononuclear Cell (PBMC) Solution

[0507] Using a syringe preloaded with 100 µL of 1,000 units/mL heparin solution (Novo Heparin for injection, 5000 units, Novo Nordisk), 50 mL of peripheral blood was collected from each healthy volunteer (adult individual). This peripheral blood was diluted two-fold in PBS (-), divided into four aliquots, and added into a Leucosep tube for lymphocyte separation (Cat. No. 227290, Greiner Bio-One) that had been loaded with 15 mL of Ficoll-Paque PLUS and subjected to centrifugation in advance. This separation tube was centrifuged (at 2150 rpm for ten minutes at room temperature), and then the mononuclear cell fraction was collected. The cells in the mononuclear cell fraction were washed once with the Dulbecco's Modified Eagle's Medium containing 10% FBS (manufactured by SIGMA, hereinafter referred to as 10% FBS/D-MEM), and then prepared to have a cell density of 4×10^6 cells/mL using 10% FBS/D-MEM. The cell suspension prepared this way was used as the human PBMC solution in the experiment below.

(1-3-2) Measurement of Cytotoxic Activity

[0508] Cytotoxic activity was assessed by the rate of cell growth inhibition using the xCELLigence Real-Time Cell Analyzer (Roche Diagnostics). The NCI-H446 human cancer cell line or the PC-10 human cancer cell line, which expresses human GPC3, was used as the target cell. NCI-H446 or PC-10 was detached from the dish, then the cells were plated into E-Plate 96 (Roche Diagnostics) in aliquots of 100 µL/well by adjusting the cells to 1×10^4 cells/well, and measurement of live cells was begun using the xCELLigence Real-Time Cell Analyzer. On the following day, the plate was removed from the xCELLigence Real-Time Cell Analyzer, and 50 µL of the respective antibodies prepared at each concentration (0.004, 0.04, 0.4, 4, or 40 nM) were added to the plate. After 15 minutes of reaction at room temperature, 50 µL of the human PBMC solution prepared in (1-2) was added (2×10^5 cells/well), and measurement of live cells was begun by setting the plate into the xCELLigence Real-Time Cell Analyzer again. The reaction was carried out under the conditions of 5% carbon dioxide gas at 37°C., and from the Cell Index value obtained 72 hours after addition of the human PBMC, the cell growth inhibition rate (%) was determined using the equation below. The Cell Index value used in the calculation was a normalized value where the Cell Index value immediately before antibody addition was defined as 1.

$$\text{Cell growth inhibition rate (\%)} = \frac{(A-B)}{A} \times 100 / (A-1)$$

[0509] A represents the mean value of the Cell Index values in wells without antibody addition (containing only the target cells and human PBMCs), and B represents the mean value of the Cell Index values in each well. The examinations were performed in triplicate.

[0510] When peripheral blood mononuclear cells (PBMCs) prepared from human blood were used as the effector cell to measure the cytotoxicity of GPC3_ERY22_rCE115, a very strong activity was observed (FIG. 2).

[Example 2] Humanization of the H Chain of the Anti-CD3 Antibody, rCE115, and Sharing of a Common L Chain

(2-1) Design of hCE115HA, the Humanized rCE115 H-Chain Variable Region

[0511] The H-chain variable region of the rCE115 anti-CD3 antibody (SEQ ID NO:42) was humanized. CDR and FR were determined as defined by Kabat (Kabat numbering).

[0512] First, a human FR sequence was selected by comparing the human antibody variable region sequences in a database to the rCE115 rat variable region sequence. The IMGT Database (www.imgt.org/) and NCBI GenBank (www.ncbi.nlm.nih.gov/genbank/) were used for the database. A humanized H-chain variable region sequence was designed by linking the H-chain CDR sequence of the rCE115 variable region with the selected human FR sequence. This yielded a humanized H-chain variable region sequence, hCE115HL (SEQ ID NO: 51).

[0513] The amino acid residue at position 93 indicated by Kabat numbering is Ala in the selected human H-chain FR3 sequence, but is Arg in the rCE115 variable region sequence. Using the database of rat and human germline sequences (IMGT Database (www.imgt.org/)), only few sequences were found to contain Arg at this site. It is reported that the

amino acid residue at position 94 indicated by Kabat numbering contributes to stabilization of the antibody structure by upper core formation (Ewert et al. Methods. 2004 October; 34 (2): 184-99). Based on such information, a humanized H-chain variable region sequence, in which the amino acid residues at Kabat positions 93 and 94 in the H-chain FR3 were substituted with those residues present in the rCE115 variable region sequence, was newly designed. This was the humanized H-chain variable region sequence, hCE115HA (SEQ ID NO:52).

(2-2) Design of the Common L Chain, L0000, for the rCE115 Anti-CD3 Antibody and the Anti-GPC3 Antibody [0514] The FR/CDR shuffling of the L-chain variable region rCE115L (SEQ ID NO:43) of the rCE115 anti-CD3 antibody and the L-chain variable region GL4 (SEQ ID NO:41) of the anti-GPC3 antibody was performed.

[0515] The FR sequence of GL4 was selected as the L-chain FR sequence. L-chain CDR2 was the same for rCE115L and GL4. The L-chain CDR1 was selected from the CDR sequences of GLA, and the L-chain CDR3 was selected from the CDR sequences of rCE115L, respectively. Furthermore, the L-chain CDR3 produced by substituting the amino acid residue Asp at Kabat position 94 of the selected L-chain CDR3 with the Val residue present in GL4 was newly designed.

[0516] A humanized L chain variable region sequence was designed by linking FR and CDR selected above. This yielded a humanized L-chain variable region sequence, L0000 (SEQ ID NO:53).

(2-3) Evaluation of the Affinity for Human GPC3

[0517] The activity to bind human GPC3 when using GL4 (SEQ ID NO:41) and L0000 (SEQ ID NO:53) as the L-chain variable regions was evaluated. This was performed using the molecular form of a single-arm antibody having a single Fab at the Fc region of a human IgG1 heterodimerized by the knobs-into-hole technique. H0000 (SEQ ID NO:40) was used for the anti-GPC3 antibody H-chain variable region.

[0518] The affinity and binding rate constants of an anti-GPC3 antibody for an antigen were measured by the multi-cycle kinetics method of a surface plasmon resonance assay using Biacore™-T200 (GE Healthcare Japan). HBS-EP+ (GE Healthcare Japan) was used for the running buffer, and an amine coupling kit (GE Healthcare Japan) was used to covalently bind Protein A/G to the CM5 chip (carboxymethyl dextran-coated chip). Each anti-GPC3 antibody was prepared so that approximately 100 RU will be captured by Protein A/G. Human GPC3 used as the analyte was prepared at 8, 16, 32, 64, and 128 nM using HBS-EP+. Measurements were carried out by first allowing Protein A/G to capture the antibody solution, and then injecting the human GPC3 solution at a flow rate of 30 µL/min for three minutes to allow reaction to take place. Then, the solution was switched to HBS-EP+ and the dissociation phase was measured for 15 minutes. After completion of the dissociation phase measurement, the sensor chip was regenerated by washing with 10 mM Gly-HCl at pH 1.5. Measurement at the concentration of 0 was similarly carried out by allowing Protein A/G to capture the antibody solution, performing a three-minute HBS-EP+ injection to allow reaction to take place, and then switching to HBS-EP+ to measure the dissociation phase for 15 minutes. After completion of the dissociation phase measurement, the sensor chip was regenerated by washing with 10 mM Gly-HCl at pH 1.5. A data analysis software

exclusively for Biacore, Biacore T200 Evaluation Software Version 1.0, was used to perform kinetic analyses to calculate the binding rate constant (ka), dissociation rate constant (kd), and the rate constant ratio from the obtained sensorgrams. The results are shown in Table 6.

TABLE 6

Variable region		Affinity for human GPC3		
H-chain variable region	L-chain variable region	KD (M)	ka (1/Ms)	kd (1/s)
H0000	GL4	4.2×10^{-9}	4.3×10^5	1.8×10^{-3}
H0000	L0000	3.6×10^{-8}	3.0×10^5	1.1×10^{-2}

(2-4) Evaluation of the Affinity for Human CD3

[0519] The activity to bind human CD3 when using hCE115HA (SEQ ID NO:52) as the H chain variable region and L0000 (SEQ ID NO:53) as the L-chain variable region was evaluated. This was performed using the molecular form of a single-arm antibody having a single Fab at the Fc region of a human IgG1 heterodimerized by the knobs-into-hole technique.

[0520] The affinity and binding rate constants of an anti-CD3 antibody for an antigen were measured by the single-cycle kinetics method of a surface plasmon resonance assay using Biacore™-T200 (GE Healthcare Japan). HBS-EP+ (GE Healthcare Japan) was used for the running buffer, and an amine coupling kit (GE Healthcare Japan) was used to covalently bind human CD3 to the CM4 chip (carboxymethyl dextran-coated chip). The anti-CD3 antibody used as the analyte was prepared at 5 and 20 µg/mL using HBS-EP+. Measurements were carried out by first injecting each of the 5- and 20-µg/mL anti-CD3 antibody solutions for three minutes continuously at a flow rate of 20 L/min to allow reaction to take place. Then, the solution was switched to HBS-EP+ and the dissociation phase was measured for 3 minutes. After completion of the dissociation phase measurement, the sensor chip was regenerated by washing with 10 mM Gly-HCl at pH 1.5. Measurement at the concentration of 0 was carried out by performing each of the three-minute HBS-EP+ injections twice successively to allow reaction to take place, and then switching to HBS-EP+ to measure the dissociation phase for 3 minutes. After completion of the dissociation phase measurement, the sensor chip was regenerated by washing with 10 mM Gly-HCl at pH 1.5. A data analysis software exclusively for Biacore, Biacore T200 Evaluation Software Version 1.0, was used to perform kinetic analyses to calculate the binding rate constant (ka), dissociation rate constant (kd), and the rate constant ratio from the obtained sensorgrams. The results are shown in Table 7.

TABLE 7

Variable region		Affinity for human CD3		
H-chain variable region	L-chain variable region	KD (M)	ka (1/Ms)	kd (1/s)
rCE115H	rCE115L	1.0×10^{-7}	5.9×10^4	6.0×10^{-3}
hCE115HA	L0000	1.2×10^{-7}	1.9×10^5	2.3×10^{-2}

(2-5) Preparation of GPC3_ERY27_hCE115

[0521] The IgG4 against a cancer antigen (GPC3) was used as the basic structure to produce the ERY27 molecule (FIG. 1B), in which the H-chain variable region of one of the Fabs has been replaced with a CD3 epsilon-binding domain, and the L chain is common to both Fabs. In this case, the IgG4 Fc used as the basic structure was a silent Fc with attenuated affinity for FcgR (an Fcy receptor). H0000 (SEQ ID NO:40) was used as the H-chain variable region of the GPC3-binding domain, and hCE115HA (SEQ ID NO:52) was used as the H-chain variable region of the CD3-binding domain. L0000 (SEQ ID NO:53) was used as the L-chain variable region. The D356K and K439E mutations introduced into the respective H chains were introduced for efficient heteromer formation of each H chain when producing heterodimeric antibodies comprising two types of H chains (WO 2006/106905). H435R is a modification that interrupts binding to Protein A, and was introduced for efficient separation of the heteromer and homomer (WO/2011/078332).

[0522] A series of expression vectors inserted with a polynucleotide encoding each of H0000-ERY27_HK (SEQ ID NO:54), hCE115HA-ERY27_HE (SEQ ID NO:55), and L0000-k0 (SEQ ID NO:56) were produced by well-known methods.

[0523] The following combination of expression vectors were introduced into FreeStyle 293-F cells for transient expression of each target molecule.

[0524] Target molecule: GPC3_ERY27_hCE115

[0525] Polypeptides encoded by the polynucleotides inserted into the expression vectors: H0000-ERY27_HK, hCE115HA-ERY27_HE, and L0000-k0

(2-6) Purification of GPC3_ERY27_hCE115

[0526] Each molecule of interest was purified by the method described in Example 1-2.

(2-7) Measurement of Cytotoxic Activity Using Human Peripheral Blood Mononuclear Cells

(2-7-1) Preparation of a Human Peripheral Blood Mononuclear Cell (PBMC) Solution

[0527] The solution was prepared by the method described in Example 1-3-1.

(2-7-2) Measurement of Cytotoxic Activity

[0528] Cytotoxic activity was measured by the method described in Example 1-3-2.

[0529] When PBMCs prepared from human blood were used as the effector cell to measure the cytotoxicity of GPC3_ERY27_hCE115, reduction of the activity was observed as a result of humanization of the H chain of rCE115 and sharing of a common L chain (FIG. 2).

[Example 3] Production and Evaluation of Humanized Bispecific Antibody Variants for Improvement of Various Properties

[0530] The T-cell-dependent cytotoxic activity of the humanized anti-human CD38 (CD3 epsilon) chain and anti-human GPC3 bispecific antibody obtained in Example 2, GPC3_ERY27_hCE115 (SEQ ID NOS: 54, 55, and 56), was lower than the T-cell-dependent cytotoxic activity of GPC3_

ERY22_rCE115 (SEQ ID NOS: 47, 48, 49, and 50). This may be due to attenuation of affinity for GPC3 and the CD38 chain as a result of humanization and sharing of a common L chain. Regarding GPC3 and CD38-chain antigens which have independent sequences, there has been no report so far on humanized bispecific antibodies whose T-cell dependent cytotoxic activity has been enhanced and whose affinity for both antigens has been improved by using a common antibody L chain. Therefore, it has been considered difficult to obtain humanized antibodies with dual specificity that show a drug efficacy equivalent to or greater than that of GPC3_ERY22_rCE115.

[0531] Under such circumstances, the Applicants produced modified humanized bispecific antibodies with modified affinity for human GPC3 and human CD38 chain by methods known to those skilled in the art, which involves comprehensively substituting amino acid residues encoded by the antibody gene to produce antibody variants against both the human GPC3 and human CD38-chain antigens, and by performing various evaluations by screening. Furthermore, similar methods were used to produce modified humanized bispecific antibodies with modified physicochemical properties. Furthermore, by combining substitutions of amino acid residues effective for modifying affinity and physicochemical properties, optimized bispecific antibodies having a TDCC activity equivalent to or greater than the T-cell dependent cellular cytotoxicity of GPC3_ERY22_rCE115 prior to humanization were produced.

[0532] Introduction of point mutations, expression and purification of antibodies, antigen affinity measurements, and determination of T-cell dependent cellular cytotoxicity in the optimization of humanized bispecific antibodies were performed by methods similar to those in Examples 1 and 2. CDR and FR were determined according to the Kabat definition (Kabat numbering).

[0533] Depending on the objective, the following were used as the antibody H-chain constant regions (the numbers indicate EU numbering): E22Hh (SEQ ID NO:57) produced by introducing L234A/L235A/N297A/D356C/T366S/L368A/Y407V/G446 deletion/K447 deletion mutations into human IgG1; E22Hk (SEQ ID NO:58) produced by introducing L234A/L235A/N297A/Y349C/T366W/G446 deletion/K447 deletion mutations and a Ser-Ser insertion mutation immediately before position 118 into human IgG1; Gldh produced by introducing D356C/T366S/L368A/Y407V/G446 deletion/K447 deletion mutations into human IgG1; none-Hi-Kn010G3 produced by introducing 118-215 deletion and C220S/Y349C/T366W/H435R mutations into human IgG1; E2270GsKsc (SEQ ID NO:60) produced by introducing L235R/S239K/N297A/E356K/R409K/H435R/L445P/G446 deletion/K447 deletion mutations into human IgG4; E2704sEpsc (SEQ ID NO:61) produced by introducing K196Q/L235R/S239K/N297A/R409K/K439E/L445P/G446 deletion/K447 deletion mutations into human IgG4; and E2702sKsc (SEQ ID NO:62) produced by introducing L235R/S239K/N297A/E356K/R409K/L445P/G446 deletion/K447 deletion mutations into human IgG4. Furthermore, human K (kappa) chain k0 (SEQ ID NO:63) and E22L (SEQ ID NO:432) produced by introducing R108A/T109S mutations into human K chain were used as the antibody L-chain constant regions.

[0534] The mutation that substitutes Cys for Asp at EU numbering position 356, the mutation that substitutes Ser for The at EU numbering position 366, the mutation that sub-

stitutes Ala for Leu at EU numbering position 368, the mutation that substitutes Val for Tyr at EU numbering position 407, the mutation that substitutes Cys for Tyr at EU numbering position 349, the mutation that substitutes Trp for Thr at EU numbering position 366, and the mutation that inserts Ser-Ser immediately before position 118 are mutations for efficient formation of heterodimeric molecules for each H chain when producing heteromeric antibodies. Similarly, the mutation that substitutes Lys for Glu at EU numbering position 356 and the mutation that substitutes Glu for Lys at EU numbering position 439 are also mutations for efficient formation of heterodimeric molecules for each H chain when producing heteromeric antibodies. They are expected to improve the efficiency of bispecific antibody production.

[0535] The mutation that substitutes Ala for Leu at EU numbering position 234, the mutation that substitutes Ala or Arg for Leu at EU numbering position 235, the mutation that substitutes Lys for Ser at EU numbering position 239, and the mutation that substitutes Ala for Asn at EU numbering position 297 are mutations for attenuating affinity for an Fcγ receptor and a complement (Clq). They are expected to suppress the binding of Fab to CD3 and Fc-mediated cross-linking of an Fcγ receptor or a complement, and avoid cytokine release syndrome that accompanies enhancement of non-specific effector functions.

[0536] The H chain introduced with deletion mutations at EU numbering positions 118 to 215 can be combined with a full-length H chain sequence to produce an antibody that has only one Fab (monovalent antibody), and it is useful for affinity evaluation.

[0537] The mutation that substitutes Lys for Arg at EU numbering position 409 and the mutation that substitutes Arg for His at EU numbering position 435 are mutations for modifying the antibody properties to be close to the properties of human IgG1 and human IgG3, respectively.

(3-1) Modifying the Affinity of a Humanized Anti-CD3 Antibody by Point Mutations

[0538] First, point mutations were introduced into FR1, FR2, FR3, CDR1, CDR2, and CDR3 of the humanized anti-human CD38 chain antibody sequence produced in Example 2, hCE115HA-ERY27_HE (SEQ ID NO:55), to prepare modified antibodies. Next, the affinity of these modified antibodies for the soluble human CD38 chain was determined. Combining sites that have an affinity-enhancing effect yielded modified antibodies having the affinities shown in Table 8.

TABLE 8

Antibody name	KD (Human CD3)
hCE115HA-E22Hh/-Hi-Kn010G3/L0000-k0	1.43E-07
TR01H083-E22Hh/none-Hi-Kn010G3/L0212-k0	5.86E-11
TR01H040-E22Hh/none-Hi-Kn010G3/L0240-k0	2.17E-09
TR01H002-E22Hh/GLS3108-k0/GL4-E22Hk/H0610-E22L	2.04E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0212-k0	2.17E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0235-k0	2.81E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0238-k0	2.91E-09
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L016-k0	2.52E-09
TR01H083-E22Hh/none-Hi-Kn010G3/L0262-k0	2.45E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0207-k0	2.60E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0241-k0	3.48E-09
CE115HA236-E22Hh/GLS3108-k0/GL4-E22Hk/H0610-E22L	3.58E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0206-k0	2.90E-09
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L019-k0	3.20E-09
TR01H080-E22Hh/none-Hi-Kn010G3/L0000-k0	3.25E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0211-k0	3.22E-09
TR01H002-E22Hh/-Hi-Kn010G3/GLC3108-k0	4.61E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0209-k0	4.25E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0208-k0	4.16E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0224-k0	5.06E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0236-k0	5.64E-09
TR01H083-E22Hh/none-Hi-Kn010G3/L0201-k0	4.42E-09
TR01H084-E2702GsKsc/none-Hi-E2704sE/L0011-k0	4.14E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0210-k0	5.06E-09
TR01H114-E2702GsKsc/none-Hi-E2704sE/L0011-k0	4.22E-09
CE115HA236-E22Hh/GLS3108-k0/GL4-E22Hk/H0610-E22L	6.08E-09
TR01H077-E22Hh/none-Hi-Kn010G3/L0200-k0	6.12E-09
TR01H071-E22Hh/none-Hi-Kn010G3/L0200-k0	6.13E-09
TR01H111-E2702GsKsc/none-Hi-E2704sE/L0011-k0	4.91E-09
TR01H081-E22Hh/none-Hi-Kn010G3/L0262-k0	5.76E-09
TR01H001-E22Hh/-Hi-Kn010G3/GLC3108-k0	8.22E-09
CE115HA179-G1dh/-Hi-Kn010G3/L0000-k0	8.35E-09
TR01H112-E2702GsKsc/none-Hi-E2704sE/L0011-k0	5.12E-09
TR01H113-E2702GsKsc/none-Hi-E2704sE/L0011-k0	5.14E-09
TR01H082-E22Hh/none-Hi-Kn010G3/L0212-k0	4.75E-09
CE115HA236-E22Hh/-Hi-Kn010G3/GLC3108-k0	9.10E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0231-k0	7.75E-09
TR01H037-E22Hh/none-Hi-Kn010G3/L0000-k0	6.93E-09
CE115HA252-E22Hh/-Hi-Kn010G3/L0000-k0	9.48E-09
TR01H083-E22Hh/none-Hi-Kn010G3/L0011-k0	6.70E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0223-k0	8.15E-09
TR01H083-E22Hh/none-Hi-Kn010G3/L0000-k0	6.83E-09
TR01H071-E22Hh/none-Hi-Kn010G3/L0000-k0	8.85E-09
TR01H087-E22Hh/none-Hi-Kn010G3/L0212-k0	5.88E-09
CE115HA178-G1dh/-Hi-Kn010G3/L0000-k0	1.09E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0237-k0	1.02E-08
TR01H083-E22Hh/none-Hi-Kn010G3/L0222-k0	9.42E-09
TR01H084-E22Hh/none-Hi-Kn010G3/L0262-k0	8.51E-09
TR01H071-E22Hh/none-Hi-Kn010G3/L0215-k0	9.51E-09
TR01H040-E22Hh/none-Hi-Kn010G3/L0218-k0	8.20E-09
TR01H081-E22Hh/none-Hi-Kn010G3/L0201-k0	9.46E-09
TR01H071-E22Hh/none-Hi-Kn010G3/L0222-k0	1.04E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0220-k0	9.15E-09
TR01H067-E22Hh/none-Hi-Kn010G3/TR01L016-k0	1.09E-08
TR01H002-E22Hh/GLS3108-k0/GL4-E22Hk/H0000-E22L	4.78E-09
TR01H067-E22Hh/none-Hi-Kn010G3/TR01L019-k0	1.21E-08
TR01H038-E22Hh/none-Hi-Kn010G3/L0000-k0	1.24E-08
TR01H061-E22Hh/none-Hi-Kn010G3/L0200-k0	1.27E-08
TR01H082-E2702GsKsc/none-Hi-E2704sE/L0011-k0	1.01E-08
CE115HA180-G1dh/-Hi-Kn010G3/L0000-k0	1.68E-08
CE115HA251-E22Hh/L0000-k0/GL4-E22Hk/H0610-E22L	1.37E-08
TR01H100-E2702GsKsc/none-Hi-E2704sE/L0011-k0	1.11E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0228-k0	1.60E-08
TR01H081-E22Hh/none-Hi-Kn010G3/L0011-k0	1.35E-08
TR01H061-E22Hh/none-Hi-Kn010G3/L0215-k0	1.54E-08
TR01H110-E2702GsKsc/none-Hi-E2704sE/L0011-k0	1.26E-08
TR01H043-E22Hh/none-Hi-Kn010G3/L0000-k0	1.52E-08
TR01H081-E22Hh/none-Hi-Kn010G3/L0000-k0	1.56E-08
CE115HA251-E22Hh/-Hi-Kn010G3/L0000-k0	2.23E-08
TR01H091-E2702GsKsc/none-Hi-E2704sE/L0011-k0	1.39E-08
CE115HA36-E22Hh/GLS3108-k0/GL4-E22Hk/H0000-E22L	6.95E-09
TR01H084-E22Hh/none-Hi-Kn010G3/L0201-k0	1.65E-08
TR01H072-E22Hh/none-Hi-Kn010G3/L0000-k0	2.03E-08
TR01H099-E2702GsKsc/none-Hi-E2704sE/L0011-k0	1.46E-08
TR01H061-E22Hh/none-Hi-Kn010G3/L0222-k0	1.88E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0239-k0	2.31E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0262-k0	1.81E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0234-k0	2.40E-08
TR01H012-E22Hh/none-Hi-Kn010G3/L0000-k0	7.94E-09
TR01H061-E22Hh/none-Hi-Kn010G3/L0000-k0	1.71E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0243-k0	2.46E-08
TR01H109-E2702GsKsc/none-Hi-E2704sE/L0011-k0	1.64E-08
TR01H047-E22Hh/none-Hi-Kn010G3/L0000-k0	2.04E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0267-k0	2.29E-08

TABLE 8-continued

TABLE 8-continued

Antibody name	KD (Human CD3)
TR01H082-E22Hh/none-Hi-Kn010G3/L0266-k0	2.29E-08
TR01H084-E22Hh/none-Hi-Kn010G3/L0011-k0	1.98E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0250-k0	2.15E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0204-k0	2.21E-08
TR01H084-E22Hh/none-Hi-Kn010G3/L0000-k0	2.13E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0213-k0	2.01E-08
hCE115HA-E22Hh/-Hi-Kn010G3/L0000-k0	1.43E-07
TR01H040-E22Hh/none-Hi-Kn010G3/L0214-k0	2.02E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0217-k0	2.07E-08
TR01H071-E22Hh/none-Hi-Kn010G3/L0226-k0	2.51E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0200-k0	2.87E-08
TR01H074-E22Hh/none-Hi-Kn010G3/L0000-k0	2.91E-08
TR01H039-E22Hh/none-Hi-Kn010G3/L0000-k0	2.61E-08
CE115HA177-G1dh//Hi-Kn010G3/L0000-k0	3.55E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0201-k0	2.81E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0263-k0	3.09E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0000-k0	3.60E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0216-k0	2.53E-08
TR01H051-E22Hh/none-Hi-Kn010G3/L0000-k0	2.91E-08
TR01H003-E22Hh/-Hi-Kn010G3/L0000-k0	4.03E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0264-k0	3.44E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0232-k0	3.86E-08
TR01H041-E22Hh/none-Hi-Kn010G3/L0000-k0	3.16E-08
CE115HA122-E22Hh/-Hi-Kn010G3/L0000-k0	4.28E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0233-k0	4.01E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0215-k0	3.37E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0203-k0	3.24E-08
TR01H015-E2702GsKsc/GCH019-E2704sEpsc/L0000-k0	2.96E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L008-k0	2.93E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0205-k0	3.42E-08
TR01H015-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	3.57E-08
TR01H064-E22Hh/none-Hi-Kn010G3/L0000-k0	3.07E-08
TR01H044-E22Hh/none-Hi-Kn010G3/L0000-k0	3.52E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0262-k0	3.98E-08
TR01H062-E22Hh/none-Hi-Kn010G3/L0000-k0	3.13E-08
CE115HA251-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	1.48E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0011-k0	3.48E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0222-k0	4.65E-08
CE115HA192-E22Hh/-Hi-Kn010G3/L0000-k0	5.05E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L010-k0	3.28E-08
TR01H025-E22Hh/none-Hi-Kn010G3/L0000-k0	3.86E-08
TR01H082-E22Hh/none-Hi-Kn010G3/TR01L023-k0	4.25E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L015-k0	3.95E-08
TR01H055-E22Hh/none-Hi-Kn010G3/L0000-k0	3.88E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0260-k0	4.53E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L009-k0	3.56E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L011-k0	3.57E-08
TR01H017-E22Hh/none-Hi-Kn010G3/L0000-k0	3.50E-08
CE115HA122-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	1.69E-08
TR01H076-E22Hh/none-Hi-Kn010G3/L0000-k0	4.78E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0258-k0	4.70E-08
TR01H046-E22Hh/none-Hi-Kn010G3/L0000-k0	4.23E-08
rCE115H-G1dh/-Hi-Kn010G3/L0000-k0	5.76E-08
TR01H082-E22Hh/none-Hi-Kn010G3/TR01L024-k0	4.76E-08
TR01H16-E22Hh/none-Hi-Kn010G3/L0000-k0	3.69E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L18-k0	4.51E-08
TR01H084-E22Hh/none-Hi-Kn010G3/L0271-k0	2.76E-08
TR01H084-E22Hh/none-Hi-Kn010G3/L0270-k0	2.76E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0000vk1-k0	4.69E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0219-k0	3.94E-08
TR01H014-E22Hh/none-Hi-Kn010G3/L0000-k0	3.87E-08
TR01H061-E22Hh/none-Hi-Kn010G3/L0226-k0	4.71E-08
TR01H048-E22Hh/none-Hi-Kn010G3/L0000-k0	4.52E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0259-k0	5.07E-08
TR01H028-E22Hh/none-Hi-Kn010G3/L0000-k0	4.80E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0201-k0	4.49E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L013-k0	4.15E-08
TR01H033-E22Hh/none-Hi-Kn010G3/L0000-k0	4.88E-08
hCE115HA-G1dh/-Hi-Kn010G3/L0000-k0	6.50E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L012-k0	4.22E-08
TR01H065-E22Hh/none-Hi-Kn010G3/L0000-k0	4.31E-08
TR01H079-E22Hh/none-Hi-Kn010G3/L0000-k0	5.05E-08
TR01H042-E22Hh/none-Hi-Kn010G3/L0000-k0	4.48E-08

TABLE 8-continued

Antibody name	KD (Human CD3)
TR01H063-E22Hh/none-Hi-Kn010G3/L0000-k0	4.35E-08
TR01H084-E22Hh/none-Hi-Kn010G3/L0272-k0	3.10E-08
CE115HA121-E22Hh/-Hi-Kn010G3/L0000-k0	6.76E-08
TR01H026-E22Hh/none-Hi-Kn010G3/L0000-k0	5.12E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0262-k0	4.92E-08
TR01H073-E22Hh/none-Hi-Kn010G3/L0000-k0	5.97E-08
TR01H045-E22Hh/none-Hi-Kn010G3/L0000-k0	5.22E-08
TR01H007-E22Hh/none-Hi-Kn010G3/L0000-k0	2.17E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0203-k0	4.07E-08
TR01H032-E22Hh/none-Hi-Kn010G3/L0000-k0	5.73E-08
TR01H006-E22Hh/none-Hi-Kn010G3/L0000-k0	2.30E-08
TR01H013-E22Hh/none-Hi-Kn010G3/L0000-k0	4.94E-08
TR01H050-E22Hh/none-Hi-Kn010G3/L0000-k0	5.76E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0200-k0	6.03E-08
TR01H015-E22Hh/none-Hi-Kn010G3/L0000-k0	6.13E-08
hCE115HA-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	6.16E-08
hCE115HA-E22stHh/none-Hi-stKn010G3/L0000-k0	5.17E-08
TR01H069-E22Hh/none-Hi-Kn010G3/L0000-k0	7.11E-08
TR01H015-E22Hh/none-Hi-Kn010G3/TR01L003-k0	6.34E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0202-k0	6.19E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0201-k0	5.93E-08
TR01H020-E22Hh/none-Hi-Kn010G3/L0000-k0	6.48E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0011-k0	5.95E-08
hCE115HA-E22Hh/-Hi-Kn010G3/L0000-k0	1.43E-07
TR01H082-E22Hh/none-Hi-Kn010G3/TR01L018-k0	4.72E-08
TR01H015-E22Hh/none-Hi-Kn010G3/TR01L005-k0	6.53E-08
TR01H052-E22Hh/none-Hi-Kn010G3/L0000-k0	6.27E-08
TR01H036-E22Hh/none-Hi-Kn010G3/L0000-k0	6.50E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0203-k0	4.79E-08
TR01H030-E22Hh/none-Hi-Kn010G3/L0000-k0	6.54E-08
TR01H015-E22Hh/none-Hi-Kn010G3/TR01L001-k0	6.56E-08
TR01H100-E22Hh/none-Hi-Kn010G3/L0011-k0	6.25E-08
TR01H029-E22Hh/none-Hi-Kn010G3/L0000-k0	6.70E-08
TR01H19-E22Hh/none-Hi-Kn010G3/L0000-k0	6.85E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0000-k0	7.37E-08
TR01H018-E22Hh/none-Hi-Kn010G3/L0000-k0	6.93E-08
TR01H027-E22Hh/none-Hi-Kn010G3/L0000-k0	6.95E-08
TR01H049-E22Hh/none-Hi-Kn010G3/L0000-k0	6.79E-08
TR01H066-E22Hh/none-Hi-Kn010G3/L0000-k0	6.02E-08
TR01H091-E22Hh/none-Hi-Kn010G3/L0011-k0	6.67E-08
rCE115H-E22Hh/none-Hi-Kn010G3/L0000-k0	8.00E-08
TR01H015-E22Hh/none-Hi-Kn010G3/TR01L002-k0	7.14E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0226-k0	8.01E-08
TR01H067-E22Hh/none-Hi-Kn010G3/TR01L018-k0	5.26E-08
TR01H093-E22Hh/none-Hi-Kn010G3/L0011-k0	6.80E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0215-k0	7.41E-08
TR01H015-E22Hh/none-Hi-Kn010G3/TR01L004-k0	7.34E-08
TR01H107-E22Hh/none-Hi-Kn010G3/L0011-k0	6.91E-08
TR01H105-E22Hh/none-Hi-Kn010G3/L0011-k0	6.95E-08
TR01H090-E22Hh/none-Hi-Kn010G3/L0011-k0	6.95E-08
TR01H108-E22Hh/none-Hi-Kn010G3/L0011-k0	6.98E-08
TR01H094-E22Hh/none-Hi-Kn010G3/L0011-k0	7.00E-08
TR01H109-E22Hh/none-Hi-Kn010G3/L0011-k0	7.06E-08
TR01H056-E22Hh/none-Hi-Kn010G3/L0000-k0	7.32E-08
TR01H031-E22Hh/none-Hi-Kn010G3/L0000-k0	7.55E-08
TR01H022-E22Hh/none-Hi-Kn010G3/L0000-k0	7.58E-08
TR01H092-E22Hh/none-Hi-Kn010G3/L0011-k0	7.21E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0000-k0	7.15E-08
TR01H067-E22Hh/none-Hi-Kn010G3/L0011-k0	7.18E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0248-k0	7.89E-08
TR01H009-E22Hh/none-Hi-Kn010G3/L0000-k0	3.15E-08
TR01H023-E22Hh/none-Hi-Kn010G3/L0000-k0	7.94E-08
TR01H096-E22Hh/none-Hi-Kn010G3/L0011-k0	7.47E-08
TR01H040-E22Hh/none-Hi-Kn010G3/TR01L007-k0	6.82E-08
TR01H054-E22Hh/none-Hi-Kn010G3/L0000-k0	7.79E-08
TR01H021-E22Hh/none-Hi-Kn010G3/L0000-k0	8.05E-08
TR01H103-E22Hh/none-Hi-Kn010G3/L0011-k0	7.72E-08
TR01H099-E22Hh/none-Hi-Kn010G3/L0011-k0	7.74E-08
rCE115H-E22Hh/none-Hi-Kn010G3/L0000vk1-k0	8.52E-08
TR01H101-E22Hh/none-Hi-Kn010G3/L0011-k0	7.87E-08
TR01H053-E22Hh/none-Hi-Kn010G3/L0000-k0	8.23E-08
TR01H035-E22Hh/none-Hi-Kn010G3/L0000-k0	8.49E-08
TR01H067-E22Hh/none-Hi-Kn010G3/TR01L015-k0	8.64E-08

TABLE 8-continued

Antibody name	KD (Human CD3)
TR01H104-E22Hh/none-Hi-Kn010G3/L0011-k0	8.26E-08
TR01H075-E22Hh/none-Hi-Kn010G3/L0000-k0	9.88E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0227-k0	1.01E-07
TR01H102-E22Hh/none-Hi-Kn010G3/L0011-k0	8.54E-08
TR01H034-E22Hh/none-Hi-Kn010G3/L0000-k0	9.11E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0222-k0	1.01E-07
rCE115H-E22Hh/rCE115L-k0/GL4-E22Hk/H0000-E22L	9.37E-08
TR01H015-E22Hh/none-Hi-Kn010G3/TR01L006-k0	9.30E-08
TR01H040-E22Hh/none-Hi-Kn010G3/L0246-k0	9.28E-08
TR01H097-E22Hh/none-Hi-Kn010G3/L0011-k0	8.76E-08
TR01H011-E22Hh/none-Hi-Kn010G3/L0000-k0	3.71E-08
TR01H010-E22Hh/none-Hi-Kn010G3/L0000-k0	3.73E-08
TR01H095-E22Hh/none-Hi-Kn010G3/L0011-k0	9.09E-08
TR01H082-E22Hh/none-Hi-Kn010G3/TR01L020-k0	1.06E-07
TR01H098-E22Hh/none-Hi-Kn010G3/L0011-k0	9.14E-08
TR01H082-E22Hh/none-Hi-Kn010G3/TR01L017-k0	1.09E-07
TR01H040-E22Hh/none-Hi-Kn010G3/L0247-k0	1.00E-07
rCE115H-E22Hh/none-Hi-Kn010G3/rCE115L-k0	1.24E-07
TR01H004-E22Hh//Hi-Kn010G3/L0000-k0	1.35E-07
TR01H067-E22Hh/none-Hi-Kn010G3/L0222-k0	7.63E-08
rCE115H-E22Hh//Hi-Kn010G3/rCE115L-k0	1.38E-07
TR01H008-E22Hh/none-Hi-Kn010G3/L0000-k0	4.22E-08
TR01H070-E22Hh/none-Hi-Kn010G3/L0000-k0	1.20E-07
TR01H106-E22Hh/none-Hi-Kn010G3/L0011-k0	1.00E-07
TR01H024-E22Hh/none-Hi-Kn010G3/L0000-k0	1.08E-07
CE115HA124-E22Hh//Hi-Kn010G3/L0000-k0	1.43E-07
TR01H040-E22Hh/none-Hi-Kn010G3/L0249-k0	1.11E-07
TR01H082-E22Hh/none-Hi-Kn010G3/L0271-k0	6.82E-08
TR01H057-E22Hh/none-Hi-Kn010G3/L0000-k0	1.12E-07
TR01H058-E22Hh/none-Hi-Kn010G3/L0000-k0	1.15E-07
TR01H068-E22Hh/none-Hi-Kn010G3/L0000-k0	1.01E-07
TR01H082-E22Hh/none-Hi-Kn010G3/L0270-k0	7.42E-08
TR01H082-E22Hh/none-Hi-Kn010G3/L0272-k0	7.44E-08
hCE115HA-E22Hh/none-Hi-Kn010G3/L0000-k0	1.24E-07
TR01H082-E22Hh/none-Hi-Kn010G3/L0268-k0	1.36E-07
hCE115HAA-E22Hh/none-Hi-Kn010G3/L0000-k0	1.08E-07
TR01H067-E22Hh/none-Hi-Kn010G3/L0226-k0	1.32E-07
TR01H067-E22Hh/none-Hi-Kn010G3/L0248-k0	1.39E-07

(3-2) Modifying the Affinity of a Humanized Anti-GPC3 Antibody

[0539] First, point mutations were introduced into CDR1, CDR2, and CDR3 of the anti-human GPC3 bispecific antibody sequence produced in Example 2, H0000-ERY27_HK (SEQ ID NO:54), to prepare modified antibodies. Next, the affinity of these modified antibodies for soluble human GPC3 was determined. Combining sites that have an affinity-enhancing effect yielded modified antibodies having the affinities shown in Table 9.

TABLE 9

Antibody name	KD (Human GPC3)
H0610-G1dh/none-Hi-Kn010G3/L0000-k0	3.97E-09
H0610-G1dh/none-Hi-Kn010G3/L0222-k0	1.40E-13
H0610-G1dh/none-Hi-Kn010G3/L0258-k0	3.52E-13
GCH054-G1dh/none-Hi-Kn010G3/L0262-k0	5.25E-13
GCH060-G1dh/none-Hi-Kn010G3/L0222-k0	6.42E-13
H0610-G1dh/none-Hi-Kn010G3/L0246-k0	1.21E-12
GCH057-G1dh/none-Hi-Kn010G3/L0222-k0	1.85E-12
GCH054-G1dh/none-Hi-Kn010G3/L0249-k0	3.61E-12
GCH055-G1dh/none-Hi-Kn010G3/L0222-k0	3.90E-12
GCH094-G1dh/none-Hi-Kn010G3/L0246-k0	4.12E-12
H0610-G1dh/none-Hi-Kn010G3/L0249-k0	6.86E-12

TABLE 9-continued

Antibody name	KD (Human GPC3)
H0610-G1dh/none-Hi-Kn010G3/TR01L017-k0	8.27E-12
H0610-G1dh/none-Hi-Kn010G3/L0265-k0	8.70E-12
H0610-G1dh/none-Hi-Kn010G3/L0261-k0	1.07E-11
GCH065-G1dh/none-Hi-Kn010G3/L0262-k0	1.18E-11
GCH054-G1dh/none-Hi-Kn010G3/L0262-k0	1.19E-11
H0610-G1dh/none-Hi-Kn010G3/L0268-k0	1.69E-11
H0610-G1dh/none-Hi-Kn010G3/TR01L020-k0	2.24E-11
GCH054-G1dh/none-Hi-Kn010G3/L0246-k0	3.15E-11
GCH054-G1dh/none-Hi-Kn010G3/L0222-k0	3.15E-11
GCH073-G1dh/none-Hi-Kn010G3/L0201-k0	3.50E-11
H0610-G1dh/none-Hi-Kn010G3/L0248-k0	5.55E-11
GCH065-G1dh/none-Hi-Kn010G3/L0201-k0	7.74E-11
H0610-G1dh/none-Hi-Kn010G3/L0226-k0	9.30E-11
H0610-G1dh/none-Hi-Kn010G3/L0093-k0	1.06E-10
GCH098-G1dh/none-Hi-Kn010G3/L0201-k0	1.11E-10
H0610-G1dh/none-Hi-Kn010G3/L0267-k0	1.79E-10
H0610-G1dh/none-Hi-Kn010G3/L0228-k0	2.02E-10
H0610-G1dh/none-Hi-Kn010G3/L0262-k0	2.11E-10
H0610-G1dh/none-Hi-Kn010G3/L0266-k0	2.13E-10
H0610-G1dh/none-Hi-Kn010G3/L0264-k0	2.19E-10
H0610-G1dh/none-Hi-Kn010G3/L0224-k0	2.43E-10
H0610-G1dh/none-Hi-Kn010G3/L0167-k0	2.11E-10
CE115HA251-E22Hh/L0000-k0/GL4-E22Hk/H0610-E22L	2.36E-10
TR01H015-E22Hh/L0000-k0/GL4-E22Hk/H0610-E22L	2.63E-10
CE115HA236-E22Hh/GLS3108-k0/GL4-E22Hk/H0610-E22L	2.67E-10
H0610-G1dh/none-Hi-Kn010G3/L0259-k0	3.34E-10
H0610-G1dh/none-Hi-Kn010G3/L0227-k0	4.08E-10
GCH065-G1dh/none-Hi-Kn010G3/L0272-k0	3.93E-10
H0610-G1dh/none-Hi-Kn010G3/L0269-k0	4.59E-10
H0610-G1dh/none-Hi-Kn010G3/L0223-k0	4.75E-10
TR01H002-E22Hh/GLS3108-k0/GL4-E22Hk/H0610-E22L	4.75E-10
GCH054-G1dh/none-Hi-Kn010G3/L0212-k0	5.17E-10
H0610-G1dh/none-Hi-Kn010G3/L0208-k0	5.30E-10
H0610-G1dh/none-Hi-Kn010G3/L0263-k0	5.64E-10
H0610-G1dh/none-Hi-Kn010G3/L0231-k0	5.89E-10
H0610-G1dh/none-Hi-Kn010G3/L0143-k0	5.73E-10
GCH055-G1dh/none-Hi-Kn010G3/L0212-k0	6.14E-10
H0610-G1dh/none-Hi-Kn010G3/L0211-k0	6.47E-10
H0610-G1dh/none-Hi-Kn010G3/L0238-k0	6.37E-10
H0610-G1dh/none-Hi-Kn010G3/L0214-k0	6.57E-10
H0610-G1dh/none-Hi-Kn010G3/L0243-k0	6.49E-10
GCH025-G1dh/none-Hi-Kn010G3/L0204-k0	6.70E-10
GCH054-G1dh/none-Hi-Kn010G3/TR01L016-k0	7.63E-10
H0610-G1dh/none-Hi-Kn010G3/L0168-k0	6.99E-10
GCH094-G1dh/none-Hi-Kn010G3/L0271-k0	6.92E-10
GCH054-G1dh/none-Hi-Kn010G3/TR01L019-k0	8.71E-10
H0610-G1dh/none-Hi-Kn010G3/L0234-k0	7.78E-10
GCH098-G1dh/none-Hi-Kn010G3/L0011-k0	8.02E-10
H0610-G1dh/none-Hi-Kn010G3/L0204-k0	7.27E-10
H0610-G1dh/none-Hi-Kn010G3/L0240-k0	8.48E-10
H0610-G1dh/none-Hi-Kn010G3/L0239-k0	8.74E-10
H0610-G1dh/none-Hi-Kn010G3/L0212-k0	9.94E-10
GCH065-G1dh/none-Hi-Kn010G3/L0011-k0	8.84E-10
H0610-G1dh/none-Hi-Kn010G3/L0200-k0	1.04E-09
H0610-G1dh/none-Hi-Kn010G3/L0124-k0	9.72E-10
GCH073-G1dh/none-Hi-Kn010G3/L0011-k0	9.10E-10
H0610-G1dh/none-Hi-Kn010G3/TR01L016-k0	1.08E-09
GCH054-G1dh/none-Hi-Kn010G3/L0201-k0	1.08E-09
H0610-G1dh/none-Hi-Kn010G3/L0090-k0	1.12E-09
H0610-G1dh/none-Hi-Kn010G3/L0209-k0	1.12E-09
H0610-G1dh/none-Hi-Kn010G3/L0201-k0	1.13E-09
H0610-G1dh/none-Hi-Kn010G3/L0161-k0	9.73E-10
H0610-G1dh/none-Hi-Kn010G3/L0206-k0	8.65E-10
H0610-G1dh/none-Hi-Kn010G3/L0186-k0	1.08E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L019-k0	1.15E-09
H0610-G1dh/none-Hi-Kn010G3/L0085-k0	1.17E-09
GCH055-G1dh/none-Hi-Kn010G3/L0200-k0	1.13E-09
H0610-G1dh/none-Hi-Kn010G3/L0154-k0	1.01E-09
H0610-G1dh/none-Hi-Kn010G3/L0229-k0	1.20E-09
GCH054-G1dh/none-Hi-Kn010G3/L0200-k0	1.18E-09
GCH094-G1dh/none-Hi-Kn010G3/L0201-k0	1.17E-09
H0610-G1dh/none-Hi-Kn010G3/L0000-k0	3.97E-09
H0610-G1dh/none-Hi-Kn010G3/L0205-k0	1.01E-09

TABLE 9-continued

Antibody name	KD (Human GPC3)
GCH099-G1dh/none-Hi-Kn010G3/L0201-k0	1.29E-09
H0610-G1dh/none-Hi-Kn010G3/L0242-k0	1.19E-09
GCH056-G1dh/none-Hi-Kn010G3/L0201-k0	1.16E-09
H0610-G1dh/none-Hi-Kn010G3/L0213-k0	1.25E-09
GCH060-G1dh/none-Hi-Kn010G3/L0200-k0	1.34E-09
GCH065-G1dh/none-Hi-Kn010G3/L0000-k0	1.41E-09
GCH100-G1dh/none-Hi-Kn010G3/L0201-k0	1.37E-09
H0610-G1dh/none-Hi-Kn010G3/L0015-k0	1.31E-09
H0610-G1dh/none-Hi-Kn010G3/L0151-k0	1.25E-09
H0610-G1dh/none-Hi-Kn010G3/L0237-k0	1.31E-09
H0610-G1dh/none-Hi-Kn010G3/L0220-k0	1.36E-09
H0610-G1dh/none-Hi-Kn010G3/L0155-k0	1.28E-09
GCH055-G1dh/none-Hi-Kn010G3/L0215-k0	1.52E-09
H0610-G1dh/none-Hi-Kn010G3/L0202-k0	1.22E-09
GCH056-G1dh/none-Hi-Kn010G3/L0215-k0	1.59E-09
H0610-G1dh/none-Hi-Kn010G3/L0012-k0	1.55E-09
GCH054-G1dh/none-Hi-Kn010G3/L0215-k0	1.62E-09
H0610-G1dh/none-Hi-Kn010G3/L0215-k0	1.64E-09
GCH098-G1dh/none-Hi-Kn010G3/L0000-k0	1.77E-09
H0610-G1dh/none-Hi-Kn010G3/L0125-k0	1.71E-09
GCH057-G1dh/none-Hi-Kn010G3/L0215-k0	1.83E-09
H0610-G1dh/none-Hi-Kn010G3/L0217-k0	1.79E-09
H0610-G1dh/none-Hi-Kn010G3/L0014-k0	1.82E-09
H0610-G1dh/none-Hi-Kn010G3/L0216-k0	1.86E-09
TR01H015-E2702GsKsc/GCH019-E2704sEpscL0000-k0	1.64E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L015-k0	2.16E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L018-k0	2.17E-09
H0610-G1dh/none-Hi-Kn010G3/L0218-k0	1.99E-09
H0610-G1dh/none-Hi-Kn010G3/L0000vk1-k0	2.16E-09
H0610-G1dh/none-Hi-Kn010G3/L0160-k0	2.12E-09
H0610-G1dh/none-Hi-Kn010G3/L0047-k0	2.23E-09
GCH073-G1dh/none-Hi-Kn010G3/L0000-k0	2.00E-09
GCH054-G1dh/none-Hi-Kn010G3/TR01L015-k0	2.45E-09
H0610-G1dh/none-Hi-Kn010G3/L0219-k0	2.28E-09
GCH094-G1dh/none-Hi-Kn010G3/L0272-k0	2.10E-09
H0610-G1dh/none-Hi-Kn010G3/L0149-k0	2.16E-09
GCH054-G1dh/none-Hi-Kn010G3/TR01L018-k0	2.59E-09
GCH054-G1dh/none-Hi-Kn010G3/L0203-k0	2.48E-09
H0610-G1dh/none-Hi-Kn010G3/L0122-k0	2.42E-09
H0610-G1dh/none-Hi-Kn010G3/L0134-k0	2.53E-09
H0610-G1dh/none-Hi-Kn010G3/L0152-k0	2.36E-09
H0610-G1dh/none-Hi-Kn010G3/L0203-k0	2.11E-09
H0610-G1dh/none-Hi-Kn010G3/L0075-k0	2.85E-09
H0610-G1dh/none-Hi-Kn010G3/L0038-k0	2.75E-09
H0610-G1dh/none-Hi-Kn010G3/L0011-k0	2.76E-09
H0610-G1dh/none-Hi-Kn010G3/L0157-k0	2.60E-09
H0610-G1dh/none-Hi-Kn010G3/L0145-k0	2.66E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L010-k0	2.92E-09
H0610-G1dh/none-Hi-Kn010G3/L0009-k0	2.99E-09
GCH099-G1dh/none-Hi-Kn010G3/L0011-k0	2.78E-09
H0610-G1dh/none-Hi-Kn010G3/L0006-k0	3.04E-09
H0610-G1dh/none-Hi-Kn010G3/L0173-k0	2.83E-09
H0610-G1dh/none-Hi-Kn010G3/L0127-k0	3.12E-09
H0610-G1dh/none-Hi-Kn010G3/L0082-k0	3.43E-09
H0610-G1dh/none-Hi-Kn010G3/L0064-k0	3.37E-09
H0610-G1dh/none-Hi-Kn010G3/L0008-k0	3.30E-09
H0610-G1dh/none-Hi-Kn010G3/L0013-k0	3.35E-09
H0610-G1dh/none-Hi-Kn010G3/L0140-k0	3.38E-09
H0610-G1dh/none-Hi-Kn010G3/L0039-k0	3.41E-09
GCH043-G1dh/none-Hi-Kn010G3/L0000-k0	3.74E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L008-k0	3.48E-09
H0610-G1dh/none-Hi-Kn010G3/L0148-k0	3.28E-09
GCH062-G1dh/none-Hi-Kn010G3/L0000-k0	3.73E-09
H0610-G1dh/none-Hi-Kn010G3/L0163-k0	3.38E-09
H0610-G1dh/none-Hi-Kn010G3/L0233-k0	3.55E-09
H0610-G1dh/none-Hi-Kn010G3/L0230-k0	4.00E-09
GCH006-G1dh/none-Hi-Kn010G3/L0000-k0	4.06E-09
H0610-G1dh/none-Hi-Kn010G3/L0032-k0	3.72E-09
H0610-G1dh/none-Hi-Kn010G3/L0181-k0	3.51E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L009-k0	3.81E-09
H0610-G1dh/none-Hi-Kn010G3/L0141-k0	3.86E-09
H0610-G1dh/none-Hi-Kn010G3/L0079-k0	4.23E-09
GCH094-G1dh/none-Hi-Kn010G3/L0270-k0	3.60E-09

TABLE 9-continued

Antibody name	KD (Human GPC3)
GCH066-G1dh/none-Hi-Kn010G3/L0000-k0	4.29E-09
GCH064-G1dh/none-Hi-Kn010G3/L0000-k0	4.14E-09
H0610-G1dh/none-Hi-Kn010G3/L0066-k0	4.20E-09
GCH027-G1dh/none-Hi-Kn010G3/L0000-k0	3.83E-09
H0610-G1dh/none-Hi-Kn010G3/L0003-k0	4.01E-09
H0610-G1dh/none-Hi-Kn010G3/L0042-k0	4.27E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L011-k0	4.02E-09
H0610-G1dh/none-Hi-Kn010G3/L0000-k0	3.97E-09
GCH015-G1dh/none-Hi-Kn010G3/L0000-k0	4.14E-09
H0610-G1dh/none-Hi-Kn010G3/L0175-k0	3.84E-09
GCH100-G1dh/none-Hi-Kn010G3/L0011-k0	3.81E-09
GCH014-G1dh/none-Hi-Kn010G3/L0000-k0	4.20E-09
GCH053-G1dh/none-Hi-Kn010G3/L0000-k0	4.05E-09
rCE115HA-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	4.28E-09
GCH094-G1dh/none-Hi-Kn010G3/L0011-k0	3.88E-09
GCH045-G1dh/none-Hi-Kn010G3/L0000-k0	4.63E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L012-k0	4.25E-09
H0610-G1dh/none-Hi-Kn010G3/L0115-k0	4.34E-09
H0610-G1dh/none-Hi-Kn010G3/L0044-k0	4.57E-09
H0610-G1dh/none-Hi-Kn010G3/L0107-k0	4.38E-09
H0610-G1dh/none-Hi-Kn010G3/L0007-k0	4.39E-09
GCH013-G1dh/none-Hi-Kn010G3/L0000-k0	4.44E-09
H0610-G1dh/none-Hi-Kn010G3/L0045-k0	4.66E-09
GCH010-G1dh/none-Hi-Kn010G3/L0000-k0	4.12E-09
GCH040-G1dh/none-Hi-Kn010G3/L0000-k0	4.80E-09
H0610-G1dh/none-Hi-Kn010G3/L0002-k0	4.43E-09
H0610-G1dh/none-Hi-Kn010G3/L0016-k0	4.44E-09
GCH007-G1dh/none-Hi-Kn010G3/L0000-k0	4.93E-09
GCH042-G1dh/none-Hi-Kn010G3/L0000-k0	4.89E-09
rCE115H-E22Hh/rCE115L-k0/GL4-E22Hk/H0000-E22L	4.57E-09
H0610-G1dh/none-Hi-Kn010G3/L0129-k0	4.54E-09
H0610-G1dh/none-Hi-Kn010G3/L0065-k0	4.79E-09
GCH016-G1dh/none-Hi-Kn010G3/L0000-k0	4.59E-09
GCH035-G1dh/none-Hi-Kn010G3/L0000-k0	4.94E-09
GCH039-G1dh/none-Hi-Kn010G3/L0000-k0	4.95E-09
GCH099-G1dh/none-Hi-Kn010G3/L0000-k0	4.24E-09
H0610-G1dh/none-Hi-Kn010G3/L0041-k0	4.85E-09
GCH019-G1dh/none-Hi-Kn010G3/L0000-k0	4.36E-09
GCH029-G1dh/none-Hi-Kn010G3/L0000-k0	5.01E-09
GCH056-G1dh/none-Hi-Kn010G3/L0011-k0	4.31E-09
H0610-G1dh/none-Hi-Kn010G3/L0147-k0	4.38E-09
GCH034-G1dh/none-Hi-Kn010G3/L0000-k0	5.09E-09
GCH003-G1dh/none-Hi-Kn010G3/L0000-k0	5.20E-09
H0610-G1dh/none-Hi-Kn010G3/L0139-k0	4.78E-09
H0610-G1dh/none-Hi-Kn010G3/L0089-k0	5.24E-09
H0610-G1dh/none-Hi-Kn010G3/L0113-k0	4.82E-09
H0610-G1dh/none-Hi-Kn010G3/L0180-k0	4.48E-09
GCH005-G1dh/none-Hi-Kn010G3/L0000-k0	5.32E-09
GCH067-G1dh/none-Hi-Kn010G3/L0000-k0	5.24E-09
H0610-G1dh/none-Hi-Kn010G3/L0187-k0	4.92E-09
H0610-G1dh/none-Hi-Kn010G3/L0043-k0	5.14E-09
H0610-G1dh/none-Hi-Kn010G3/L0117-k0	4.92E-09
GCH061-G1dh/none-Hi-Kn010G3/L0000-k0	5.13E-09
GCH022-G1dh/none-Hi-Kn010G3/L0000-k0	4.92E-09
H0610-G1dh/none-Hi-Kn010G3/L0091-k0	5.43E-09
GCH023-G1dh/none-Hi-Kn010G3/L0000-k0	4.94E-09
H0610-G1dh/none-Hi-Kn010G3/L0062-k0	5.28E-09
H0610-G1dh/none-Hi-Kn010G3/L0136-k0	5.04E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L003-k0	5.08E-09
H0610-G1dh/none-Hi-Kn010G3/L0069-k0	5.32E-09
H0610-G1dh/none-Hi-Kn010G3/L0123-k0	5.08E-09
GCH025-G1dh/none-Hi-Kn010G3/L0000-k0	5.05E-09
GCH100-G1dh/none-Hi-Kn010G3/L0000-k0	5.39E-09
H0610-G1dh/none-Hi-Kn010G3/L0046-k0	5.45E-09
H0610-G1dh/none-Hi-Kn010G3/L0144-k0	4.84E-09
GCH026-G1dh/none-Hi-Kn010G3/L0000-k0	5.17E-09
H0610-G1dh/none-Hi-Kn010G3/L0138-k0	5.24E-09
GCH056-G1dh/none-Hi-Kn010G3/L0000-k0	5.03E-09
H0610-G1dh/none-Hi-Kn010G3/L0129-k0	5.28E-09
GCH032-G1dh/none-Hi-Kn010G3/L0000-k0	5.74E-09
H0610-G1dh/none-Hi-Kn010G3/TR01L005-k0	5.37E-09
GCH012-G1dh/none-Hi-Kn010G3/L0000-k0	5.40E-09
GCH055-G1dh/none-Hi-Kn010G3/L0000-k0	5.60E-09

TABLE 9-continued

Antibody name	KD (Human GPC3)
H0610-G1dh/none-Hi-Kn010G3/L0104-k0	5.90E-09
GCH059-G1dh/none-Hi-Kn010G3/L0000-k0	5.70E-09
GCH054-G1dh/none-Hi-Kn010G3/L0000-k0	5.30E-09
GCH008-G1dh/none-Hi-Kn010G3/L0000-k0	5.55E-09
H0610-G1dh/none-Hi-Kn010G3/L0232-k0	5.38E-09
H0610-G1dh/none-Hi-Kn010G3/L0126-k0	5.62E-09
GCH094-G1dh/none-Hi-Kn010G3/L0000-k0	5.89E-09
H0610-G1dh/none-Hi-Kn010G3/L0132-k0	5.65E-09
H0610-G1dh/none-Hi-Kn010G3/L0106-k0	5.66E-09
GCH054-G1dh/none-Hi-Kn010G3/L0011-k0	5.25E-09
H0610-G1dh/none-Hi-Kn010G3/L0109-k0	5.70E-09
H0610-G1dh/none-Hi-Kn010G3/L0063-k0	6.03E-09
GCH068-G1dh/none-Hi-Kn010G3/L0000-k0	6.23E-09
GCH057-G1dh/none-Hi-Kn010G3/L0000-k0	5.61E-09
H0610-G1dh/none-Hi-Kn010G3/L0137-k0	5.87E-09

(3-3) Modification of pI by Point Mutations

[0540] In commercial production of bispecific antibodies, a high level of purity is required. When using ion-exchange chromatography, modifying the molecular isoelectric point (pI) has been reported to be effective (PLOS One. 2013; 8 (2): e57479). Therefore, point mutations for pI modifications were introduced into CDR1, CDR2, and CDR3 of the humanized anti-human GPC3 antibody sequence produced in Example 2, H0000-ERY27_HK (SEQ ID NO:54), to prepare modified antibodies. Next, the affinity of these modified antibodies for soluble human GPC3 was determined.

[0541] As a result, amino acid modifications that can lower the pI while maintaining the affinity for human GPC3 were found to be amino acids at positions 19, 43, 53, and 61 according to Kabat numbering.

[0542] Combination of sites showing effects of maintaining the affinity for human GPC3 and lowering the pI yielded antibodies having the affinities and pI values shown in Table 10.

TABLE 10

Antibody name (homomeric antibody)	Calculated pI value (homomeric antibody)	Antibody name (single-arm antibody)	human GPC3 KD (single-arm antibody)	Mutation sites based on H0610-E2704sEpsc
H0610-E2704sEpsc/L0000-k0	7.8	H0610-G1dh/none-Hi-Kn010G3/L0000-K0	4.16E-09	—
GCH054-E2704sEpsc/L0011-k0	6.2	GCH054-G1dh/none-Hi-Kn010G3/L0011-k0	5.25E-09	K19T/Q43E/P52aG/K53E/G55P/Q61E
GCH065-E2704sEpsc/L0011-k0	6.4	GCH065-G1dh/none-Hi-Kn010G3/L0011-k0	8.84E-10	K19T/Q43E/P52aG/K53P/G55P/Q61E
GCH094-E2704sEpsc/L0011-k0	6.2	GCH094-G1dh/none-Hi-Kn010G3/L0011-K0	4.54E-09	K19T/I37V/P40A/Q43E/I48M/P52aG/ K53E/G55P/Q61E

(3-4) Modifying the Extracellular Matrix-Binding Ability by Point Mutation

[0543] It has been reported that non-specific binding to the extracellular matrix (ECM) and such may have effects on pharmacokinetics (MAbs. 2012 November-December; 4 (6): 753-60). Therefore, the ECM-binding ability of the modified antibodies obtained in the Examples was determined by the method described in Reference Example 4. As a result, the humanized anti-human CD38 chain and anti-human GPC3 bispecific antibody, GPC3_ERY27_hCE115 (SEQ ID NOS: 54, 55, and 56), were confirmed to have high ECM-binding

abilities. Therefore, any of the point mutations examined in Examples 3-1, 3-2, and 3-3 for the humanized anti-human CD38 chain antibody sequence hCE115HA-ERY27_HE (SEQ ID NO: 55) was investigated to be a combination for reducing the ECM-binding ability. As a result, amino acids at positions 11, 16, 52a, 53, 98, and 100 by Kabat numbering were found to contribute to the maintenance of affinity for CD38 and to have influence on the reduction of the ECM-binding ability, and antibodies with a reduced ECM-binding ability in comparison to that of an antibody variant of the humanized anti-human CD38 chain and anti-human GPC3 bispecific antibody, GPC3_ERY27_hCE115, were obtained (Table 11).

TABLE 11

Antibody name	ECM binding ratio (standard = 1)
GPC3_ERY22_CE115 (rCE115H-E22Hh/rCE115L-k0/ GL4-E22Hk/H0000-E22L)	4.0
GPC3_ERY27 (hCE115HA-E22Hh/L0000-k0/GL4- E22Hk/H0000-E22L)	50.9
CE115HA236-E22Hh/GLS3108-k0/GL4-E22Hk/H0610- E22L	429.9
CE115HA236-E22Hh/GLS3108-k0/GL4-E22Hk/H0000- E22L	414.8
CE115HA251-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	346.9
CE115HA251-E22Hh/L0000-k0/GL4-E22Hk/H0610-E22L	334.4
TR01H002-E22Hh/GLS3108-k0/GL4-E22Hk/H0610-E22L	301.1
TR01H002-E22Hh/GLS3108-k0/GL4-E22Hk/H0000-E22L	216.9
TR01H011-E22Hh/L0000-k0/GL4-E22Hk/H0610-E22L	185.7
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0208-k0	50.4
CE115HA122-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L	47.0
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0211-k0	15.5
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0206-k0	15.4
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0209-k0	7.4
rCE115H-E22Hh/rCE115L-k0/GL4-E22Hk/H0610-E22L	4.6
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0204-k0	4.4
TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0212-k0	3.3
TR01H113-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0	2.5
TR01H082-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0	1.7
TR01H113-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0	1.6

TABLE 11-continued

Antibody name	ECM binding ratio (standard = 1)
rCE115H-E22Hh/rCE115L-k0/L0000-E22Hk/H0610-E22L	1.4
TR01H084-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0	1.3
TR01H084-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0	1.2
TR01H082-E2702GsKsc/GCH094-E2704sEpsc/L0201-k0	1.1
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0000-k0	0.8
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0201-k0	0.8

TABLE 11-continued

Antibody name	ECM binding ratio (standard = 1)
TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0203-k0	0.8
TR01H082-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0	0.7
TR01H109-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0	0.7
TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0222-k0	0.6
TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0201-k0	0.5
TR01H109-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0	0.4
TR01H113-E2702sKsc/GCH065-E2704sEpsc/L0011-k0	0.3
MRAH-G1d/MRAL-k0 (standard)	1

ligand will be used in commercial production, and thus point mutations for SuRe™ ligand binding were introduced into CDR2 of the humanized anti-CD3 antibody H-chain variants, TR01H082-E2702GsKsc and TR01H084-E2702GsKsc (SEQ ID NO:398 and 399), to prepare modified antibodies. The binding ability of these modified antibodies to the SuRe™ ligand was determined by the method described in Reference Example 5. As a result, amino acids at positions 19, 57, and 59 by Kabat numbering were found to contribute to the maintenance of the affinity for CD38 and to have influence on the SuRe™ ligand-binding ability, and antibodies with a reduced SuRe™ ligand-binding ability in comparison to that of TR01H082-E2702GsKsc/L0011-k0 (SEQ ID NOs: 398 and 410) or TR01H084-E2702GsKsc/L0011-k0 (SEQ ID NOs: 399 and 410) were obtained (Table 12).

TABLE 12

Antibody name	SuRe™ binding (RU)	Mutation sites based on CE115HA000
TR01H084-E2702GsKsc/L0011-k0	5065.8	R16G/A52aD/N53Q/D72A/L78I/G98A/Y100G/A102I
TR01H082-E2702GsKsc/L0011-k0	4469.2	V11L/A52aD/N53Q/G98A/Y100G
TR01H090-E2702GsKsc/L0011-k0	3606.3	V11L/R16G/A52aD/N53Q/G98A/Y100G
TR01H093-E2702GsKsc/L0011-k0	2459.7	V11L/A52aD/N53Q/K64Q/G98A/Y100G
TR01H094-E2702GsKsc/L0011-k0	2351.9	V11L/A52aD/N53Q/K64S/G98A/Y100G
TR01H114-E2702GsKsc/L0011-k0	1485.5	R16G/A52aD/N53Q/T57S/D72A/L78I/G98A/Y100G/A102I
TR01H092-E2702GsKsc/L0011-k0	1159.5	V11L/A52aD/N53Q/K64A/G98A/Y100G
TR01H100-E2702GsKsc/L0011-k0	383.0	V11L/A52aD/N53Q/T57S/G98A/Y100G
TR01H111-E2702GsKsc/L0011-k0	50.7	R16G/R19K/A52aD/N53Q/D72A/L78I/G98A/Y100G/A102I
TR01H110-E2702GsKsc/L0011-k0	29.5	R19K/A52aD/N53Q/G98A/Y100G
TR01H091-E2702GsKsc/L0011-k0	27.5	V11L/R19K/A52aD/N53Q/G98A/Y100G
TR01H091-E2702GsKsc/L0011-k0	15.0	V11L/R19K/A52aD/N53Q/G98A/Y100G
TR01H112-E2702GsKsc/L0011-k0	8.8	R16G/A52aD/N53Q/T57Q/D72A/L78I/G98A/Y100G/A102I
TR01H113-E2702GsKsc/L0011-k0	7.0	R16G/A52aD/N53Q/Y59V/D72A/L78I/G98A/Y100G/A102I
TR01H096-E2702GsKsc/L0011-k0	2.7	V11L/A52aD/N53Q/T57G/G98A/Y100G
TR01H109-E2702GsKsc/L0011-k0	2.2	V11L/A52aD/N53Q/Y59V/G98A/Y100G
TR01H098-E2702GsKsc/L0011-k0	1.6	V11L/A52aD/N53Q/T57P/G98A/Y100G
TR01H107-E2702GsKsc/L0011-k0	1.4	V11L/A52aD/N53Q/Y59Q/G98A/Y100G
TR01H103-E2702GsKsc/L0011-k0	1.4	V11L/A52aD/N53Q/Y59G/G98A/Y100G
TR01H104-E2702GsKsc/L0011-k0	1.0	V11L/A52aD/N53Q/Y59I/G98A/Y100G
TR01H105-E2702GsKsc/L0011-k0	0.8	V11L/A52aD/N53Q/Y59L/G98A/Y100G
TR01H099-E2702GsKsc/L0011-k0	0.6	V11L/A52aD/N53Q/T57Q/G98A/Y100G
TR01H102-E2702GsKsc/L0011-k0	0.5	V11L/A52aD/N53Q/Y59F/G98A/Y100G
TR01H101-E2702GsKsc/L0011-k0	0.5	V11L/A52aD/N53Q/T57V/G98A/Y100G
TR01H108-E2702GsKsc/L0011-k0	0.4	V11L/A52aD/N53Q/Y59T/G98A/Y100G
TR01H097-E2702GsKsc/L0011-k0	0.1	V11L/A52aD/N53Q/T57L/G98A/Y100G
TR01H106-E2702GsKsc/L0011-k0	0.0	V11L/A52aD/N53Q/Y59P/G98A/Y100G
TR01H095-E2702GsKsc/L0011-k0	-0.2	V11L/A52aD/N53Q/T57F/G98A/Y100G

(3-5) Modifying the Binding Ability to the SuRe™ Ligand by Point Mutations

[0544] An example where the binding of an antibody to Protein A depends on its variable region sequence (VH3) is known (J Biomol Tech. 2011 July; 22 (2): 50-2). In the Protein A purification of the humanized anti-human CD38 chain and anti-human GPC3 bispecific antibody, removal of the homomeric anti-CD3 antibody is important for suppressing non-specific reactions via CD3. Therefore, it is considered desirable to suppress the binding of the homomeric anti-CD3 antibody to Protein A. Presumably, the SuRe™

(3-6) Production of Optimized Bispecific Antibodies by Combining Point Mutations that Lead to Improvement of Various Properties

[0545] Optimized modified antibodies can be produced by combining the point mutations that lead to improvement of various properties as described in Examples 3-1 to 3-5. As examples of such modified antibodies, the antibodies described in Table 13 were produced, and they were subjected to the T-cell-dependent cellular cytotoxicity (TDCC) evaluation using methods similar to those of Example 1. The results are shown in FIGS. 4 to 9. As a result, optimized humanized anti-human CD38 chain and anti-human GPC3 bispecific antibodies showing a T-cell-dependent cellular

cytotoxicity equivalent to or greater than that of GPC3_ERY22_rCE115 prior to humanization were obtained.

TABLE 13

Sample number in TDCC assay	Antibody name
1	GPC3_ERY22_CE115 (rCE115H-E22Hh/rCE115L-k0/GL4-E22Hk/H0000-E22L)
2	GPC3_ERY27 (hCE115HA-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L)
3	CE115HA251-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L
4	CE115HA236-E22Hh/GLS3108-k0/GL4-E22Hk/H0000-E22L
5	TR01H002-E22Hh/GLS3108-k0/GL4-E22Hk/H0000-E22L
6	CE115HA122-E22Hh/L0000-k0/GL4-E22Hk/H0000-E22L
7	rCE115H-E22Hh/rCE115L-k0/L0000-E22Hk/H0610-E22L
8	rCE115H-E22Hh/rCE115L-k0/GL4-E22Hk/H0610-E22L
13	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0000-k0
14	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0201-k0
15	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0203-k0
16	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0204-k0
17	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0206-k0
18	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0208-k0
19	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0209-k0
20	TR01H040-E2702GsKsc/H0610-E2704sEpsc/L0211-k0
21	rCE115H-E2702GsKsc/H0610-E2704sEpsc/L0000-k0
22	TR01H061-E2702GsKsc/H0610-E2704sEpsc/L0000-k0
23	TR01H068-E2702GsKsc/H0610-E2704sEpsc/L0000-k0
24	TR01H071-E2702GsKsc/H0610-E2704sEpsc/L0000-k0
25	TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0201-k0
26	TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0212-k0
27	TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0222-k0
28	TR01H067-E2702GsKsc/GCH054-E2704sEpsc/L0000-k0
29	TR01H082-E2702GsKsc/GCH094-E2704sEpsc/L0201-k0
30	TR01H082-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0
31	TR01H084-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0
32	TR01H084-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0
33	TR01H082-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0
34	TR01H109-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0
35	TR01H109-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0
36	TR01H113-E2702GsKsc/GCH094-E2704sEpsc/L0011-k0
37	TR01H113-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0
38	TR01H113-E2702GsKsc/GCH065-E2704sEpsc/L0011-k0

[0546] Examples 3-1 to 3-6 showed that the following amino acid residues, for example, are important for maintaining the properties of the optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies showing a T-cell-dependent cellular cytotoxicity equivalent to or greater than that of GPC3_ERY22_rCE115 prior to humanization.

[0547] In anti-human CD38 chain antibodies, the examples are Leu at position 11, Gly at position 16, Asp at position 52a, Gln at position 53, Ala at position 72, Ile at position 78, Ala at position 98, Gly at position 100, and Ile at position 102. In anti-human GPC3 antibodies, the examples are Thr at position 19, Glu at position 43, Gly at position 52a, Pro or Glu at position 53, Pro at position 55, and Glu at position 61. Furthermore, in common antibody L chains, the examples are Pro at position 25, Pro at position 27a, Pro at position 27b, Ile at position 33, Gln at position 34, Arg or Trp at position 56, and Tyr at position 89. (All positions are indicated by Kabat numbering).

[Example 4] Evaluation of the In Vivo Efficacy

[0548] Some of the above-described antibodies were evaluated for their in vivo efficacy using tumor-bearing models.

[0549] Evaluation of the in vivo efficacy was carried out on representative antibodies from among those shown in Table 13, which have been confirmed to have cytotoxic activities from the in vitro assay described in Example 3-6. In the in vivo efficacy evaluation, any influence caused by differences in the microenvironment due to tumor aggregate formation on the evaluation results was taken into consideration. Therefore, two types of human cancer cell lines having different sensitivities to the antibody drug efficacy, i.e., PC-10 and NCI-H446, were used for the evaluation, even though the GPC3 expression levels of these cell lines were nearly equal. The cell lines were transplanted into the NOD scid mice, and the NOD scid mice with confirmed tumor formation were subjected to transplantation of T cells grown by in vitro culturing of human PBMCs. The mice (referred to as a T-cell injected model) were treated by administration of the optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies.

[0550] More specifically, in drug efficacy tests of the optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies using the PC-10 T-cell injected model, the tests below were performed. T cells were expansively cultured using PBMCs separated from blood collected from healthy volunteers and T cell activation/expansion kit/human (MACS Miltenyi biotec). The human cancer cell line PC-10 (1×10^7 cells) was mixed with Matrigel™ Basement Membrane Matrix (BD), and transplanted to the inguinal subcutaneous region of NOD scid mice (CLEA Japan, female, 6W). The day of transplantation was defined as day 0. On the day before transplantation, the anti-asialo-GMI antibody (Wako Pure Chemicals) was administered intraperitoneally to the mice at 0.2 mg/mouse. On days 13 to 15 after the transplantation, the mice were separated into groups according to their body weight and tumor size, and the anti-asialo-GM1 antibody was administered again intraperitoneally to the mice at 0.2 mg/mouse. On the following day, T cells obtained by the aforementioned expansive culturing were transplanted intraperitoneally at 3×10^7 cells/mouse. Four hours after T-cell transplantation, the optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies were administered intravenously through the caudate vein at 1 mg/kg. The optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies were administered only once.

[0551] As a result, anti-tumor activities were more clearly observed in the optimized anti-human CD3& chain and anti-human GPC3 bispecific antibody-administered group than in the solvent-administered group (FIG. 10A and FIG. 10B).

[0552] Drug efficacy tests for the optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies on the NCI-H446 T-cell injected model were performed by similar methods. The optimized anti-human CD38 chain and anti-human GPC3 bispecific antibodies were administered once intravenously through the caudate vein at 5 mg/kg against NCI-H446.

[0553] As a result, anti-tumor activities were more clearly observed in the optimized anti-human CD3& chain and anti-human GPC3 bispecific antibody-administered group than in the solvent-administered group (FIG. 11A and FIG. 11B).

REFERENCE EXAMPLES

[Reference Example 1] Production of Antibody Expression Vectors, and Antibody Expression and Purification

[0554] Amino acid substitutions were introduced by methods known to those skilled in the art such as using the QuikChange Site-Directed Mutagenesis Kit (Stratagene), PCR, or the In-fusion Advantage PCR cloning kit (TAKARA) to construct expression vectors. Nucleotide sequences of the obtained expression vectors were determined by a method known to those skilled in the art. The produced plasmids were transiently introduced into cells of the human embryonic kidney cancer-derived cell line HEK293H (Invitrogen) or FreeStyle293 (Invitrogen) to express antibodies. From the obtained culture supernatants, antibodies were purified using the rProtein A Sepharose™ Fast Flow (GE Healthcare) by a method known to those skilled in the art. Absorbance at 280 nm of the purified antibody solutions was measured using a spectrophotometer, and antibody concentrations were calculated from the determined values using an absorption coefficient calculated by the PACE method (Protein Science 1995; 4:2411-2423).

[Reference Example 2] the ADCC Activity of Each Test Antibody Using Human Peripheral Blood Mononuclear Cells as the Effector Cell

[0555] The ADCC activity of each test antibody was determined according to the method below.

[0556] Human peripheral blood mononuclear cells (hereinafter referred to as human PBMC) were used as the effector cell to measure the ADCC activity of each test antibody as below.

(1) Preparation of a Human PBMC Solution

[0557] From a healthy volunteer (adult male) of Chugai Pharmaceutical Co. Ltd., 50 mL of peripheral blood was collected using a syringe preloaded with 200 µL of a 1000 unit/mL heparin solution (Novo-Heparin Injection 5000 units, Novo Nordisk). The peripheral blood was diluted two-fold with PBS (-), divided into four aliquots, and added into a Leucosep lymphocyte separation tube (Greiner Bio-one) that had been loaded with 15 mL of Ficoll-Paque PLUS and subjected to centrifugation in advance. This separation tube containing aliquots of peripheral blood was centrifuged at 2150 rpm for ten minutes at room temperature, and then the mononuclear cell fraction was collected. The cells in each fraction were washed once with Dulbecco's Modified Eagle's Medium (SIGMA) containing 10% FBS (hereinafter referred to as 10% FBS/D-MEM) and then suspended in 10% FBS/D-MEM at a cell density of 5×10^6 cells/mL. After incubation in an incubator at 37° C. for one hour, the cells were washed once with 10% FBS/D-MEM, and the cells were suspended in 10% FBS/D-MEM to produce a cell density of 2×10^5 cells/mL. The cell suspension was subjected to the experiment below as the target cell.

(2) Chromium Release Assay (ADCC Activity)

[0558] The ADCC activity was evaluated from the specific chromium release rate according to the chromium release method. First, antibody solutions prepared at each concentration (0, 0.004, 0.04, 0.4, 4, and 40 µg/mL) were added to a 96-well U-bottomed plate at 50 µL per well. Next, the

target cells were seeded at 50 µL per well (1×10^4 cells/well), and this was allowed to stand at room temperature for 15 minutes. The human PBMC solution prepared in (1) was added at 100 µL per well (5×10^5 cells/well), and the plate was left to stand in a 5% carbon dioxide gas incubator at 37° C. for four hours, followed by centrifugation. The radioactivity of 100 µL of culture supernatant in each well of the plate was measured using a gamma counter. The specific chromium release rate was determined based on the following equation:

$$\text{Specific chromium release rate (\%)} = \frac{(A-C)}{(B-C)} \times 100$$

[0559] In this equation, A represents the mean value of radioactivity (cpm) of 100 µL of culture supernatant in each well; B represents the mean value of radioactivity (cpm) of 100 µL of culture supernatant in the well where 100 µL of a 2% aqueous NP-40 solution (Nonidet P-40, Nacalai Tesque) and 50 µL of 10% FBS/D-MEM had been added to the target cells; and C represents the mean value of radioactivity (cpm) of 100 µL of culture supernatant in the well where 150 µL of 10% FBS/D-MEM had been added to the target cells. The examinations were performed in triplicate and the mean values and standard deviations of the specific chromium release rates (%) in the above-mentioned examination reflecting the ADCC activity were calculated for each of the test antibodies.

[Reference Example 3] Assessment of Tm of the Modified Antibodies by Differential Scanning Fluorimetry

[0560] In this examination, the Tm (thermal denaturation temperature) value of the modified antibodies was assessed by differential scanning fluorimetry using Rotor-Gene Q (QIAGEN). It has been reported that this method has a favorable correlation with Tm assessment using a differential scanning calorimeter widely known as a method for evaluating thermal stability of antibodies (Journal of Pharmaceutical Science 2010; 4:1707-1720).

[0561] The 5000×-concentrated SYPRO™ orange (Molecular Probes) was diluted with PBS (Sigma), and then admixed with the antibody solutions to prepare measurement samples. Twenty-µL aliquots of each sample were placed into measurement tubes, and the temperature was increased from 30° C. to 99° C. at a temperature elevation rate of 240° C./hr. Changes in fluorescence accompanying the temperature elevation were detected at 470 nm (excitation wavelength)/555 nm (fluorescence wavelength).

[0562] The data were analyzed using the Rotor-Gene Q Series software (QIAGEN) to calculate the temperature at which fluorescence transition was observed, and this temperature was defined as the Tm.

[Reference Example 4] Assessment of the ECM-Binding Ability

[0563] The assessment was carried out according to the method described in WO 2012093704. Specifically, BD Matrigel (BD Biosciences, #356237) was prepared at 2 mg/mL using TBS (Takara, #T903), and this was dispensed into a 96-well measurement plate (Meso Scale Discovery, #L15XB-3 (High Bind)) at 5 µL per well and then allowed to stand overnight in a cool place. Then, 150 µL of an ECL blocking buffer (PBS containing 0.05% Tween20, 0.5% BSA, and 0.01% sodium azide) was dispensed into each

well of the plate, and this was allowed to stand at room temperature for two hours or more.

[0564] A goat anti-human IgG (Y) (Invitrogen, #628400) was ruthenium-labeled with MSD SULFO-TAG NHS Ester (Meso Scale Discovery, #R91AN-2) by following the attached instructions. This was diluted in an ECL dilution buffer (PBS containing 0.01% Tween20, 0.1% BSA, and 0.01% sodium azide) to have a final concentration of 2 µg/mL. Furthermore, the standard antibody and the test antibodies were diluted in PBS-T (PBS containing 0.05% Tween 20 and 0.01% sodium azide) to have a final concentration of 3 µg/mL.

[0565] To a 96-well reaction plate (Thermo scientific, Nunc #145399), 10 µL of the ECL dilution buffer, 20 µL of the standard antibody and test antibody (3 µg/mL), and 30 µL of the ruthenium-labeled antibody (2 µg/mL) were added sequentially, and this was allowed to react for one hour at room temperature with stirring in the dark.

[0566] The ECL blocking buffer was removed from the 96-well measurement plate by tilting, 50 µL of the sample solution from the 96-well reaction plate was added, and this was allowed to stand in the dark at room temperature for one hour. This was followed by removal of the sample solution from the 96-well measurement plate by tilting, and immediately after addition of 150 µL of 2×T buffer (4×MSD Read Buffer T (Meso Scale Discovery) diluted two-fold using the ECL dilution buffer), ECL measurements were taken. SEC-TOR Imager 2400 (Meso Scale Discovery) was used for taking the measurements.

[0567] Analyses were carried out by dividing the fluorescence intensity of the test antibody by the fluorescence intensity of the standard antibody to calculate and compare the intensities by defining the value for the standard antibody to be 1.

[Reference Example 5] Assessment of the SuRe™ Ligand-Binding Ability

[0568] The ability to bind to the SuRe™ ligand was assessed by using Biacore™-T200 (GE Healthcare Japan). HBS-EP+ (GE Healthcare Japan) was used for the running buffer, and an amine coupling kit (GE Healthcare Japan) was used to covalently bind the Mab Select SuRe™ Ligand (GE Healthcare Japan) to the CM5 chip (carboxymethyl dextran-coated chip). The antibody used as the analyte was prepared at 5 µg/mL using HBS-EP+. Measurements were carried out by first injecting the 5-µg/mL antibody solution at a flow rate of 10 µL/min for 3 minutes, then switching to HBS-EP+, and measuring the response (RU) after allowing the flow to continue for 0.5 minutes. After completion of the measurements, the sensor chip was regenerated by washing with 10 mM Gly-HCl at pH 1.5. For the control flow cell, a similar experiment was performed without covalent bonding of the ligand to the chip, and the affinity for the SuRe™ ligand was analyzed by taking the difference between the responses (RU).

[0569] Sequences corresponding to the SEQ ID NOs mentioned herein are shown in the Table below.

TABLE 14

SEQ ID NO:	Name
1	GPC3 nucleotide sequence (NM_001164617.1)
2	GPC3 amino acid sequence (NP_001158089.1)
3	Signal sequence
4	T cell receptor α-chain peptide (CAA26636.1)
5	T cell receptor β-chain peptide (C25777)
6	T cell receptor γ1-chain peptide (A26659)
7	T cell receptor γ2-chain peptide (AAB63312.1)
8	T cell receptor δ-chain peptide (AAA61033.1)
9	CD3 γ-chain nucleotide (NM_000073.2)
10	CD3 δ-chain nucleotide (NM_000732.4)
11	CD3 ε-chain nucleotide (NM_000733.3)
12	CD3 γ-chain peptide (NP_000064.1)
13	CD3 δ-chain peptide (NP_000723.1)
14	CD3 ε-chain peptide (NP_000724.1)
15-22	Peptide linker
23	Human Cγ1
24	Human Cγ2
25	Human Cγ3
26	Human Cγ4
27	FcγRI nucleotide (NM_000566.3)
28	FcγRI peptide (NP_000557.1)
29	FcγRIIA nucleotide (BC020823.1)
30	FcγRIIA peptide (AAH20823.1)
31	FcγRIIB nucleotide (BC146678.1)
32	FcγRIIB peptide (AAI46679.1)
33	FcγRIIIA nucleotide (BC033678.1)
34	FcγRIIIA peptide (AAH33678.1)
35	FcγRIIIB nucleotide (BC128562.1)
36	FcγRIIIB peptide (AAI28563.1)
37	Fc region (addition of A to the N terminus of RefSeq accession number AAC82527.1)
38	Fc region (addition of A to the N terminus of RefSeq accession number AAB59393.1)
39	Fc region (addition of A to the N terminus of RefSeq accession number AAB59394.1)
40	H0000, GPC3 H-chain variable region
41	GL4, GPC3 L-chain variable region
42	rCE115H, CE115 H-chain variable region
43	rCE115L, CE115 L-chain variable region
44	G1dh
45	ERY22_Hk

TABLE 14-continued

SEQ ID NO: Name

46	ERY22_Hh
47	GL4-ERY22_Hk
48	H0000-ERY22_L
49	rCE115H-ERY22_Hh
50	rCE115L-k0
51	hCE115HL (Heavy chain of humanized CE115)
52	hCE115HA (Heavy chain of humanized CE115)
53	L0000 (Light chain of humanized CE115)
54	H0000-ERY27_HK
55	hCE115HA-ERY27_HE
56	L0000-k0
57	E22Hh
58	E22Hk
59	Hi-Kn010G3
60	E2702GsKsc
61	E2704sEpsc
62	E2702sKsc
63	k0
64	CE115HA177
65	CE115HA178
66	CE115HA179
67	CE115HA180
68	hCE115HAAa
69	TR01H006
70	TR01H007
71	TR01H008
72	TR01H009
73	TR01H010
74	TR01H011
75	TR01H012
76	TR01H013
77	TR01H014
78	TR01H015
79	TR01H016
80	TR01H017
81	TR01H018
82	TR01H019
83	TR01H020
84	TR01H021
85	TR01H022
86	TR01H023
87	TR01H024
88	TR01H025
89	TR01H026
90	TR01H027
91	TR01H028
92	TR01H029
93	TR01H030
94	TR01H031
95	TR01H032
96	TR01H033
97	TR01H034
98	TR01H035
99	TR01H036
100	TR01H037
101	TR01H038
102	TR01H039
103	TR01H040
104	TR01H041
105	TR01H042
106	TR01H043
107	TR01H044
108	TR01H045
109	TR01H046
110	TR01H047
111	TR01H048
112	TR01H049
113	TR01H050
114	TR01H051
115	TR01H052
116	TR01H053
117	TR01H054
118	TR01H055
119	TR01H056
120	TR01H057

TABLE 14-continued

 SEQ ID NO: Name

121	TR01H058
122	TR01H061
123	TR01H062
124	TR01H063
125	TR01H064
126	TR01H065
127	TR01H066
128	TR01H067
129	TR01H068
130	TR01H069
131	TR01H070
132	TR01H071
133	TR01H072
134	TR01H073
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139	TR01H079
140	TR01H080
141	TR01H081
142	TR01H082
143	TR01H083
144	TR01H084
145	TR01H090
146	TR01H091
147	TR01H092
148	TR01H093
149	TR01H094
150	TR01H095
151	TR01H096
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154	TR01H099
155	TR01H100
156	TR01H101
157	TR01H102
158	TR01H103
159	TR01H104
160	TR01H105
161	TR01H106
162	TR01H107
163	TR01H108
164	TR01H109
165	TR01H110
166	TR01H111
167	TR01H112
168	TR01H113
169	TR01H114
170	GCH003
171	GCH005
172	GCH006
173	GCH007
174	GCH008
175	GCH010
176	GCH012
177	GCH013
178	GCH014
179	GCH015
180	GCH016
181	GCH019
182	GCH022
183	GCH023
184	GCH025
185	GCH026
186	GCH027
187	GCH029
188	GCH032
189	GCH034
190	GCH035
191	GCH039
192	GCH040
193	GCH042
194	GCH043
195	GCH045

TABLE 14-continued

 SEQ ID NO: Name

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198	GCH055
199	GCH056
200	GCH057
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203	GCH061
204	GCH062
205	GCH064
206	GCH065
207	GCH066
208	GCH067
209	GCH068
210	GCH073
211	GCH094
212	GCH098
213	GCH099
214	GCH100
215	H0610
216	L0000vk1
217	L0002
218	L0003
219	L0006
220	L0007
221	L0008
222	L0009
223	L0011
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225	L0013
226	L0014
227	L0015
228	L0016
229	L0032
230	L0038
231	L0039
232	L0041
233	L0042
234	L0043
235	L0044
236	L0045
237	L0046
238	L0047
239	L0062
240	L0063
241	L0064
242	L0065
243	L0066
244	L0069
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249	L0089
250	L0090
251	L0091
252	L0093
253	L0104
254	L0106
255	L0107
256	L0109
257	L0113
258	L0115
259	L0117
260	L0120
261	L0122
262	L0123
263	L0124
264	L0125
265	L0126
266	L0127
267	L0129
268	L0132
269	L0134
270	L0136

TABLE 14-continued

SEQ ID NO: Name

271	L0137
272	L0138
273	L0139
274	L0140
275	L0141
276	L0143
277	L0144
278	L0145
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288	L0161
289	L0163
290	L0167
291	L0168
292	L0173
293	L0175
294	L0180
295	L0181
296	L0186
297	L0187
298	L0200
299	L0201
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311	L0213
312	L0214
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316	L0218
317	L0219
318	L0220
319	L0222
320	L0223
321	L0224
322	L0226
323	L0227
324	L0228
325	L0229
326	L0230
327	L0231
328	L0232
329	L0233
330	L0234
331	L0235
332	L0236
333	L0237
334	L0238
335	L0239
336	L0240
337	L0241
338	L0242
339	L0243
340	L0246
341	L0247
342	L0248
343	L0249
344	L0250
345	L0258

TABLE 14-continued

SEQ ID NO: Name

346	L0259
347	L0260
348	L0261
349	L0262
350	L0263
351	L0264
352	L0265
353	L0266
354	L0267
355	L0268
356	L0269
357	L0270
358	L0271
359	L0272
360	TR01L001
361	TR01L002
362	TR01L003
363	TR01L004
364	TR01L005
365	TR01L006
366	TR01L007
367	TR01L008
368	TR01L009
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371	TR01L012
372	TR01L013
373	TR01L015
374	TR01L016
375	TR01L017
376	TR01L018
377	TR01L019
378	TR01L020
379	TR01L023
380	TR01L024
381	CE115HA122-E22Hh
382	CE115HA236-E22Hh
383	CE115HA251-E22Hh
384	GCH054-E2704sEpsc
385	GCH065-E2704sEpsc
386	GCH094-E2704sEpsc
387	H0610-E2704sEpsc
388	hCE115HA-E22Hh
389	rCE115H-E22Hh
390	rCE115H-E2702GsKsc
391	TR01H002-E22Hh
392	TR01H015-E22Hh
393	TR01H040-E2702GsKsc
394	TR01H061-E2702GsKsc
395	TR01H067-E2702GsKsc
396	TR01H068-E2702GsKsc
397	TR01H071-E2702GsKsc
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399	TR01H084-E2702GsKsc
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405	H0000-E22L
406	H0610-E22L
407	rCE115L-k0
408	GLS3108-k0
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411	L0201-k0
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413	L0204-k0
414	L0206-k0
415	L0208-k0
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419	L0222-k0
420	TR01H001

TABLE 14-continued

SEQ ID NO: Name

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422	TR01H003
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424	rCE115H
425	CE115HA121
426	CE115HA122
427	CE115HA124
428	CE115HA192
429	CE115HA236
430	CE115HA251
431	CE115HA252
432	E22L

INDUSTRIAL APPLICABILITY

[0570] The present invention provides novel multispecific antigen-binding molecules that maintain the strong anti-tumor activity possessed by BiTE and the excellent safety property of not inducing a cytokine storm or such independently from cancer antigen, and also have long half-lives in blood. Cytotoxicity-inducing agents that comprise an anti-

gen-binding molecule of the present invention as an active ingredient can target glycan 3-expressing cells and tumor tissues containing these cells and induce cell injury. Administration of a multispecific antigen-binding molecule of the present invention to patients makes it possible to have a desirable treatment which not only has a high level of safety but also a reduced physical burden, and is highly convenient.

SEQUENCE LISTING

Sequence total quantity: 432
 SEQ ID NO: 1 moltype = DNA length = 2398
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 source 1..2398
 mol_type = other DNA
 organism = Homo sapiens
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FEATURE          Location/Qualifiers
source           1..603
mol_type = protein
organism = Homo sapiens
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DLQVCLPKGP TCCSRKMEEK YQLTARLNME QLQSASMEL KFLIIQNAAV FQEAFEIVVR 120
HAKNYTNAMF KNNYPSLTPQ AFEFVGEGFT DVSLYILGSD INVDDMVNEL FDSDLFPVITY 180
QLMPNGLPDS ALDINECLRG ARRDLKVFGN FPKLIIMTQVS KSLQVTRIFL QALNLGIEVI 240
NTTDHFKFSK DCGRMLTRMV YCSYQCGLMM VKPCGGYCNV VMQGCMAGVV EIDKYWREYI 300
LSLEELVNGM YRIYDMENVL LGLFSTIHD S I Q Y V Q K N A G K L W H F K P I F F L 360
CIGLDLQIGK LCAHSQROQ RSAVYPEDLF IDKKVLVKAH VEHEETLSSR RRELIQKLKS 420
FISFYSALPG YICSHSPVAE ND TLCWNGQE LVERYSQKAA RNMGNQFNL HELKMKGPEP 480
VVSQIIDKLK HINQLLRTMS MPKGRVLDKN LDEEGFESGD CGDDEDECIG GSGDGMIVKV 540
NQLRFLAELA YDLDVDDAPG NSQQATPKDN EISTFHNLGN VHSPLKLTS MAISVVCF 600
LVH                                         603

SEQ ID NO: 3      moltype = AA length = 19
FEATURE          Location/Qualifiers
REGION           1..19
note = An artificially generated sequence
source           1..19
mol_type = protein
organism = synthetic construct
SEQUENCE: 3
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SEQ ID NO: 4      moltype = AA length = 141
FEATURE          Location/Qualifiers
source           1..141
mol_type = protein
organism = Homo sapiens
SEQUENCE: 4
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SAVAWSNKSDFACANAFNNS IIPEPDTEFFPS PESSIONCDVKLV EKSFETDTNL NFQNLNSVIGF 120
RILLLKVGAF NLLMTLRLWS S                                         141

SEQ ID NO: 5      moltype = AA length = 179
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mol_type = protein
organism = Homo sapiens
SEQUENCE: 5
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VSAEAWGRAD CGFTSESYQQ GVLSATILYE ILLGKATLYA VLVSALVLM MVKRKDSRG 179

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organism = Homo sapiens
SEQUENCE: 6
DKQLDADVSP KPTIFLPSIA ETKLQKAGTY LCLLEKFFF D VIKIHWQEKK NTILGSQEG 60
NTMKTNNDYTM KFSWLTVPEK SLDEKEHRCIV RHENNKNQVD QEIIFFPIKT DVITMDPKDN 120
CSKDANDTLL LQLTNTSAYY MYLLLLLKS VYFAIITCCL LRRTAFCCNG EKS 173

SEQ ID NO: 7      moltype = AA length = 204
FEATURE          Location/Qualifiers
source           1..204
mol_type = protein
organism = Homo sapiens
SEQUENCE: 7
KQLDADVSPK PTIFLPSIAE TKLQKAGTYL CLLEKFFF D I K I H W Q E K K S N T I L G S Q E G N 60
TMKTNNDYMK FSWLTVPEES LDKEHRCIVR HENNKNGIDQ E I I F P P I K T D V T T V D P K D S Y 120
SKDANDTTLV DPKYNSKDA NDVITMDPKD NW SKDANDTLL Q L Q L T N T S A Y Y M Y L L L L K S 180
VYFAIITCCL LLGRTAFCCN GEKS                                         204

SEQ ID NO: 8      moltype = AA length = 177
FEATURE          Location/Qualifiers
source           1..177
mol_type = protein

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organism = Homo sapiens

SEQUENCE: 8

PSYTGGYADK LIFGKGKTRVT VEPRSQPHTK PSVFVVMKNGT NVACLVKEFY PKDIRNLVS 60
 SKKITEFDPA IVISPGSKYVN AVKLGKYEDS NSVTCVQHD NKTVHSTDVE VKTDSTDHVVK 120
 PKETENTKQP SKSCHKPKAI VHTEKVNMSM LTVLGLRMLF AKTVAVNFLL TAKLFFL 177

SEQ ID NO: 9 moltype = DNA length = 1311
 FEATURE Location/Qualifiers
 source 1..1311
 mol_type = other DNA
 organism = Homo sapiens

SEQUENCE: 9

agtcttagctt ctgcacaggc tggctggctg gctggctgt aagggtcgat ccacgctttt 60
 agccggaggac agagactgac atggaaacagg ggaaggccct ggctgtccat atccctggcta 120
 tcattcttc tcaaggtaat ttggcccaatg caatcaaagg aaaccacttg gttaagggtt 180
 atgactatca agaaagatctc tcggacttc tgacttgcgat tgccagaacgg aaaaatata 240
 catggtttaa agatgggaatg atgatcggtt ctcctaaatgc agataaaaaa aatggaaatc 300
 ttggaaagtaat tgccaaaggac ctcggaggaa tgatcaggatg taaaaggatca cagaacaatg 360
 caaaactactt ccaagtgtat tacagaatgt gtcagaactg cattgaacta aatgcagcca 420
 ccatacttgc ttggatcttgcgatc tcaaggatgt cgttcgttgc gttgggtt 480
 acttcattgc tggacaggat ggaggctccg agtcgagatc ttccagacaag cagacttgt 540
 tgcccaatgtt ccacgttctca cagccccctca aggatcgaa agatgaccatc agccggcc 600
 ttcaaggaaat ccagttgagg aggaatttgcgatc ctcaggactc agagtagtcc aggtgttctc 660
 ctccatttc tggcccaatgtt ccaaggatcat gcattttggat aagctcttgcgatc 720
 ttccggccatc aatcttagact tcggatccc agagatgaca aatggaaag aaggccatc 780
 agagacaaat tgggggttcaat taaaatataa aaaaatataa aatccatgttgc 840
 agccggccacctt attggggaaaat attgtaaaatg aaaaatgaaaatcaata accccctggat 900
 ttgtatcataat ttttttttttgcgatc ttgtatcataat ttttttttttgcgatc 960
 atgggtcaatcatttgcgatc aatcttagact tcggatccc tccacccggatc cccatgttgc 1020
 ctggcccttgcgatc ttccggccatc aatcttagact tcggatccc tccacccggatc 1080
 agacagtctg gctctgtcactt ccaggctgaa atgcgttgcgatc accatctggatc 1140
 cctctgcctc ctgggttcaat gcgatttgcgatc tggctcggatc tccggccatc 1200
 aggccacacac taccacacactt ggctatcatttttgcgatc ttgtatcataat ttttttttgc 1260
 ttgttgcgatc gctggctcgcgatc aactctgtcactt ccaggctgaaatc 1311

SEQ ID NO: 10 moltype = DNA length = 771
 FEATURE Location/Qualifiers
 source 1..771
 mol_type = other DNA
 organism = Homo sapiens

SEQUENCE: 10

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 ggtggggggat ggcacgttcc acccaggctg atagttcggtt gacccgggtt tatctactgg 120
 atgagttcgat ctggggatgtt gaacatagca cgtttcttc tggctggatc ttggcttaccc 180
 ttctctcgatc agtgagcccccc ttcaggatcat ctatagagga acttgaggac agagtgtttt 240
 tgaatggcata taccaggatcat acatgggttag agggaaacggt gggaaacactg ctcccgacata 300
 ttacaaggactt gggacccatggaa aacacccatcc gggacccacgg aggaataatgat aggtgtaaatg 360
 ggacagatatacataaggac aaaaatctca cccgttcaatgc tccatataatgc atgtggccaga 420
 gctgtgtggat gctggatccat gccaccgtgg ctggcatcat tgcactgtat gtcattggca 480
 ctctgtcttc tggctttggat gtcatttcgtt tgctggatca tggatgttgcgatc 540
 gggctggccatc cacacaatgttgcgatc ttgttggatca atggacaggat cttatcgatcc ctccggatc 600
 gagatgtatc tcaatgttgcgatc ttgttggatca gaaactgggc tccggacaaatg tgaacctggat 660
 actgggttgcgatc tctagaagatc gccattacca actgttacccctt cccttcttgc tcaatgttgcgatc 720
 aatatatccctt ctggatccatc aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa a 771

SEQ ID NO: 11 moltype = DNA length = 1534
 FEATURE Location/Qualifiers
 source 1..1534
 mol_type = other DNA
 organism = Homo sapiens

SEQUENCE: 11

tattgtcaga gtccttcttgcgatc ttggcccttcttgcgatc aggaaggctg tggggacccag ctttcttcaatgc 60
 ccaggccaggat tggggcccttc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 120
 agaggcccttc tctacttcttc tggggggatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 180
 gtcctgttcaatgcgatc gaaaatgttgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 240
 gtcctgttcaatgcgatc gaaaatgttgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 300
 atcaggatgttgcgatc gtttccatgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 360
 atataaaatgcgatc tccatcttc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 420
 tggatgttgcgatc tccatcttc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 480
 agggccatc gtttccatgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 540
 ttatgttgcgatc tccatcttc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 600
 ggcaaggatgttgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 660
 tggatgttgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 720
 aaaaatccctcc acgttccatgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 780
 aaaaatccctcc acgttccatgcgatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc 840
 gggggccatc tggggccatc gtttccatgcgatc aaaaatccctcc acgttccatgcgatc tggggccatc gtttccatgcgatc 900

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cccgctggcc caggctcct ctccagttccc cctgcgactc cctgtttcct gggcttagtct 960
tggacccac gagagagaat cggttcctcg cctcatggtg aactcgccg ccacagctg 1020
atcccccgat cccttcctcc tgccttcctc gctggtaacc agtctaaaatttgcgtct 1080
tcccttcctt ttgaagcatc atcagtatgc acaccctcac agctggctg ccctttgcc 1140
aggatattta ttgtgtctat tcactccctt ccctttggat gtaacttctc cgttcagttc 1200
cccttccttc ttgcgtgtaa gttgtccccc atcccaaagt attccatcta cttttctatc 1260
ggcgtccccc ttgcagcccc tctctgggg tggactgggaaatgttgc agaggccctg 1320
cccccgttac agatcttggc cctyagccag cccctgtctc ctccctcccc caacactccc 1380
taccaacccc ctaatccccct actccctcca ccccccctcc actgttaggg actggatgg 1440
catttgcatac tccgtaaatg tgctgtgtc ctcagctgag agagaaaaaaa ataaaactgt 1500
tttggctgca agaaaaaaaaaaa aaaaaaaaaaaa aaaa 1534

SEQ ID NO: 12 moltype = AA length = 182
FEATURE Location/Qualifiers
source 1..182
mol_type = protein
organism = Homo sapiens
SEQUENCE: 12
MEQGKGLAVL ILAIIILQGT LAQSKGNHL VKVYDYQEDG SVLLTCDAEA KNITWFKDGK 60
MIGFLTEDKK KWNLGSNAKD PRGMYQCKGS QNKSKPLQVY YRMCQNIEL NAATISGFLF 120
AEIVSIFVLA VGVYFIAGQD GVRQSRASDK QTLLPNDQLY QPLKDREDDQ YSHLQGNQLR 180
RN 182

SEQ ID NO: 13 moltype = AA length = 171
FEATURE Location/Qualifiers
source 1..171
mol_type = protein
organism = Homo sapiens
SEQUENCE: 13
MEHSTFLSGL VLATLLSQVS PFKIPIEELLE DRVFVNCNTS ITWVEGTVGT LLSDITRLDL 60
GKRILDPRGI YRCNGTDIYK DKESTVQVHY RMCQSCVELD PATVAGIIVT DVIATLLLAL 120
GVPCFAGHET GRLSGAADTQ ALLRNDQVYQ PLRDRDDAQY SHLGGNWARN K 171

SEQ ID NO: 14 moltype = AA length = 207
FEATURE Location/Qualifiers
source 1..207
mol_type = protein
organism = Homo sapiens
SEQUENCE: 14
MQSGTHWRVL GLCLLSVGVW GQDGNEEMGG ITQTPYKVSI SGTTVILTCP QYPGSEILWQ 60
HNNDKNIGGDE DDKNIGSDED HLSLKEFSEL EQSGYYVCYR RGSKPEDANF YLYLRARVCE 120
NCMEMDVMSV ATIVIVDICI TGGLLLLVYY WSKNRKAKAK PVTRGAGAGG RQRGQNKER 180
PPVNPNDYEP IRKGQRDLYS GLNQRRI 207

SEQ ID NO: 15 moltype = AA length = 4
FEATURE Location/Qualifiers
REGION 1..4
source note = An artificially generated sequence
1..4
mol_type = protein
organism = synthetic construct
SEQUENCE: 15
GGGS 4

SEQ ID NO: 16 moltype = AA length = 4
FEATURE Location/Qualifiers
REGION 1..4
source note = An artificially generated sequence
1..4
mol_type = protein
organism = synthetic construct
SEQUENCE: 16
SGGG 4

SEQ ID NO: 17 moltype = AA length = 5
FEATURE Location/Qualifiers
REGION 1..5
source note = An artificially generated sequence
1..5
mol_type = protein
organism = synthetic construct
SEQUENCE: 17
GGGGS 5

SEQ ID NO: 18 moltype = AA length = 5
FEATURE Location/Qualifiers

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REGION	1..5				
source	note = An artificially generated sequence				
	1..5				
	mol_type = protein				
	organism = synthetic construct				
SEQUENCE: 18					
SGGGG					5
SEQ ID NO: 19	moltype = AA length = 6				
FEATURE	Location/Qualifiers				
REGION	1..6				
source	note = An artificially generated sequence				
	1..6				
	mol_type = protein				
	organism = synthetic construct				
SEQUENCE: 19					
GGGGGS					6
SEQ ID NO: 20	moltype = AA length = 6				
FEATURE	Location/Qualifiers				
REGION	1..6				
source	note = An artificially generated sequence				
	1..6				
	mol_type = protein				
	organism = synthetic construct				
SEQUENCE: 20					
GGGGGG					6
SEQ ID NO: 21	moltype = AA length = 7				
FEATURE	Location/Qualifiers				
REGION	1..7				
source	note = An artificially generated sequence				
	1..7				
	mol_type = protein				
	organism = synthetic construct				
SEQUENCE: 21					
GGGGGGS					7
SEQ ID NO: 22	moltype = AA length = 7				
FEATURE	Location/Qualifiers				
REGION	1..7				
source	note = An artificially generated sequence				
	1..7				
	mol_type = protein				
	organism = synthetic construct				
SEQUENCE: 22					
GGGGGGG					7
SEQ ID NO: 23	moltype = AA length = 330				
FEATURE	Location/Qualifiers				
source	1..330				
	mol_type = protein				
	organism = Homo sapiens				
SEQUENCE: 23					
ASTKGPSVFP LAPSSKSTSG GTAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS	60				
GLYSLSSVVT VPSSNFGTQT YICCNVNHKPS NTKVDKKVER KSCDKTHTCP PCPAPELLGG	120				
PSVFLFPKPK KDTLMISRTP EVTCVVVDVS HEDPEVKPNW YVDGVENVNA KTKPREEQYN	180				
STYRVSVLTL VHQNWLNGK EYKCKVSNKA LPAPIEKTIIS KAKGQPREPQ VYTLPPSRDE	240				
LTKNQVSLTC LVKGFYPSDI AVEWESNGOP ENNYKTTTPPV LDSDGSFFLY SKLTVDKSRW	300				
QQGNVFSCSV MHEALHNHYT QKSLSLSPGK	330				
SEQ ID NO: 24	moltype = AA length = 326				
FEATURE	Location/Qualifiers				
source	1..326				
	mol_type = protein				
	organism = Homo sapiens				
SEQUENCE: 24					
ASTKGPSVFP LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS	60				
GLYSLSSVVT VPSSNFGTQT YTCNVDHKPS NTKVDKTVER KCCVECPPCP APPVAGPSVF	120				
LFPPKPKDNL MISRTPEVTC VVVDVSHEDP EVQFNWYVDG VEVHNAKTKP REEQFNSTFR	180				
VVSVLTVVHQ DWLNGKEYKC KVSNKGLPAP IEKTISKTKG QPREPQVYTL PPSREEMTKN	240				
QVSLTCLVKG FYPSDIAVEW ESNGQPENNY KTPPMULDSD GSFFLYSKLT VDKSRWQQGN	300				
VFSCSVMHEA LHNHYTQKSL SLSPGK	326				
SEQ ID NO: 25	moltype = AA length = 377				
FEATURE	Location/Qualifiers				

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source          1..377
               mol_type = protein
               organism = Homo sapiens
SEQUENCE: 25
ASTKGPSVFP LAPCSRSTSG GTAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT VPSSSLGTQT YTCVNHHKPS NTKVDKRVEL KTPLGDTHTH CPRCPEPKSC 120
DTPPPCPRCP EPKSCDTPPP CPRCPEPKSC DTPPPCPRCP APELLGGPSV FLFPPPKD 180
LMISRTPEVT CVVVDVSHED PEVQFKWYWD GVEVHNNAKTK PREEQYNSTF RVVSVLTVLH 240
QDWLNGKEYK CKVSNKALPA PIEKTISKTK GQPREPQVYT LPPSREEMTK NQVSLTCLVK 300
GFYPSDIAVE WESSGOPENN YNTTPPMQLDS DGSFFFLYSKL TVDKSRWQQG NIFSCSVMHE 360
ALHNRFQKLSLSPGK                                     377

SEQ ID NO: 26          moltype = AA  length = 327
FEATURE
source          1..327
               mol_type = protein
               organism = Homo sapiens
SEQUENCE: 26
ASTKGPSVFP LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT VPSSSLGTKT YTCVNVDHKPS NTKVDKRVES KYGPPCPSCP APEFLGGPSV 120
FLFPPPKD LMISRTPEVT CVVVDVSQED PEVQFNWYWD GVEVHNNAKTK PREEQFNSTY 180
RVVSVLTVLH QDWLNGKEYK CKVSNKGKPS SIEKTISKAK GQPREPQVYT LPPSQEEMTK 240
NQVSLTCLVK GFYPSDIAVE WESNGQOPENN YKTTPPVLDs DGSFFFLYSRL TVDKSRWQEG 300
NVFSCSVMHE ALHNHYTQKS LSLSLGK                     327

SEQ ID NO: 27          moltype = DNA  length = 2268
FEATURE
source          1..2268
               mol_type = other DNA
               organism = Homo sapiens
SEQUENCE: 27
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tgacaactct gtcctttgg gttccaggat atggccaatg ggacaccacaa aaggcagtga 120
tcactttgcg ccctccatgg gtcagcgat tccaaaggatg aaccgtaaacc ttgcactgtg 180
agggtcgatc tctgcgttgg agcagctca cacatgttgc tctcaatggc acagccactc 240
agacctcgac ccccgatc aaaaatccat ctggccaggat caatgcacatg ggtgaatata 300
gggtgccagat aggtcttcgatggcgaaatg accccatata gctggaaatc cacagaggct 360
ggctactact gcaggatctcc agcagatgt tcacggaaagg agaacctctg gccttggat 420
gtcatgcgtg gaaggatataatg ctgggttata atgtgttttta ctatcgaat ggc当地ggct 480
ttaagttttt ccactggat tctaaatctca ccattctgaa aaccacatata agtcaatata 540
gcacccatcca ttgctcaggat atggaaatggc atcgctacatc atcagcggatc atatctgtca 600
ctgtgaaaga gtcatttcca gtcaggatgc tgaatgcacatc tggatcatcc ccactctgg 660
aggggtaattt ggtcaccctt agctgtgaaatc caaaatgttgc ctggcaagg ctgggttgc 720
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aataccaaat actaactgtc agaaagaaatg actctgggtt atactgttgc gaggtgcacca 840
cagaggatgg aaatgtccat aagcgccatcc ctggatgttgc gttcaatgg ctggccctcc 900
agttaccaac ttctgtctgg ttcatgttgc ttcttatctt ggcaggatggc ataatgttt 960
tagtgaacatc ttgttctgg gtgacaatataatc gtaaaatggaaatggaaaatgggggggg 1020
atttagaaat ctctttggat tctggatcatc agaaatggatc aattttccatc cttcaagaaatg 1080
acagacatcc agaagaagatc ctgaaatgtc aggaacaaaaa agaagaacatc ctgcaggaaatg 1140
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ctcaaagaaat tctttttttttaatc aatggatgttgc tggatcatc attacatatgtt 1440
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agaagatataa gacagagaaaaa cacagacaaaaa gagacacacaaatc cttccaaatgtt 1560
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acttcacttgc ggatacatatc ttggaaatgtgg tcatcaaaatgc tgacttggaaatggccatc 1680
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tcataacccc tctttttttttaatc tcttgcgttgc tggatgttgc tggatgttgc 1860
taggaaggatc gatgttgc tttttttttttaatc tcttgcgttgc tggatgttgc tggatgttgc 1920
ggactgttgc aaatggatgttgc tttttttttttaatc tcttgcgttgc tggatgttgc tggatgttgc 1980
gcctccatccatc tttttttttttaatc tcttgcgttgc tggatgttgc tggatgttgc 2040
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agctgaggatc aaaaactgttgc tttttttttttaatc tcttgcgttgc tggatgttgc tggatgttgc 2160
ggaggaggatc gacacaaaaatc aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaa 2220
aaaaaaaaaaaaatc aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa 2220

SEQ ID NO: 28          moltype = AA  length = 374
FEATURE
source          1..374
               mol_type = protein
               organism = Homo sapiens
SEQUENCE: 28

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MWFLTTLLLW	VPVDGQVDTT	KAVITLQPPW	VSVFQEETVT	LHCEVLHLPG	SSSTQWFILNG	60
TATQTQTPSY	RITSASVNDS	GEYRCQRGLS	GRSDPIQLEI	HRGWLLLQVS	SRVFTEGEPPL	120
ALRCHAWDKK	LVYNVLYYRN	KGAFKFFHWN	SNLTILKTNI	SHNGTYHCSCG	MGKHRYTSAG	180
ISVTVKELFP	APVLNASVTS	PLLEGNLVTL	SCBTKLLLQR	PGLQLYFSFY	MGSKTLRGRN	240
TSSEYQILTA	RREDSGLYWC	EAATEDGNVL	KRSPELELQV	LGLQLPTPVW	FHVLFYLAVG	300
IMPLVNTVLW	VTIRKELKRK	KKWDLEISLD	SGHEKKVISS	LQEDRHLEEE	LKCQEQQKEEQ	360
LQEGVHRKEP	QGAT					374

SEQ ID NO: 29	moltype = DNA length = 1396					
FEATURE	Location/Qualifiers					
source	1..1396					
	mol_type = other DNA					
	organism = Homo sapiens					
SEQUENCE: 29						
gggtatgacta	tggagaccca	aatgtctcg	aatgtatgtc	ccagaaaacct	gtggctgttt	60
caaccattga	cagtttgc	gctgttggt	tctgcagaca	gtcaagctgc	tccccaaag	120
gctgtgtga	aacttgagcc	cccgtggatc	aacgtgtcc	aggaggactc	tgtgtactcg	180
acatgccagg	gggtcgccag	ccctgagatc	gactccatc	agtggttcca	caatggaaat	240
ctcatttccca	cccacacgca	gcccgcgtac	agggttcagg	ccaacaacaa	tgacaggccc	300
gagtagacgt	gccagactgg	ccagaggccc	ctcagcgacc	ctgtgtcatc	gactgtgttt	360
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aaatcccaaga	aatttccca	tttggatcc	accttctcca	tccccacaagc	aaaccacagt	540
cacagtgggt	attaccatc	cacaggaaac	ataggctaca	cgctgttcc	atccaaaggct	600
gtgaccatca	ctgttcaagt	gcccagcatg	ggcagcttct	caccaatggg	ggtcatttgt	660
gtctgtgtca	ttggactgtc	tgtagcagcc	attgttgc	ctgtgtggc	cttgatctac	720
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tttagaaatag	ctttaacttt	gcttaaacta	caaacacaag	caaaacttca	cggggtcata	1140
ctacatacaa	gcataagcaa	aacttaactt	ggatcatttc	ttgttaatgc	ttatgttaga	1200
aataagagaa	cccccaagccaa	tcacaaaggc	cctactaaaca	tataatttag	tgacttaggaa	1260
ctttctaaga	agataccatc	cccccaaaaaaa	caattatgt	atggaaaaac	aaccgatttc	1320
cttttatttt	cttccacatt	ttcccaataaa	atacttgctt	gtgactaaaa	aaaaaaaaaa	1380
aaaaaaaaaaaa	aaaaaaaa					1396

SEQ ID NO: 30	moltype = AA length = 316					
FEATURE	Location/Qualifiers					
source	1..316					
	mol_type = protein					
	organism = Homo sapiens					
SEQUENCE: 30						
MTMETQMSQN	VCPRNLWLLQ	PLTVLLLAS	ADSQAAPPKA	VLKLEPPWIN	VLQEDSVTLT	60
CQGARSPESD	SIQWFHNGNL	IPTHTQPSYR	FKANNNNDSGE	YTCQTCQTS	SDPVHLTVLS	120
EWLVLQTPHL	EPQEGEGETIML	RCHSWKDKPL	VKVTFQNGK	SQKFSLDPT	FSIPQANHSH	180
SGDYHCTGNI	GYTLFSSKPV	TITVQVPSMG	SSSPMGVIVA	VVIATAVAII	VAAVVALIYC	240
RKKRISANST	DPVKAQFEP	PGRQMIAIRK	RQLEETNNNDY	ETADGGYMTL	NPRAPTDDDK	300
NYLTLPPND	HVNSNN					316

SEQ ID NO: 31	moltype = DNA length = 1497					
FEATURE	Location/Qualifiers					
source	1..1497					
	mol_type = other DNA					
	organism = Homo sapiens					
SEQUENCE: 31						
tgtgactgt	gtgtctgggg	cgccagctcg	ctccagggg	tgtatggaaat	cctgtcattc	60
ttacacttgc	tttccactga	gagtgtactgg	gtctgtccatc	agttccccca	gccttgggggt	120
catatgttc	tgtgcacgc	tgtgttattc	ctggcttcgt	ttgtgtggac	acctgtcgatc	180
ccccccaaagg	ctgtgtgaa	actcgagccc	cagtggatca	acgtgtccca	ggaggactct	240
gtgactctga	catggccgggg	gactcacago	cctgagagcg	actccattca	gtgggtccac	300
aatggaaatc	tccatccac	ccacacgcag	cccaagtcata	ggttcaaggc	caacaacaaat	360
gacagcgggg	agtacacgtc	ccagacggcc	ccagacggcc	tacgcgacc	tgtgtactcg	420
actctgttctt	ctgagttggct	gggtgtccag	accccttacc	tggagttcca	ggaggggagaa	480
accatctgtc	tgaggtgcca	cagctggaaag	gacaaggctc	ttgtcaaggt	cacattttcc	540
cagaatggaa	aatccaagaa	atttccgt	tcggatccca	acttctccat	cccacaagca	600
aaccacatgc	acagtggta	ttacccatc	acaggaaaaca	taggttacac	gtgtactca	660
tccaaagectg	tgaccatcac	tgttcaagct	cccaagcttct	caccgtatgg	gttcatttgt	720
gtctgtgtca	ctgggattgc	tgtgtcgcc	attgttgc	ctgtgtggc	cttgatctac	780
tgcaggaaaa	acggggatcc	agccaatccc	actaatctcg	atgaggctga	caaagttggg	840
gctgagaaca	caatcaccta	ttcacttctc	atgcaccccg	atgtctcgaa	agagcctgtat	900
gaccagaacc	gtatttagtc	tccattgtct	tgcattggaa	tttggagaaga	aaatcagaga	960
gsgaagatct	ggtatccct	ggccctaaat	ccccctgggg	aggacaggga	gtgtgtcgag	1020
ttccaaaaga	gaaggtttct	tccagagtca	tctacactgag	tcctgaagct	ccctgtctcg	1080

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aaagccacag acaatatggt cccaaataac cgactgcacc ttctgtgctt cagctctct	1140
tgcacatcaag gtccttcggc tccacatcca cacagccaat ccaattaatc aaaccactgt	1200
tattaaacaga taatagcaac ttgggaaatg cttatgttac aggttacgtg agaacaatca	1260
tgttaatcta tatgatcca gaaatgttaa aatagactaa cctctaccag cacattaaaa	1320
gtgattgttt ctgggtgata aaattattga tgatTTTtat ttctttaaaa ttcttataaa	1380
gatcatatat tactttata ataaaacatt ataaaacaa aaaaaaaaaa aaaaaaaaaa	1440
aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa aaaaaaaaaa	1497

SEQ ID NO: 32	moltype = AA length = 291
FEATURE	Location/Qualifiers
source	1..291
	mol_type = protein
	organism = Homo sapiens
SEQUENCE: 32	
MGILSFLPVL ATESDWADCK SPQPWHGMLL WTAVLFLAPV AGTPAAPPKA VLKLEPQWIN	60
VLIQEDSVTLT CRGTHSPESD SIQWFHNGNL IPTHTQPSYR FKANNNDSGE YTQQTGQTS	120
SDPVHLTVLS EWLVQLQPHL EFQEGETIVL RCHSWKDPL VKVTFFQNKG SKKFSRSDPN	180
FSIPQANHSH SGDYHCTGNI GYTLYSSKPV TITVQAPSSE PMGIIVAVVT GIAVAAIVAA	240
VVALIYCRKK RISANPTNPD EADKVGAEIT ITYSLLMHPD ALEEPDDQNR I	291

SEQ ID NO: 33	moltype = DNA length = 2137
FEATURE	Location/Qualifiers
source	1..2137
	mol_type = other DNA
	organism = Homo sapiens
SEQUENCE: 33	
cttgtccact ccagtgtggc atcatgtggc agtgcgtcct cccaaactgt ctgtacttc	60
tagtttcaggc tggcatcgcc actgaagatc tcccaaaggc tgggtgttc ctggagcctc	120
aatggtagac ggtgctcgag aaggacagtg tgactctgaa gtgccaggaa gcctactccc	180
ctgaggacaa ttccacacag tggtttcaca atgagagcct catctcaagc caggcctcga	240
gtcactctcat tgacgctggc acatggcggc acatgggaga gtacagggtc cagacaacc	300
tctccaccct cagtgaccggc tgcaactgt aagtccatcat cggctggctg ttgtccagg	360
cccctcggtt ggtgtcaag gaggaaagacc ctattcacct gaggtgtcac agctgaaaga	420
acactgtctc gcataagggtc acatatttac agaatggcaa aggccaggaa tattttcatc	480
ataatttcgtt ctcttacattt cccaaaggcc caactcaagaa cagccggctcc tacttctgca	540
ggggggctgtt tggggataaa aatgtgttcc agagactgt gaacatcccc atcaactcaag	600
gttttgcgtt gtcaaccatc tcatacttcc ttccacccgtt gtaccaactc tctttctgt	660
tggtgatgtt actcctttt gcagtgacca caggactata ttctctgtg aagacaaca	720
ttcgaaggcgc aacaatggacc tggaaaggcc ataaattttaa atggagaaag gaccctcaag	780
acaatattgacc cccatccat gggggataa agagcgttcc cagccggatc tctgaacatt	840
tctctggatt tgcaacccca tcatacttcgc gcctcttcac aagcggcagg aaacatagaa	900
ctcagagccca gatcccttat ccaactctcg acttttcctt ggtctccagt ggaaggaaaa	960
agccccatgtt ctcaaggccat ggaaggccccca gtggactgtcgtt gcatccatcg aaatttgaat	1020
ttcagagatca cacaacactc tttttgtcc caacccgttcc ctcacagccaa agcaacaata	1080
caggctagggtt atggtaatcc tttaaacata caaaaaattgc tcgtgttata aatttccatc	1140
tttagaggggg aaaaaaaaaac aatttttccat caaaaatggg ataaatggg ttaatgttg	1200
aggcggcggc atacagatgtt ggaaactgtc tggggatctt ggggatccag tgggaccaat	1260
gaaaggcatggc ctggggataaa cggatgttcc caggatgttcc taaggggatgtt gttccatct	1320
gagccccagat ataaagggtt ctctcttagaa cattttccgtt agtggaaata acaggaaatc	1380
atgggggtgtt cgtatgttcc agtgcgttccatc gggactctat cagaactggc ccatttcacaa	1440
gtatataacgc atagatgttcc ttaatgttccatc gggactctat cagaactggc ccatttcacaa	1500
aggattggccggc tggggggatgg ggtggaaatgg aatgttccatc aacaaacccctg tggactgtc	1560
ccaaatgttccatc aactatctca gcatcgttccatc gggactctat cagaactggc ccatttcacaa	1620
aaccaaccac aaggccacacag gaaggaaagc gcaggagggtt aaaaatgttccatc tttggccagg	1680
gtatgttccatc aactatctca gcatcgttccatc gggactctat cagaactggc ccatttcacaa	1740
aatttccatc tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt	1800
atgttactatc gatgttccatc gggactctat cagaactggc ccatttcacaa	1860
tttccatc tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt	1920
tctgtttttt tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt	1980
tttccatc tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt tttttttttt	2040
aaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa	2100
aaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa aaaaaaaaaaaaaaaa	2137

SEQ ID NO: 34	moltype = AA length = 254
FEATURE	Location/Qualifiers
source	1..254
	mol_type = protein
	organism = Homo sapiens
SEQUENCE: 34	
MWQLLLPTAL LLLVSAGMRT EDLPKAVVFL EPQWYRVLEK DSVTLKCQGA YSPEDNSTQW	60
FHNESLIISSQ ASSYFIDAAT VDDSGEYRCQ TNLSTLSDPV QLEVHIGWLL LQAPRWVFKE	120
EDPIHLRCHS WKNTALHKVT YLQNGKGRKY FHHNSDFYIP KATLKDSGSY FCRGLVGSKN	180
VSSSETVNITI TQGLSVSTIS SFFPPGYQVS FCLVMVLLFA VDTGLYFSVK TNIRSSTRDW	240
KDHKFKWRKD PQDK	254

SEQ ID NO: 35	moltype = DNA length = 820
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FEATURE	Location/Qualifiers
source	1..820
	mol_type = other DNA
	organism = Homo sapiens
SEQUENCE: 35	
cactccagt	tggcatcatg tggcagctgc tcccccacat tgctctgcata cttcttagtt 60
cagctggcat	cgggactgaa gatctccca aggtctggat gtctctggag cctcaatgggt 120
acacgcgtct	tgagaaggac agtgcgtact tgaatgtccca gggagctac tccccctgagg 180
acaattccac	acatgtggttt cacaatgaga gcctcatctc aaggcaggcc tcgagctact 240
tcattgcgc	tgcccacatc aacgcacatg gagagatcagat gtgcacagaca aacctctcca 300
ccttcgtga	ccttcgtca atatcggtcg cttagaactg atatcggtcg cttagaactg 360
ggtgtgggtt	caaggaggaa gacccattt acctcggttg tcacagctgg aagaacactg 420
ctctgcataa	ggtcacatat ttacagaatg gcaaagacag gaagtattt catataatt 480
ctgacttcca	cattccaaa gccacacta aagatagcggg ctcctactc tgcagggggc 540
ttgttgggg	tttttttttggat ttctcgatca ctgtgaacat caccatact caagggttgg 600
caatgtcac	catctcatca ttcttcacat ctgtgttccat agtcttttc tgctttgtga 660
ttgttactct	tttttgcgtg gacacaggac tatatttctc ttttgcgtg gacacaggac 720
gctcaacaag	agactggaa gaccataaa ttaaatggag aaaggaccct caagacaaat 780
gaccccccatac	ccatgggatg aataagagca gtggcagcag 820
SEQ ID NO: 36	moltype = AA length = 233
FEATURE	Location/Qualifiers
source	1..233
	mol_type = protein
	organism = Homo sapiens
SEQUENCE: 36	
MWQLLLPTAL	LLLVSAGMRT EDLPAVVFLEK EPQWYSVLEK DSVTLKCQGA YSPEDNSTQW 60
FHNESLISQQ	ASSYFIDAAT VNDSEYRCQ TNLSLSDPV QLEVHIGWLL LQAPRWFVKE 120
EDPIHLRCHS	WKNTALHKVT YLQNKGDRKY FHHNSDFHIP KATLKDSGSY FCRLVGSKN 180
VSSETVNITI	TQGLAVSTIS SFSPPGYQVS FCLVMVLFLA VDTGLYFSVK TNI 233
SEQ ID NO: 37	moltype = AA length = 330
FEATURE	Location/Qualifiers
source	1..330
	mol_type = protein
	organism = Homo sapiens
SEQUENCE: 37	
ASTKGPSVFP	LAPSSKSTSG GTAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT	VPSSNFTQT YTCNVDHKPS NTKVDKTKVER KCCVECPFC APPVAGPSVF 120
PSVFLFPPKP	KDTLMISRTP EVTCVVVDVS HEDPEVKFWN YVDGVEVHNA KTKPREEQYN 180
STYRVVSVLT	VLHQDWLNGK EYKCKVSNKA LPAPIEKTS KAKGQPREPQ VYTLPPSRDE 240
LTKNQVSLTC	LVKGFYPSDI AVEWESNGQP ENNYKTTPPV LDSDGSFFLY SKLTVDKSRW 300
QQGNVFSCSV	MHEALHNHYT QKSLSLSPGK 330
SEQ ID NO: 38	moltype = AA length = 326
FEATURE	Location/Qualifiers
source	1..326
	mol_type = protein
	organism = Homo sapiens
SEQUENCE: 38	
ASTKGPSVFP	LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT	VPSSNFTQT YTCNVDHKPS NTKVDKTKVER KCCVECPFC APPVAGPSVF 120
LFPKPKDML	MISRTPEVTC VVVDVSHEDP EVQFNWYVDG VEVHNAKTKP REEQFNSTFR 180
VVSVLTVVHQ	DWLNGKEYKC KVSNKGLPAP IEKTISKTKG QPREPQVYTL PPSREEMTKN 240
QVSLTCLVKG	FYPYPSDIAVEW ESNQOPENNY KTTTPMLSD GSFFFLYSKLT VDKSRWQGN 300
VFSCSVMHEA	LHNHYTQKSL SLSPGK 326
SEQ ID NO: 39	moltype = AA length = 327
FEATURE	Location/Qualifiers
source	1..327
	mol_type = protein
	organism = Homo sapiens
SEQUENCE: 39	
ASTKGPSVFP	LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT	VPSSSLGTKT YTCNVDHKPS NTKVDKTRVES KYGPPCPSCP APEFLGGPSV 120
FIFPPKPKDT	LMISRTPEVTC CVVVDVQSED PEVQFNWYVDG VEVHNAKTKP REEQFNSTY 180
RVSVLTVLH	QDWLNGKEYK CKVSNKGLPS SIEKTISKAK QGPREPQVYT LPPSQEEMTK 240
NQVSLTCLVK	GFYPSDIAVEW WESNGQOPENNY YKTTPPVLDS DGSFFFLYSRLL TVDKSRWQEG 300
NVFSCSVMHE	ALHNHYTQKS LSLSLGK 327
SEQ ID NO: 40	moltype = AA length = 115
FEATURE	Location/Qualifiers
REGION	1..115
note	An artificially generated sequence
source	1..115
	mol_type = protein

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organism = synthetic construct
 SEQUENCE: 40
 QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDPKTGDTAY 60
 SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQGTL VTVSS 115

 SEQ ID NO: 41 moltype = AA length = 112
 FEATURE Location/Qualifiers
 REGION 1..112
 note = An artificially generated sequence
 source 1..112
 mol_type = protein
 organism = synthetic construct

 SEQUENCE: 41
 DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTLYLHW YQQKPGQAPR LLIYKVSNRF 60
 SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCSQNTHVP PTFGQGTKLE IK 112

 SEQ ID NO: 42 moltype = AA length = 122
 FEATURE Location/Qualifiers
 source 1..122
 mol_type = protein
 organism = Rattus norvegicus

 SEQUENCE: 42
 EVQLVETGGS LVQPGKSLKL TCATSGFTFS NAWMHWVRQS PEKQLEWVAQ IKAKSNNYAT 60
 YYAESVKGRF TISRDDSKSN IYLQMNSLKE EDTAVYYCRY VHGYAYGVN AWGQGTSVTV 120
 SS 122

 SEQ ID NO: 43 moltype = AA length = 112
 FEATURE Location/Qualifiers
 source 1..112
 mol_type = protein
 organism = Rattus norvegicus

 SEQUENCE: 43
 DVVMTQTPVS MSVSLGGQVS ISCRSSQSLV HNNNGNTYVSW YIQQKPSQSPQ LLIYKVSNRF 60
 SGISDRFSGS GSGTDFTLKI SRVEPDDLGV YYCGQGTQDP YTFGAGTKLE LK 112

 SEQ ID NO: 44 moltype = AA length = 328
 FEATURE Location/Qualifiers
 REGION 1..328
 note = An artificially generated sequence
 source 1..328
 mol_type = protein
 organism = synthetic construct

 SEQUENCE: 44
 ASTKGPSVFP LAPSSKSTSG GTAALGCLVK DYPFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
 GLYSLSSSVT VPSSSLGTQT YICNVNHKPS NTKVDKKVEP KSCDKTHTCP PCPAPELLGG 120
 PSVFLFPKP KDTLMISRTP EVTCVVVDVS HEDPEVKFNW YVDGVEVHNNA KTKPREEQYN 180
 STYRVSVLVT VLHQDWLNKG EYKCKVSNKA LPAPIEKTK KAKGQPREPQ VYTLPSPCRE 240
 LTKNQVSLSC AVKGFYPSDI AVEWESNGQP ENNYKTTPPV LDSDGSFFLV SKLTVDKSRW 300
 QQGNVFSCSV MHEALHNHYT QKSLSLSP 328

 SEQ ID NO: 45 moltype = AA length = 338
 FEATURE Location/Qualifiers
 REGION 1..338
 note = An artificially generated sequence
 source 1..338
 mol_type = protein
 organism = synthetic construct

 SEQUENCE: 45
 SSASTKGPSV FPLAPSSKST SGGTAALGCL VKDYFPEPVTVS VSWNSGALTGS GVHTFPVLQ 60
 SSGLYSLSSV TVTVPSSSLGT QTYICNVNHK PSNTKVDKVK EPKSCDKTHTCP CPPCPAPEAA 120
 GGPSVFLFP KPDKDTLMISR TPEVTCVVVD VSHEDPEVKFNW NWFYDGVENHNNA KTKPREEQ 180
 YASTYRVSVT LTVLHQDWLNKG EYKCKVSNKA LPAPIEKTK ISKAKGQPREPQ PQVCTLPPSR 240
 DELTKNQVSLC WCLVKGFYPS DIAVEWESNGQ QPENNYKTTPPV PVLDSDGSFFLYS KLTVDKSRW 300
 RWQGNVFSCSV SVMHEALHNHYT QTQKSLSLSP HHHHHHHHHH 338

 SEQ ID NO: 46 moltype = AA length = 336
 FEATURE Location/Qualifiers
 REGION 1..336
 note = An artificially generated sequence
 source 1..336
 mol_type = protein
 organism = synthetic construct

 SEQUENCE: 46
 ASTKGPSVFP LAPSSKSTSG GTAALGCLVK DYPFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
 GLYSLSSSVT VPSSSLGTQT YICNVNHKPS NTKVDKKVEP KSCDKTHTCP PCPAPELLGG 120
 PSVFLFPKP KDTLMISRTP EVTCVVVDVS HEDPEVKFNW YVDGVEVHNNA KTKPREEQY 180
 A 120

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STYRVSLSLT VLHQDWLNGK EYKCKVSNKA LPAPIEKTI KAKGQPREGQ VYTLPPSRCE 240
LTKNQVSLSC AVKGFYPSDI AVEWESNGQP ENNYKTTPV LDSDGSFFLV SKLTVDKSRW 300
QQGNVFSCSV MHEALHNHYT QKSLSLSPDY KDDDK 336
SEQ ID NO: 47 moltype = AA length = 450
FEATURE Location/Qualifiers
REGION 1..450
note = An artificially generated sequence
source 1..450
mol_type = protein
organism = synthetic construct
 SEQUENCE: 47
DIVMTQSPLS LPVTPGE PAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFGS GSGTDFTLKI SRVEAEDVG V YCSQNTFVG PTFGQQGTKE IKSSASTKGP 120
SVFPLAPSSK STSGGTAALG CLVKDPSVKT VTVSWNSGAL TSGVHTFP AV LQSSGLYLSL 180
SVVTPVPSL GTQTYICCNVN HKPSNTKVDK KVEPKSCDKT HTCPCPAPE AAGGPSPVFLF 240
PPPKPDLMI SRTPEVTCVV VDVSHEDPEV KFNWYVDGVE VHNAKTKPRE EQYASTYRVV 300
SVLTVLHQDW LNGKEYKCKV SNKALPAPIE KTISKAKQGP REPQVCTLPP SRDELTKNQV 360
SLWCLVKGFY PSDIAVEWES NGQPENNYKT TPPVLDSDGS FFLYSKLTVD KSRWQQGNVF 420
SCSVMHEAL NHYTQKSLSL SPHHHHHHHHH 450
SEQ ID NO: 48 moltype = AA length = 222
FEATURE Location/Qualifiers
REGION 1..222
note = An artificially generated sequence
source 1..222
mol_type = protein
organism = synthetic construct
 SEQUENCE: 48
QVQLVQSGAE VKKPGASV KV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDPKTGDTAY 60
SQFKGRVTL TADKSTSTAY MELSSLSED TAVYYCTR FY SYTYWGQGTL VTVSSASVAA 120
PSVFIFFPSD EQLKSGTASV VCLLNNFYPR EAKVQWKVDN ALQSGNSQES VTEQDSKDST 180
YSLSTTLTS KADYEKHKVY ACEVTHQGLS SPVTKSFNRG EC 222
SEQ ID NO: 49 moltype = AA length = 458
FEATURE Location/Qualifiers
REGION 1..458
note = An artificially generated sequence
source 1..458
mol_type = protein
organism = synthetic construct
 SEQUENCE: 49
EQLVETGGS DVQPGKSLKL TCATSGFTFS NAWMHWRQD PEKQLEWVAQ IKA KSNNYAT 60
YYAESVKG RF TISRDDSKSN IYLQMNSLKE EDTAVYYCRY VHYGAYYGV D AWGQGTSVT 120
SSASTKGPSV FPLAPSSKST SGGAALGCL V KVDYFPEPV T VSWNSGAL TS GVHTPAV LQ 180
SSGLYSLSSV VTVPSSSLGT QTYICCNVNHK PSNTKVDK V EPKSCDKTHT CPPCPAPEA 240
GGPSVFLPPP KPKDITMIS TPEVTCVVVD VSHE DPEVKF NWYV DGV EHV NAKT KPRE EQ 300
YASTYRVVSV LTVLHQDWLN GKEYCKVSN KALPAPIE KT ISKAKQGP RE PQVY TL PPSR 360
CETLKNQVSL SCAVKGFYPS DIAVEWESNG QPEN NYKTT P VLVLDSDGSFF LVSKLTVDK 420
RWQGNVFSC SVMHEALHNH YTQKSLSLSP DYKDDDDK 458
SEQ ID NO: 50 moltype = AA length = 219
FEATURE Location/Qualifiers
REGION 1..219
note = An artificially generated sequence
source 1..219
mol_type = protein
organism = synthetic construct
 SEQUENCE: 50
DVVMQTPTVS MSVSLGGQVS ISCRSSQSLV HNNGNTYVSW YIQKPSQSPQ LLIYKVSNRF 60
SGISDRFGS GSGTDFTLKI SRVEPDDLG V YYCGQGTQDP YTFGAGTKLE LKRTVAAPSV 120
FIFPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180
STSTLTL SKAD YEKHKVYACE VTHQGLSSPV TKS FNRGEC 219
SEQ ID NO: 51 moltype = AA length = 122
FEATURE Location/Qualifiers
REGION 1..122
note = An artificially generated sequence
source 1..122
mol_type = protein
organism = synthetic construct
 SEQUENCE: 51
QVOLVESGGG VVQPGRSRLR SCAASGFTFS NAWMHWRQD PGKGLEWVAQ IKA KSNNYAT 60
YYAESVKG RF TISRDDSKNS LYLMQNSLKT EDTAVYYCAR VHYGAYYGV D AWGQGTTVT 120
SS

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SEQ ID NO: 52
FEATURE
REGION
source
SEQUENCE: 52
QVOLVESGG VVQPGRSRLR SCAASGFTFS NAWMHWRQA PGKGLEWVAQ IKAKSNNYAT YYAESVKGRF TISRDDSKNS LYLMQMSNLKT EDTAVYYCRY VHGYAYYGVD AWGQGTTVT
SS

SEQ ID NO: 53
FEATURE
REGION
source
SEQUENCE: 53
DIVMTQSPLS LPVTPGEPAS ISCRSSQLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK

SEQ ID NO: 54
FEATURE
REGION
source
SEQUENCE: 54
QVOLVQSGAE VKKPGASVVK SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDPKTGDTAY SQKFKGVRFL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQGTL VTUVSASTKG PSVPFLAPCS RSTSESTAAL GCLVKDFYFPV PVTWSNLSGA LTSGVHHTFPV VLQSSGLYSL SSVTVPSSS LGTKTYTCNV DHKPSNTKVD KRVESKYGP CPPCPAPEFR GGPKVFLFP KPKDTLMSR TPEVTCVVVD VSQEDPEVQF NWYVGDGEVHN NAKTKPREEQ FASTYRVVSV LTVLHQDWLN GKEYKCKVSN KGLPSSIEKT ISAKAGQPRE PQVYTLPPSQ KEMTKNQVSL TCLVKGFYPS DIAVEWESNG QPENNYKTTPT PVLDSDGSFF LYSRSLTVDKS RWQEGNVFSC SVMHEALTHNR YTQKSLSSLH HHHHHHHHH

SEQ ID NO: 55
FEATURE
REGION
source
SEQUENCE: 55
QVOLVESGG VVQPGRSRLR SCAASGFTFS NAWMHWRQA PGKGLEWVAQ IKAKSNNYAT YYAESVKGRF TISRDDSKNS LYLMQMSNLKT EDTAVYYCRY VHGYAYYGVD AWGQGTTVT SSASTKGPSV FPLAPCSRST SESTAALGCL VKDYFPEPV TWSNMSALTS GVHTPPAVLQ SSGLYSLSSV TVPSSSLGT KTYTCNVDHK PSNTKVDKRV ESKYGGPPCPP CPAPEFRGGP KVFLFPPKPK DTLMISRTPE VTCVVVDVSQ EDPEVQFNWY VDGVEVHNK TKPREEQFAS TYRVSVLTV LHQDWLNKE YKCKVSNKGL PSSIEKTIISK AKGQPREPQV YTLPSSQEEM TKNQVSLTCL VKGFYPSDIA VEWESESNGPQ NNYKTTPPVL DSDGSFFLYS RLTVDKSRWQ EGNVFSCSVM HEALHNHYTQ ESLSLSLDYK DDDDD

SEQ ID NO: 56
FEATURE
REGION
source
SEQUENCE: 56
DIVMTQSPLS LPVTPGEPAS ISCRSSQLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IKRTVAAPSV FIFPPSDEQL KSGTASVVC LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL SSTLTLASKAD YEKHVKVYACE VTHQGLSSPV TKSFNRGEC

SEQ ID NO: 57
FEATURE
REGION
source
SEQUENCE: 57
moltype = AA length = 122
Location/Qualifiers
1..122
note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

moltype = AA length = 112
Location/Qualifiers
1..112
note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

moltype = AA length = 448
Location/Qualifiers
1..448
note = An artificially generated sequence
1..448
mol_type = protein
organism = synthetic construct

moltype = AA length = 455
Location/Qualifiers
1..455
note = An artificially generated sequence
1..455
mol_type = protein
organism = synthetic construct

moltype = AA length = 219
Location/Qualifiers
1..219
note = An artificially generated sequence
1..219
mol_type = protein
organism = synthetic construct

moltype = AA length = 336
Location/Qualifiers
1..336
note = An artificially generated sequence
1..336
mol_type = protein

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SEQUENCE: 57          organism = synthetic construct
ASTKGPSVFP LAPSSKSTSG GTAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT VPSSSLGTQT YICCNVNHKPS NTKVDKKVEP KSCDKTHTCP PCPAPEAAGG 120
PSVFLFPKP KDTLMISRTP EVTCVVVDVS HEDPEVKFNW YVDGVEHNA KTKPREEQYA 180
STYRVVSVLT VLHQDWLNGK EYKCKVSNKA LPAPIEKTAKAGQPREPQ VYTLPPSRCE 240
LTKNQVSLSC AVKGFYPSDI AVEWESNGQP ENNYKTTPPV LDSDGSFFLV SKLTVDKSRW 300
QQGNVFSCSV MHEALHNHYT QKSLSLSPDY KDDDDK 336

SEQ ID NO: 58          moltype = AA length = 338
FEATURE
REGION
1..338
note = An artificially generated sequence
source
1..338
mol_type = protein
organism = synthetic construct
SEQUENCE: 58
SSASTKGPSV FPLAPSSKST SGGTAALGCL VKDYFPEPVTV SWSNGALTS GVHTFPAVLQ 60
SSGLYSLPSS VTPSSSLGT QTYYICCNVNHK PSNTKVDKKV EPKSCDKHTC CPPCPAPEAA 120
GGPSVFLPSS KPKDTLMISR TPEVTCVVVD VSHEDEPEVKF NWYVDGVEHV NAKTKPREEQ 180
YASTYRVVSV LTVLHQDWLN GKEYKCKVSN KALPAPIEKT ISKAKGQPRE PQVCTLPPSR 240
DELTKNQVSL WCLVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LYSKLTVDKS 300
RWQQGNVFSC SVMHEALHNH YTQKSLSLSP HHHHHHHHHH 338

SEQ ID NO: 59          moltype = AA length = 230
FEATURE
REGION
1..230
note = An artificially generated sequence
source
1..230
mol_type = protein
organism = synthetic construct
SEQUENCE: 59
EPKSSDKTHT CPPCPAPELL GGGSVFLFPP KPKDTLMISR TPEVTCVVVD VSHEDEPEVKF 60
NWYVDGVEHV NAKTKPREEQ YNSTYRVVSV LTVLHQDWLN GKEYKCKVSN KALPAPIEKT 120
ISKAKGQPRE PQVCTLPPSR DELTKNQVSL WCLVKGFYPS DIAVEWESNG QPENNYKTTP 180
PVLDSDGSFF LYSKLTVDKS RWQQGNVFSC SVMHEALHNH YTQKSLSLSP 230

SEQ ID NO: 60          moltype = AA length = 325
FEATURE
REGION
1..325
note = An artificially generated sequence
source
1..325
mol_type = protein
organism = synthetic construct
SEQUENCE: 60
ASTKGPSVFP LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT VPSSSLGTQT YTCNVNDHKPS NTKVDKRVES KYGPPCPCCP APEFRGGPKV 120
FLFPPPKDT LMISRTPEVT CVVVDVSQED PEVQFNWYVD GVEVHNAKTK PREEQFASTY 180
RVSVSLTVLH QDWLNGKEYK CKVSNKGLPS SIEKTISKAK GQPREQQVYT LPPSQEMTK 240
NQVSLTCLVK GFYPSDI AVEWESNG PENN YKTTPPVLDs DGSSFLYSLK TVDKSRWQEG 300
NVPSCSVMHE ALHNRYTQKS LSLSP 325

SEQ ID NO: 61          moltype = AA length = 325
FEATURE
REGION
1..325
note = An artificially generated sequence
source
1..325
mol_type = protein
organism = synthetic construct
SEQUENCE: 61
ASTKGPSVFP LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60
GLYSLSSVVT VPSSSLGTQT YTCNVNDHKPS NTKVDKRVES KYGPPCPCCP APEFRGGPKV 120
FLFPPPKDT LMISRTPEVT CVVVDVSQED PEVQFNWYVD GVEVHNAKTK PREEQFASTY 180
RVSVSLTVLH QDWLNGKEYK CKVSNKGLPS SIEKTISKAK GQPREQQVYT LPPSQEMTK 240
NQVSLTCLVK GFYPSDI AVEWESNG PENN YKTTPPVLDs DGSSFLYSLK TVDKSRWQEG 300
NVPSCSVMHE ALHNRYTQKS LSLSP 325

SEQ ID NO: 62          moltype = AA length = 325
FEATURE
REGION
1..325
note = An artificially generated sequence
source
1..325
mol_type = protein
organism = synthetic construct
SEQUENCE: 62
ASTKGPSVFP LAPCSRSTSE STAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPAVLQSS 60

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GLYSLSSVVT VPSSSLGTKT YTCNVDHKPS NTKVDKRVES KYGPPCPCP APEFRGGPKV	120
FLFPPPKDFT LMISRTPEVT CVVUDVSQED PEVQFNWYVD GVEVHNAKTK PREEQFASTY	180
RVSVLTVLH QDWLNKEYK CKVSNKGLPS SIEKTISKAK GQPREPQVYT LPPSQKEMTK	240
NQVSLTCLVK GFYPSDIAVE WESNGQPENN YKTTPPVLDS DGSFFLYSKL TVDKSRWQEG	300
NVFSCSVMHE ALHNHYTQKS LSLSP	325
 SEQ ID NO: 63	
FEATURE	moltype = AA length = 107
REGION	Location/Qualifiers
1..107	
source	note = An artificially generated sequence
1..107	
mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 63	
RTVAAPSVFIFTPSDEQLKS GTASVVCLLN NFYPREAKVQ WKVDNALQSG NSQESVTEQD	60
SKDSTYSLSS TLTLSKADYE KHKVYACEVT HQGLSSPVTK SFNRGEC	107
 SEQ ID NO: 64	
FEATURE	moltype = AA length = 122
REGION	Location/Qualifiers
1..122	
source	note = An artificially generated sequence
1..122	
mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 64	
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT	60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGVQ AWGQGTTVT	120
SS	122
 SEQ ID NO: 65	
FEATURE	moltype = AA length = 122
REGION	Location/Qualifiers
1..122	
source	note = An artificially generated sequence
1..122	
mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 65	
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT	60
YYAESVKGRF TISRDDSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGVQ AWGQGTTVT	120
SS	122
 SEQ ID NO: 66	
FEATURE	moltype = AA length = 122
REGION	Location/Qualifiers
1..122	
source	note = An artificially generated sequence
1..122	
mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 66	
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT	60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGVQ AWGQGTTVT	120
SS	122
 SEQ ID NO: 67	
FEATURE	moltype = AA length = 122
REGION	Location/Qualifiers
1..122	
source	note = An artificially generated sequence
1..122	
mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 67	
QVQLVESGGG VVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT	60
YYAESVKGRF TISRADSKNS LYLMQNSLKT EDTAVYYCRY VHYGAYYGVQ AWGQGTTVT	120
SS	122
 SEQ ID NO: 68	
FEATURE	moltype = AA length = 122
REGION	Location/Qualifiers
1..122	
source	note = An artificially generated sequence
1..122	
mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 68	
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT	60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYGAYYGVQ AWGQGTTVT	120
SS	122

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SEQ ID NO: 69      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 69
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 70      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 70
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 71      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 71
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 72      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 72
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAPSVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSAYYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 73      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 73
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSAYYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 74      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 74
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGASYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 75      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122

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source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 75
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYYYSYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 76      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 76
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSQNYAT 60
YYAPSVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGV 120
SS                           122

SEQ ID NO: 77      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 77
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAPSVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGV 120
SS                           122

SEQ ID NO: 78      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 78
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGV 120
SS                           122

SEQ ID NO: 79      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 79
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGV 120
SS                           122

SEQ ID NO: 80      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 80
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYGAYYGV 120
SS                           122

SEQ ID NO: 81      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

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SEQUENCE: 81
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 82      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 82
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 83      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 83
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAPSVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 84      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 84
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSQNYAT 60
YYAPSVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 85      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 85
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSQNYAT 60
YYAPSVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 86      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 86
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 87      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 87
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS 122

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SEQ ID NO: 88      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = An artificially generated sequence
source            1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 88
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKSKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 89      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = An artificially generated sequence
source            1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 89
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKSKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 90      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = An artificially generated sequence
source            1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 90
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKSKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 91      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = An artificially generated sequence
source            1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 91
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKSKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 92      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = An artificially generated sequence
source            1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 92
QVQLVESGGG LVQPGRSLRL SCSASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 93      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = An artificially generated sequence
source            1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 93
QVQLVESGGG VVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 94      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122

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source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 94
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 95      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 95
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKSKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 96      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 96
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKSKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 97      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 97
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 98      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 98
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 99      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 99
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 100     moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

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SEQUENCE: 100
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMPWVRQA PGKGLEWVAQ IKGKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 101      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 101
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 102      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 102
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 103      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 103
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 104      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 104
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS 122

SEQ ID NO: 105      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 105
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 106      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 106
QVQLVESGGG LVQPGRSLRL SCSASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

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SEQ ID NO: 107      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 107
QVQLVESGGG LVQPGRSLRL SCSASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 108      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 108
QVQLVESGGG LVQPGRSLRL SCSASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 109      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 109
QVQLVESGGG LVQPGRSLRL SCSASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 110      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 110
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSNNYAT 60
YYAESVKGRF TISRDDSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 111      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 111
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 112      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 112
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 113      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122

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source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 113
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 114      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 114
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 115      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 115
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 116      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 116
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 117      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 117
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 118      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 118
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 119      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

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SEQUENCE: 119
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 120      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 120
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 121      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 121
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYSASYGVD YWGQGTTVT 120
SS 122

SEQ ID NO: 122      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 122
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLMQNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT 120
SS 122

SEQ ID NO: 123      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 123
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLMQNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 124      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 124
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLMQNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 125      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 125
QVQLVESGGG LVQPGRSLRL SCSASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLMQNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS 122

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SEQ ID NO: 126      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 126
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS IYLQMNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 127      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 127
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 128      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 128
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 129      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 129
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD AWGQGTTVT 120
SS                           122

SEQ ID NO: 130      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 130
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 131      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 131
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 132      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122

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source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 132
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 133      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 133
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 134      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 134
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 135      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 135
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT 120
SS                           122

SEQ ID NO: 136      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 136
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 137      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 137
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAASYGVD YWGQGTTVT 120
SS                           122

SEQ ID NO: 138      moltype = AA  length = 124
FEATURE           Location/Qualifiers
REGION            1..124
source          note = An artificially generated sequence
                1..124
                mol_type = protein
                organism = synthetic construct

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SEQUENCE: 138
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAAEMVR DDIWGQGTTV 120
TVSS                                         124

SEQ ID NO: 139      moltype = AA  length = 128
FEATURE           Location/Qualifiers
REGION            1..128
source             note = An artificially generated sequence
                  1..128
                  mol_type = protein
                  organism = synthetic construct

SEQUENCE: 139
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAAEMVR DSYGVDIWGQ 120
GTTVTVSS                                         128

SEQ ID NO: 140      moltype = AA  length = 128
FEATURE           Location/Qualifiers
REGION            1..128
source             note = An artificially generated sequence
                  1..128
                  mol_type = protein
                  organism = synthetic construct

SEQUENCE: 140
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSNNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAVRRRL SSYGVDIWGQ 120
GTTVTVSS                                         128

SEQ ID NO: 141      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct

SEQUENCE: 141
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D I WGQGTTVTV 120
SS                                         122

SEQ ID NO: 142      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct

SEQUENCE: 142
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVTV 120
SS                                         122

SEQ ID NO: 143      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct

SEQUENCE: 143
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D I WGQGTTVTV 120
SS                                         122

SEQ ID NO: 144      moltype = AA  length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct

SEQUENCE: 144
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D I WGQGTTVTV 120
SS                                         122

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SEQ ID NO: 145      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 145
QVQLVESGGG LVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 146      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 146
QVQLVESGGG LVQPGRSLKL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 147      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 147
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVAGR F TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 148      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 148
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVQGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 149      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 149
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVSGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 150      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 150
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAF 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 151      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122

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source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 151
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAG 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 152      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 152
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAL 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 153      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 153
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAP 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 154      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 154
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAQ 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 155      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 155
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAS 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 156      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 156
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAV 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 157      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source          note = An artificially generated sequence
                1..122
                mol_type = protein
                organism = synthetic construct

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SEQUENCE: 157
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YFAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 158      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 158
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YGAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 159      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 159
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YIAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 160      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 160
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YIAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 161      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 161
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YPAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 162      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 162
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YPAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 163      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
source             note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 163
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YTAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT 120
SS 122

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SEQ ID NO: 164      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 164
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 165      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 165
QVQLVESGGG VVQPGRSLKL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYAAGYGV D AWGQGTTVT 120
SS                           122

SEQ ID NO: 166      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 166
QVQLVESGGG VVQPGGSLKL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 167      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 167
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAQ 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 168      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 168
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 169      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
                  1..122
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 169
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAS 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGV D IWGQGTTVT 120
SS                           122

SEQ ID NO: 170      moltype = AA  length = 115
FEATURE          Location/Qualifiers
REGION           1..115

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source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 170
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGDPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 171      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 171
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAE 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 172      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 172
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 173      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 173
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SDKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 174      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 174
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGETPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 175      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 175
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGDPDTAY 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 176      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 176
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAE 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 177      moltype = AA length = 115
FEATURE          Location/Qualifiers

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REGION          1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 177
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 178      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 178
QVQLVQSGAE VKKPGASVKG SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGKTPDTAY 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 179      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 179
QVQLVQSGAE VKKPGASVKG SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
EESFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 180      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 180
QVQLVQSGAE VKKPGASVKG SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
EQKFKQDRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 181      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 181
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGKTPDTAY 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 182      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 182
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGETPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 183      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source          note = An artificially generated sequence
                1..115
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 183
QVQLVQSGAE VKKPGASVKG SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 184      moltype = AA length = 115

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FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 184
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 185      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 185
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SESFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 186      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 186
QVQLVQSGAE VKKPGASVTV SCKASGYEFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 187      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 187
QVQLVQSGAE VKKPGASVTV SCKASGYTFE DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 188      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 188
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SEKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 189      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 189
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFEGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

SEQ ID NO: 190      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
source            note = An artificially generated sequence
                  1..115
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 190
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVEL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS      115

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SEQ ID NO: 191      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 191
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELESLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS     115

SEQ ID NO: 192      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 192
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSELTSED TAVYYCTRFY SYTYWGQQGTL VTVSS     115

SEQ ID NO: 193      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 193
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTEED TAVYYCTRFY SYTYWGQQGTL VTVSS     115

SEQ ID NO: 194      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 194
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED EAVYYCTRFY SYTYWGQQGTL VTVSS     115

SEQ ID NO: 195      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 195
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAEYYCTRFY SYTYWGQQGTL VTVSS     115

SEQ ID NO: 196      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 196
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
EQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS     115

SEQ ID NO: 197      moltype = AA length = 115
FEATURE          Location/Qualifiers
REGION           1..115
note = An artificially generated sequence
source            1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 197
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SEKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS     115

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SEQ ID NO: 198      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 198
QVQLVQSGAE VKKPGASVTW SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS 115

SEQ ID NO: 199      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 199
QVQLVQSGAE VKKPGASVTW SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SQKFEGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS 115

SEQ ID NO: 200      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 200
QVQLVQSGAE VKKPGASVTW SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SEKFEGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS 115

SEQ ID NO: 201      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 201
QVQLVQSGAE VKKPGASVTW SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SQKFEGRVTL TADKSTSTAY MELSSLTED TAVYYCTRFY SYTYWGQQGTL VTVSS 115

SEQ ID NO: 202      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 202
QVQLVQSGAE VKKPGASVTW SCKASGYTFE DYEMHWIRQP PGEGLEWIGA IDGETPDTAY 60
SQKFEGRVTL TADKSTSTAY MELSSLTED TAVYYCTRFY SYTYWGQQGTL VTVSS 115

SEQ ID NO: 203      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 203
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWVRQPA PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQQGTL VTVSS 115

SEQ ID NO: 204      moltype = AA  length = 115
FEATURE
REGION
1..115
note = An artificially generated sequence
source
1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 204
QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWMGA IDGKTPDTAY 60

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SQFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS      115
SEQ ID NO: 205      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 205
QVQLVQSGAE VKKPGASV р SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGKTPDTAY 60
SQFKGRVTL TADKSTSTAY MELSSLRSED TAVYYCTRFY SYTYWGGT VTVSS      115

SEQ ID NO: 206      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 206
QVQLVQSGAE VKKPGASV р SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGPTPDTAY 60
SEKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS      115

SEQ ID NO: 207      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 207
QVQLVQSGAE VKKPGASV р SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGWTPTAY 60
SEKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS      115

SEQ ID NO: 208      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 208
QVQLVQSGAE VKKPGASV р SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGYTPDTAY 60
SEKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS      115

SEQ ID NO: 209      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 209
QVQLVQSGAE VKKPGASV р SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGKTPDTAY 60
SEKFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS      115

SEQ ID NO: 210      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 210
QVQLVQSGAE VKKPGASV р SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGPTPDTAY 60
SQKFGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS      115

SEQ ID NO: 211      moltype = AA length = 115
FEATURE
REGION          Location/Qualifiers
1..115
note = An artificially generated sequence
source           1..115
mol_type = protein
organism = synthetic construct
SEQUENCE: 211

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QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWVRQA PGEGLEWMGA IDGETPDTAY 60
SEFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS 115

SEQ ID NO: 212 moltype = AA length = 115
FEATURE Location/Qualifiers
REGION 1..115
note = An artificially generated sequence
source 1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 212 QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWVRQA PGEGLEWMGA IDGPTPDTAY 60
SEFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS 115

SEQ ID NO: 213 moltype = AA length = 115
FEATURE Location/Qualifiers
REGION 1..115
note = An artificially generated sequence
source 1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 213 QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWVRQA PGEGLEWMGA IDGWTPTAY 60
SEFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS 115

SEQ ID NO: 214 moltype = AA length = 115
FEATURE Location/Qualifiers
REGION 1..115
note = An artificially generated sequence
source 1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 214 QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWVRQA PGEGLEWMGA IDGYTPDTAY 60
SEFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS 115

SEQ ID NO: 215 moltype = AA length = 115
FEATURE Location/Qualifiers
REGION 1..115
note = An artificially generated sequence
source 1..115
mol_type = protein
organism = synthetic construct

SEQUENCE: 215 QVQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWIRQP PGQGLEWI GA IDGKTPDTAY 60
SQFKGRVTL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGGT VTVSS 115

SEQ ID NO: 216 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 216 DIQMTQSPSS LSASVGDRVT ITCRSSQSLV HSNRNTYLHW YQQKPGKAPK LLIYKVSNRF 60
SGVPSRFSGS GSGTDFTFTI SSLQPEDIAT YYCGQGTQVP YTFGQGTKVE IK 112

SEQ ID NO: 217 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 217 DIVMTQSPLS LPVTPGEPAS ISCRASQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 218 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

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SEQUENCE: 218
DIVMTQSPLS LPVTPGEPAS ISCRISQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 219      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 219
DIVMTQSPLS LPVTPGEPAS ISCRPSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 220      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 220
DIVMTQSPLS LPVTPGEPAS ISCRTSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 221      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 221
DIVMTQSPLS LPVTPGEPAS ISCRVSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 222      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 222
DIVMTQSPLS LPVTPGEPAS ISCRSPQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 223      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 223
DIVMTQSPLS LPVTPGEPAS ISCRSSQPLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 224      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 224
DIVMTQSPLS LPVTPGEPAS ISCRSSQSFV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 225      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein

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SEQUENCE: 225          organism = synthetic construct
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 226          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 226          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 227          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 227          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 228          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 228          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 229          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 229          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV RSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 230          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 230          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV HFNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 231          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112
                      mol_type = protein
                      organism = synthetic construct
SEQUENCE: 231          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSIV HHNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 232          moltype = AA length = 112
FEATURE               Location/Qualifiers
REGION                1..112
source                 note = An artificially generated sequence
                      1..112

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mol_type = protein
organism = synthetic construct
SEQUENCE: 232
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HLNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 233      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 233
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HMNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 234      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 234
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HNNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 235      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 235
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HPNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 236      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 236
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HQNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 237      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 237
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HVNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 238      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 238
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HWNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 239      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence

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source          1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 239
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRDTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 240      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 240
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRDTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 241      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 241
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRFTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 242      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 242
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRFTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 243      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 243
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRHTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 244      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 244
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRMTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 245      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 245
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRWTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 246      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112

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source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 246
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 247      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 247
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYVHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 248      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 248
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLEW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 249      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 249
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLLW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 250      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 250
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLMW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 251      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 251
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLNW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 252      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 252
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLQW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 253      moltype = AA length = 112
FEATURE           Location/Qualifiers

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REGION          1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 253
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSHRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 254      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 254
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSHRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 255      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 255
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSHRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 256      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 256
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSMRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 257      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 257
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSTRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 258      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 258
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSWRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 259      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 259
DIVMTQSPSLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRA 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 260      moltype = AA length = 112

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FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 260
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRG 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 261      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 261
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRI 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 262      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 262
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRK 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 263      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 263
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRL 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 264      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 264
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRM 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 265      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 265
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRN 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 266      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION          1..112
source           note = An artificially generated sequence
                 1..112
                 mol_type = protein
                 organism = synthetic construct
SEQUENCE: 266
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRP 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

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SEQ ID NO: 267      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 267
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 268      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 268
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 269      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 269
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 270      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 270
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 271      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 271
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
GGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 272      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 272
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
HGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 273      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 273
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
IGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK       112

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SEQ ID NO: 274      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 274
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
LGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 275      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 275
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
MGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 276      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 276
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
PGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 277      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 277
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
QGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 278      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 278
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 279      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 279
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
VGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 280      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 280
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60

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WGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 281	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 281	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCAQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 282	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 282	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCEQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 283	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 283	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCFQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 284	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 284	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCIQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 285	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 285	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCLQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 286	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 286	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCNQGTQVP YTFGQGTKLE IK	112
<hr/>	
SEQ ID NO: 287	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 287	

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DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 288      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 288
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCVQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 289      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 289
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 290      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 290
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQSTQVP YTFGQGTKLE IK 112

SEQ ID NO: 291      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 291
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQTTQVP YTFGQGTKLE IK 112

SEQ ID NO: 292      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 292
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQFP YTFGQGTKLE IK 112

SEQ ID NO: 293      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 293
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQHP YTFGQGTKLE IK 112

SEQ ID NO: 294      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

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SEQUENCE: 294
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQNP YTFGQGTKLE IK 112

SEQ ID NO: 295      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 295
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQNP YTFGQGTKLE IK 112

SEQ ID NO: 296      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 296
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQWP YTFGQGTKLE IK 112

SEQ ID NO: 297      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 297
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQWP YTFGQGTKLE IK 112

SEQ ID NO: 298      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 298
DIVMTQSPLS LPVTPGEPAS ISCRPSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 299      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 299
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 300      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 300
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRWTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 301      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein

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SEQUENCE: 301
organism = synthetic construct
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 302
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 302
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 303
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 303
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
WGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 304
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 304
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 305
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 305
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 306
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 306
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
WGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 307
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 307
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 308
moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = An artificially generated sequence
source 1..112

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mol_type = protein
organism = synthetic construct
SEQUENCE: 308
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 309      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 309
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 310      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 310
DIVMTQSPLS LPVTPGEPAS ISCRPSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 311      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 311
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 312      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 312
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 313      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 313
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 314      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 314
DIVMTQSPLS LPVTPGEPAS ISCRSSQPLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 315      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence

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source          1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 315
DIVMTQSPS LPVTPGEPAS ISCRPSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 316      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 316
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 317      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 317
DIVMTQSPS LPVTPGEPAS ISCRSSQSLV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 318      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 318
DIVMTQSPS LPVTPGEPAS ISCRSSQPLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 319      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 319
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIOW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 320      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 320
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRL 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 321      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 321
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
PGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 322      moltype = AA  length = 112
FEATURE           Location/Qualifiers
REGION            1..112

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source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 322
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCIQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 323      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 323
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCLQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 324      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 324
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCVQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 325      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 325
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQSTQVP YTFGQGTKLE IK      112

SEQ ID NO: 326      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 326
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQTTQVP YTFGQGTKLE IK      112

SEQ ID NO: 327      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 327
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQWP YTFGQGTKLE IK      112

SEQ ID NO: 328      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 328
DIVMTQSPLS LPVTPGEPAS ISCRSSSES LV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK      112

SEQ ID NO: 329      moltype = AA length = 112
FEATURE          Location/Qualifiers

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REGION          1..112
source          note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 329
DIVMTQSPLS LPVTPGEPAS ISCRSSQELV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 330      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 330
DIVMTQSPLS LPVTPGEPAS ISCRSSQEPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 331      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 331
DIVMTQSPLS LPVTPGEPAS ISCRSSQEPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 332      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 332
DIVMTQSPLS LPVTPGEPAS ISCRSSQEPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
WGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 333      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 333
DIVMTQSPLS LPVTPGEPAS ISCRSSQEPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 334      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 334
DIVMTQSPLS LPVTPGEPAS ISCRSSSEPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 335      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source            note = An artificially generated sequence
                1..112
                mol_type = protein
                organism = synthetic construct
SEQUENCE: 335
DIVMTQSPLS LPVTPGEPAS ISCRSSSESPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK        112

SEQ ID NO: 336      moltype = AA length = 112

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FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 336	
DIVMTQSPLS LPVTPGEPAS ISCRSSSESPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 337	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 337	
DIVMTQSPLS LPVTPGEPAS ISCRSSSESPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF	60
WGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 338	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 338	
DIVMTQSPLS LPVTPGEPAS ISCRSSSESPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 339	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 339	
DIVMTQSPLS LPVTPGEPAS ISCRSSSEPPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF	60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 340	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 340	
DIVMTQSPLS LPVTPGEPAS ISCRSSSQSPV HSNRNTYIQW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCSQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 341	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 341	
DIVMTQSPLS LPVTPGEPAS ISCRSSSQSPV HSNRNTYIQW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCIQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 342	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
source	note = An artificially generated sequence 1..112 mol_type = protein organism = synthetic construct
SEQUENCE: 342	
DIVMTQSPLS LPVTPGEPAS ISCRPSQSPV HSNRNTYIQW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IK	112

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SEQ ID NO: 343      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 343
DIVMTQSPS LPVTPGEPAS ISCRPSQSPV HSNRNTYIOW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCSQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 344      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 344
DIVMTQSPS LPVTPGEPAS ISCRPSQSPV HSNRNTYIOW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCIQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 345      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 345
DIVMTQSPS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 346      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 346
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 347      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 347
DIVMTQSPS LPVTPGEPAS ISCRPSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 348      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 348
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIOW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQGTKLE IK       112

SEQ ID NO: 349      moltype = AA  length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = An artificially generated sequence
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 349
DIVMTQSPS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCAQGTQVP YTFGQGTKLE IK       112

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SEQ ID NO: 350      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 350
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCAQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 351      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 351
DIVMTQSPLS LPVTPGEPAS ISCRPSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCAQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 352      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 352
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCAQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 353      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 353
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCVQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 354      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 354
DIVMTQSPLS LPVTPGEPAS ISCRPSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCVQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 355      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 355
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCVQGTQVP YTFGQGTKLE IK           112

SEQ ID NO: 356      moltype = AA  length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct
SEQUENCE: 356
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60

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SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 357	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 357	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRQTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 358	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 358	
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRQTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 359	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 359	
DIVMTQSPLS LPVTPGEPAS ISCRSSQPLV HSNRQTYLHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 360	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 360	
DIVMTQSPLS LPVTPGEPAS ISCRQASESLV HSNRNTYLYHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 361	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 361	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLYHW YLQKPGQSPQ LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK	112
SEQ ID NO: 362	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 362	
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLYHW YQQKPGQAPR LLIYKVSNRF	60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKVE IE	112
SEQ ID NO: 363	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = An artificially generated sequence
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 363	

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DIVMTQSPLS LPVTPGEPAS ISQCASESLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKVE IE 112

SEQ ID NO: 364      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 364
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKVE IE 112

SEQ ID NO: 365      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 365
DIVMTQSPLS LPVTPGEPAS ISQCASESLV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKVE IE 112

SEQ ID NO: 366      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 366
DIVMTQSPLS LPVTPGEPAS ISQCASESLV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 367      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 367
DIVMTQSPLS LPVTPGEPAS ISQCASESLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 368      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 368
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IK 112

SEQ ID NO: 369      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 369
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKVE IE 112

SEQ ID NO: 370      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = An artificially generated sequence
source             1..112
mol_type = protein
organism = synthetic construct

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SEQUENCE: 370
DIVMTQSPLS LPVTPGEPAS ISQCQASESLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTTKVE IE 112

SEQ ID NO: 371      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 371
DIVMTQSPLS LPVTPGEPAS ISCRSSSQSLV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTTKVE IE 112

SEQ ID NO: 372      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 372
DIVMTQSPLS LPVTPGEPAS ISQCQASESLV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
RGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTTKVE IE 112

SEQ ID NO: 373      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 373
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTTKLE IK 112

SEQ ID NO: 374      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 374
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTTKLE IK 112

SEQ ID NO: 375      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 375
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIOW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTTKLE IK 112

SEQ ID NO: 376      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 376
DIVMTQSPLS LPVTPGEPAS ISQCQASESPV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTTKLE IK 112

SEQ ID NO: 377      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
source             note = An artificially generated sequence
                  1..112
mol_type = protein

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SEQUENCE: 377          organism = synthetic construct
DIVMTQSPLS LPVTPGEPAS ISQCQASESPV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQQGKLE IK           112

SEQ ID NO: 378          moltype = AA length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 378          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISQCQASESPV HSNRNTYIOW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQQGKLE IK           112

SEQ ID NO: 379          moltype = AA length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 379          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQQGKLE IK           112

SEQ ID NO: 380          moltype = AA length = 112
FEATURE
REGION
1..112
note = An artificially generated sequence
source
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 380          moltype = AA length = 112
DIVMTQSPLS LPVTPGEPAS ISQCQASESPV HSNRNTYLHW YLQKPGQSPQ LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCTQGTQVP YTFGQQGKLE IK           112

SEQ ID NO: 381          moltype = AA length = 458
FEATURE
REGION
1..458
note = An artificially generated sequence
source
1..458
mol_type = protein
organism = synthetic construct

SEQUENCE: 381          moltype = AA length = 458
QVQLVESGGG VVQPGRSRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAN IKAKSNNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQMSNLKT EDTAVYYCERY VHYGAYYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPSSKST SGGTAALGCL VKDYFPEPVT VSWNSGALTS GVHTPPAVLQ 180
SSGLYSLSSV VTVPSSSLGT QTYICNVNHK PSNTKVDKKV EPKSCDKTHT CPPCPAPEAA 240
GGPSVFLPFP KPKDTLMISR TPEVITCVVVD VSHEDPFVVF NWYVDGVEVH NAKTKPREEQ 300
YASTYRVVSV LTVLHQDWLN GKEYKCKVSN KALPAPIEKT ISKAKGQPRE PQVYTLPPSR 360
CELTKNQVSL SCAVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LVSKLTVDKS 420
RWQQGNVFSC SVMHEALHNN YTQKSLSLSP DYKDDDDK                         458

SEQ ID NO: 382          moltype = AA length = 458
FEATURE
REGION
1..458
note = An artificially generated sequence
source
1..458
mol_type = protein
organism = synthetic construct

SEQUENCE: 382          moltype = AA length = 458
QVQLVESGGG VVQPGRSRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQMSNLKT EDTAVYYCERY VHYYAYYGVQ AWGQGTTVT 120
SSASTKGPSV FPLAPSSKST SGGTAALGCL VKDYFPEPVT VSWNSGALTS GVHTPPAVLQ 180
SSGLYSLSSV VTVPSSSLGT QTYICNVNHK PSNTKVDKKV EPKSCDKTHT CPPCPAPEAA 240
GGPSVFLPFP KPKDTLMISR TPEVITCVVVD VSHEDPFVVF NWYVDGVEVH NAKTKPREEQ 300
YASTYRVVSV LTVLHQDWLN GKEYKCKVSN KALPAPIEKT ISKAKGQPRE PQVYTLPPSR 360
CELTKNQVSL SCAVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LVSKLTVDKS 420
RWQQGNVFSC SVMHEALHNN YTQKSLSLSP DYKDDDDK                         458

SEQ ID NO: 383          moltype = AA length = 458
FEATURE
REGION
1..458
note = An artificially generated sequence

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source          1..458
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 383
QVQLVESGGG VVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWAQ IAKASNYYAT 60
YYAEHSVGRF TISRDSKNS LYLMQNSLKT EDTAVYYC TRFY VHYYYYYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPSSKST SGGTAALGCL VKDYFPEPVY VSWNSGALTS GVHTFPAVLQ 180
SSGLYSLSV VTVPSSSLGT QTYICVNHNK PSNTKVDKVV EPKSCDKTH' CPPCPAPEAA 240
GGPSVFLFPP KPKDTLMISR TPEVTCVVVD VSHEDPEVKF NWYVDGVEVH NAKTKPREEQ 300
YASTYRVVSV LTVLHQDWLN GKEYKCKVSN KALPAPIEKT ISKAKGQPRE PQVYTLPPSR 360
CELTKNQVSL SCAVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LVSKLTVDKS 420
RWQQGNVFSC SVMHEALHNH YTQKSLSLSP DYKDDDDK 458

SEQ ID NO: 384      moltype = AA length = 440
FEATURE           Location/Qualifiers
REGION            1..440
note = An artificially generated sequence
source             1..440
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 384
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGTPDTAY 60
SEKFKGRTVL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQGTL VTVSSASTKG 120
PSVFPLAPCS RSTSESTAAL GCLVKDYPFPE PVTVSWNSGA LTSGVHTFPV VLQSSGLYSL 180
SSVTVPSSS LGTQTYTCNV DHKPSNTKVD KRVESKYGPP CPPCPAPEFR GGPKVFLFPP 240
KPKDTLMISR TPEVTCVVVD VSQEDPEVQF NWYVDGVEVH NAKTKPREEQ FASTYRVVSV 300
LTVLHQDWLN GKEYKCKVSN KGLPSSIEKT ISKAKGQPRE PQVYTLPPSQ EEMTKNQVSL 360
TCLVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LYSKLTVDKS RWQEGNVFSC 420
SVMHEALHNH YTQESLSSLSP 440

SEQ ID NO: 385      moltype = AA length = 440
FEATURE           Location/Qualifiers
REGION            1..440
note = An artificially generated sequence
source             1..440
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 385
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGEGLEWIGA IDGTPDTAY 60
SEKFKGRTVL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQGTL VTVSSASTKG 120
PSVFPLAPCS RSTSESTAAL GCLVKDYPFPE PVTVSWNSGA LTSGVHTFPV VLQSSGLYSL 180
SSVTVPSSS LGTQTYTCNV DHKPSNTKVD KRVESKYGPP CPPCPAPEFR GGPKVFLFPP 240
KPKDTLMISR TPEVTCVVVD VSQEDPEVQF NWYVDGVEVH NAKTKPREEQ FASTYRVVSV 300
LTVLHQDWLN GKEYKCKVSN KGLPSSIEKT ISKAKGQPRE PQVYTLPPSQ EEMTKNQVSL 360
TCLVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LYSKLTVDKS RWQEGNVFSC 420
SVMHEALHNH YTQESLSSLSP 440

SEQ ID NO: 386      moltype = AA length = 440
FEATURE           Location/Qualifiers
REGION            1..440
note = An artificially generated sequence
source             1..440
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 386
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQA PGEGLEWMGA IDGTPDTAY 60
SEKFKGRTVL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQGTL VTVSSASTKG 120
PSVFPLAPCS RSTSESTAAL GCLVKDYPFPE PVTVSWNSGA LTSGVHTFPV VLQSSGLYSL 180
SSVTVPSSS LGTQTYTCNV DHKPSNTKVD KRVESKYGPP CPPCPAPEFR GGPKVFLFPP 240
KPKDTLMISR TPEVTCVVVD VSQEDPEVQF NWYVDGVEVH NAKTKPREEQ FASTYRVVSV 300
LTVLHQDWLN GKEYKCKVSN KGLPSSIEKT ISKAKGQPRE PQVYTLPPSQ EEMTKNQVSL 360
TCLVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LYSKLTVDKS RWQEGNVFSC 420
SVMHEALHNH YTQESLSSLSP 440

SEQ ID NO: 387      moltype = AA length = 440
FEATURE           Location/Qualifiers
REGION            1..440
note = An artificially generated sequence
source             1..440
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 387
QVQLVQSGAE VKKPGASVTV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDGKTPDTAY 60
SQKFKGRTVL TADKSTSTAY MELSSLTSED TAVYYCTRFY SYTYWGQGTL VTVSSASTKG 120
PSVFPLAPCS RSTSESTAAL GCLVKDYPFPE PVTVSWNSGA LTSGVHTFPV VLQSSGLYSL 180
SSVTVPSSS LGTQTYTCNV DHKPSNTKVD KRVESKYGPP CPPCPAPEFR GGPKVFLFPP 240

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KKPKDTLMISR	TPEVTCVVVD	VSQEDPEVQF	NWYVDGVEVH	NAKTKPREEQ	FASTYRVVSV	300
LTVLHQDWLN	GKEYKCVSN	KGLPSSIEKT	ISAKAGQPRE	PQVYTLPPSQ	EEMTKNQVSL	360
TCLVKGFYPS	DAIWEWESNG	QPENNYKTTP	PVLSDSGSFF	LYSKLTVDKS	RWQEGNVFSC	420
SVMHEALHNH	YTQESLSLSP					440

SEQ ID NO: 388	moltype = AA length = 458	
FEATURE	Location/Qualifiers	
REGION	1..458	
	note = An artificially generated sequence	
source	1..458	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 388		
QVQLVESGGG VVQPGRLRLR	SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNYYAT	60
YYAESVKGFR TISRDDSKNS	LYLQMNSLKT EDTAVYYCRY VHGYGAYGVD AWGQGTTVT	120
SSASTKGPSV FPLAPSSKST	SGGTAALGCL VKDYFPEPV VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV VTVPSSSLGT	QTYICNVNHK PSNTKVDDKKV EPKSCDKTHT CPPCPAPEAA	240
GGPSVFLFPP KPKDTLMISR	TPEVTCVVVD VSHEDPEVKF NWYVDGVEVH NAKTKPREEQ	300
YASTYRVVSV LTVLHQDWLN	GKEYKCVSN KALPAPIEKT ISAKAGQPRE PQVYTLPPSR	360
CETLTKNQVSL SCAVKGFYPS	DAIWEWESNG QPENNYKTTP PVLSDGSFF LVSKLTVDKS	420
RWQQGNVFSC SVMHEALHNH	YTQKSLSLSP DYKDDDDK	458

SEQ ID NO: 389	moltype = AA length = 458	
FEATURE	Location/Qualifiers	
REGION	1..458	
	note = An artificially generated sequence	
source	1..458	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 389		
EVQLVETGGS LVQPGKSLKL	TCATSGFTFS NAWMHWVRQS PEKQLEWVAQ IKAKSNYYAT	60
YYAESVKGFR TISRDDSKSN	IYLQMNSLKE EDTAVYYCRY VHGYGAYGVD AWGQGTSVT	120
SSASTKGPSV FPLAPSSKST	SGGTAALGCL VKDYFPEPV VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV VTVPSSSLGT	QTYICNVNHK PSNTKVDDKKV EPKSCDKTHT CPPCPAPEAA	240
GGPSVFLFPP KPKDTLMISR	TPEVTCVVVD VSHEDPEVKF NWYVDGVEVH NAKTKPREEQ	300
YASTYRVVSV LTVLHQDWLN	GKEYKCVSN KALPAPIEKT ISAKAGQPRE PQVYTLPPSR	360
CETLTKNQVSL SCAVKGFYPS	DAIWEWESNG QPENNYKTTP PVLSDGSFF LVSKLTVDKS	420
RWQQGNVFSC SVMHEALHNH	YTQKSLSLSP DYKDDDDK	458

SEQ ID NO: 390	moltype = AA length = 447	
FEATURE	Location/Qualifiers	
REGION	1..447	
	note = An artificially generated sequence	
source	1..447	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 390		
EVQLVETGGS LVQPGKSLKL	TCATSGFTFS NAWMHWVRQS PEKQLEWVAQ IKAKSNYYAT	60
YYAESVKGFR TISRDDSKSN	IYLQMNSLKE EDTAVYYCRY VHGYGAYGVD AWGQGTSVT	120
SSASTKGPSV FPLAPCSRST	SESTAALGCL VKDYFPEPV VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV VTVPSSSLGT	KTYTCNVDHK PSNTKVDDKKV ESKYGPCCPP CPAPEFRGGP	240
KVLFPPPKP KPKDTLMISR	TPEVTCVVVD EDPEVQFNWY VDGVEVHNAK TKPREEQFAS	300
TYRVSVSLTV LHQDWLNGKE	YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPSSQEM	360
TKNQVSLTCL VKGFYPSDIA	VEWESNGQPE NNYKTPPPVLDSDGSFFLYS KLTVDKSRWQ	420
EGNVFSCSVM HEALHNRYTQ	KSLSLSP	447

SEQ ID NO: 391	moltype = AA length = 458
FEATURE	Location/Qualifiers
REGION	1..458
	note = An artificially generated sequence
source	1..458
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 391	

QVQLVESGGG VVQPGRLRLR	SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNYYKK	60
YYAESVKGFR TISRDDSKNS	LYLQMNSLKT EDTAVYYCRY VHYYYYYGVQ AWGQGTTVT	120
SSASTKGPSV FPLAPSSKST	SGGTAALGCL VKDYFPEPV VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV VTVPSSSLGT	QTYICNVNHK PSNTKVDDKKV EPKSCDKTHT CPPCPAPEAA	240
GGPSVFLFPP KPKDTLMISR	TPEVTCVVVD VSHEDPEVKF NWYVDGVEVH NAKTKPREEQ	300
YASTYRVVSV LTVLHQDWLN	GKEYKCVSN KALPAPIEKT ISAKAGQPRE PQVYTLPPSR	360
CETLTKNQVSL SCAVKGFYPS	DAIWEWESNG QPENNYKTTP PVLSDGSFF LVSKLTVDKS	420
RWQQGNVFSC SVMHEALHNH	YTQKSLSLSP DYKDDDDK	458

SEQ ID NO: 392	moltype = AA length = 458
FEATURE	Location/Qualifiers
REGION	1..458

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source          note = An artificially generated sequence
                1..458
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 392
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQQYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPCSRST SGGTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPALQ 180
SSGLYSLSSV VTVPSSSLGT QTYICNVNHK PSNTKVDKKV EPKSCDKTHT CPPCPAPEAA 240
GGPSVFLPP KPDKDTLMISRTPE TPEVTCVVVDVQFNWY EDPEVQFVHNAK TKPREEQFAS 300
YASTYRVSVS LTVLHQDWLNL GKEYKCKVSN KALPAPIEKT ISKAKQPRE PQVYTLPPSR 360
CELTKNQVSL CCAVKGFYPS DIAVEWESNG QPENNYKTPP VLDSDGSFF LVSKLTVDKS 420
RWQQGNVFSC SVMHEALHNH YTQKSLSLSP DYKDDDDK 458

SEQ ID NO: 393      moltype = AA length = 447
FEATURE          Location/Qualifiers
REGION           1..447
source          note = An artificially generated sequence
                1..447
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 393
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQQYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPALQ 180
SSGLYSLSSV VTVPSSSLGT KTYTCNVNDHK PSNTKVDKRV ESKYGPCCPP CPAPEFRGGP 240
KVFLFPKPK DTLMISRTPE VTCVVVDVQS EDPEVQFVHNAK TKPREEQFAS 300
TYRVSVSLTV LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM 360
TKNQVSLTCL VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ 420
EGNVFSCSVM HEALHNRYTQ KSLSLSP 447

SEQ ID NO: 394      moltype = AA length = 447
FEATURE          Location/Qualifiers
REGION           1..447
source          note = An artificially generated sequence
                1..447
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 394
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQQYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPALQ 180
SSGLYSLSSV VTVPSSSLGT KTYTCNVNDHK PSNTKVDKRV ESKYGPCCPP CPAPEFRGGP 240
KVFLFPKPK DTLMISRTPE VTCVVVDVQS EDPEVQFVHNAK TKPREEQFAS 300
TYRVSVSLTV LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM 360
TKNQVSLTCL VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ 420
EGNVFSCSVM HEALHNRYTQ KSLSLSP 447

SEQ ID NO: 395      moltype = AA length = 447
FEATURE          Location/Qualifiers
REGION           1..447
source          note = An artificially generated sequence
                1..447
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 395
QVQLVESGGG LVQPGRSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQQYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPALQ 180
SSGLYSLSSV VTVPSSSLGT KTYTCNVNDHK PSNTKVDKRV ESKYGPCCPP CPAPEFRGGP 240
KVFLFPKPK DTLMISRTPE VTCVVVDVQS EDPEVQFVHNAK TKPREEQFAS 300
TYRVSVSLTV LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM 360
TKNQVSLTCL VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ 420
EGNVFSCSVM HEALHNRYTQ KSLSLSP 447

SEQ ID NO: 396      moltype = AA length = 447
FEATURE          Location/Qualifiers
REGION           1..447
source          note = An artificially generated sequence
                1..447
                mol_type = protein
                organism = synthetic construct

SEQUENCE: 396
QVQLVESGGG VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQQYAT 60
YYAESVKGRF TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYSASYGVD AWGQGTTVT 120
SSASTKGPSV FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPALQ 180

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SSGLYSLSSV	VTVPSSSLGT	KTYTCNVDHK	PSNTKVDKRV	ESKYGPPCPP	CAPAEPFRGGP	240
KVFLFPPKPK	DTLMISRTPE	VTCVVVDVSQ	EDPEVQFNWY	VDGVEVHNAK	TKPREEQFAS	300
TYRVSVLTV	LHQDWLNGKE	YKCKVSNKGL	PSSIEKTISK	AKGQPREPQV	YTLPPSQKEM	360
TKNQVSLTCL	VKGFYPSDIA	VEWESNGQPE	NNYKTTPPVVL	DSDGSFFLYS	KLTVDKSRWQ	420
EGNVFSCSVM	HEALHNRYTQ	KSLSLSP				447

SEQ ID NO: 397	moltype = AA length = 447	
FEATURE	Location/Qualifiers	
REGION	1..447	
note	An artificially generated sequence	
source	1..447	
mol_type	protein	
organism	synthetic construct	
SEQUENCE: 397		
QVQLVESGGV	LVQPGRLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSNNYAT	60
YYAESVKGRF	TISRADSKNS IYLQMNSLKT EDTAVYYCRY VHYAASYGVD IWGQGTTVT	120
SSASTKGPSV	FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV	VTVPSSSLGT KTYTCNVDHK PSNTKVDKRV ESKYGPPCPP CPAPEFRGGP	240
KVFLFPPKPK	DTLMISRTPE VTCVVVDVSQ EDPEVQFNWY VDGVEVHNAK TKPREEQFAS	300
TYRVSVLTV	LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM	360
TKNQVSLTCL	VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ	420
EGNVFSCSVM	HEALHNRYTQ KSLSLSP	

SEQ ID NO: 398	moltype = AA length = 447	
FEATURE	Location/Qualifiers	
REGION	1..447	
note	An artificially generated sequence	
source	1..447	
mol_type	protein	
organism	synthetic construct	
SEQUENCE: 398		
QVQLVESGGV	LVQPGRLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQQNYAT	60
YYAESVKGRF	TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT	120
SSASTKGPSV	FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV	VTVPSSSLGT KTYTCNVDHK PSNTKVDKRV ESKYGPPCPP CPAPEFRGGP	240
KVFLFPPKPK	DTLMISRTPE VTCVVVDVSQ EDPEVQFNWY VDGVEVHNAK TKPREEQFAS	300
TYRVSVLTV	LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM	360
TKNQVSLTCL	VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ	420
EGNVFSCSVM	HEALHNRYTQ KSLSLSP	

SEQ ID NO: 399	moltype = AA length = 447	
FEATURE	Location/Qualifiers	
REGION	1..447	
note	An artificially generated sequence	
source	1..447	
mol_type	protein	
organism	synthetic construct	
SEQUENCE: 399		
QVQLVESGGV	VVQPGGSLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT	60
YYAESVKGRF	TISRDDSKNS IYLQMNSLKT EDTAVYYCRY VHYAAGYGVD IWGQGTTVT	120
SSASTKGPSV	FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV	VTVPSSSLGT KTYTCNVDHK PSNTKVDKRV ESKYGPPCPP CPAPEFRGGP	240
KVFLFPPKPK	DTLMISRTPE VTCVVVDVSQ EDPEVQFNWY VDGVEVHNAK TKPREEQFAS	300
TYRVSVLTV	LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM	360
TKNQVSLTCL	VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ	420
EGNVFSCSVM	HEALHNRYTQ KSLSLSP	

SEQ ID NO: 400	moltype = AA length = 447
FEATURE	Location/Qualifiers
REGION	1..447
note	An artificially generated sequence
source	1..447
mol_type	protein
organism	synthetic construct
SEQUENCE: 400	

QVQLVESGGV	LVQPGRLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKDKSQNYAT	60
YYAESVKGRF	TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYAAGYGVD AWGQGTTVT	120
SSASTKGPSV	FPLAPCSRST SESTAALGCL VKDYFPEPVT VSWNSGALTS GVHTFPAVLQ	180
SSGLYSLSSV	VTVPSSSLGT KTYTCNVDHK PSNTKVDKRV ESKYGPPCPP CPAPEFRGGP	240
KVFLFPPKPK	DTLMISRTPE VTCVVVDVSQ EDPEVQFNWY VDGVEVHNAK TKPREEQFAS	300
TYRVSVLTV	LHQDWLNGKE YKCKVSNKGL PSSIEKTISK AKGQPREPQV YTLPPSQKEM	360
TKNQVSLTCL	VKGFYPSDIA VEWESNGQPE NNYKTTPPVVL DSDGSFFLYS KLTVDKSRWQ	420
EGNVFSCSVM	HEALHNRYTQ KSLSLSP	

SEQ ID NO: 401	moltype = AA length = 447
FEATURE	Location/Qualifiers

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REGION
source note = An artificially generated sequence
SEQUENCE: 401 mol_type = protein
organism = synthetic construct

QVQLVESGGG VVQPGGSSLRL SCAASGFTFS NAWMHWRQQA PGKGLEWVAQ IKDKSQQNYAT
YVAESVKGRF TISRADSKNS IYLOQMSNLKT EDTAVYYCRY VHYAAGYGV D WGQGTTVT
SSASTKGPSV FPLAPCSRST SESTAALGLC VKDYFPPEVT VSWNSGALT GVHTPAVLQ
SSGLYSLSSV VTVPSLSSLTG KTYTCVNVDHK PSNTVKDVRV ESKYGPPCPP CPAPEFRGGP
KVFLFPPKPK DTLMISRTPE VTCVVVVDVSQ EDPEVQFNWY VDGVEVHNKA TKPREEQFAS
TYRVSVLTV LHQDWLNGKE YKCKVSNKGL PSSIEKTQAK GQPREPQV YTLPFSQKEM
TKNQVSLTCL VKGFYPSDIA VEWESENQPE NNYKTPPVL DSDGSFFLYS KLTVDKSRWQ
EGNVFSCSVM HEALHNRYTQ KSLSLSP 60
120
180
240
300
360
420
480
447

SEQ ID NO: 402 moltype = AA length = 447
FEATURE Location/Qualifiers
REGION 1..447
source note = An artificially generated sequence
1..447
mol_type = protein
organism = synthetic construct

SEQUENCE: 402
QVQLVESGGG VVQPGGSSLRL SCAASGFTFS NAWMHWRQQA PGKGLEWVAQ IKDKSQQNYAT
YVAESVKGRF TISRADSKNS IYLOQMSNLKT EDTAVYYCRY VHYAAGYGV D WGQGTTVT
SSASTKGPSV FPLAPCSRST SESTAALGLC VKDYFPPEVT VSWNSGALT GVHTPAVLQ
SSGLYSLSSV VTVPSLSSLTG KTYTCVNVDHK PSNTVKDVRV ESKYGPPCPP CPAPEFRGGP
KVFLFPPKPK DTLMISRTPE VTCVVVVDVSQ EDPEVQFNWY VDGVEVHNKA TKPREEQFAS
TYRVSVLTV LHQDWLNGKE YKCKVSNKGL PSSIEKTQAK GQPREPQV YTLPFSQKEM
TKNQVSLTCL VKGFYPSDIA VEWESENQPE NNYKTPPVL DSDGSFFLYS KLTVDKSRWQ
EGNVFSCSVM HEALHNHYTQ KSLSLSP 60
120
180
240
300
360
420
480
447

SEQ ID NO: 403 moltype = AA length = 450
FEATURE Location/Qualifiers
REGION 1..450
source note = An artificially generated sequence
1..450
mol_type = protein
organism = synthetic construct

SEQUENCE: 403
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF
SGVPDRFSGS GSGTDFTLKI SRVEAEDVG YYCSQNTHVP PTFGQGTKLE IKSSASTKGP
SVFPLAPSSK STSGGTAALG CLVKDYFPPEP VTVSWNSGAL TSGVHTFPAV LQSSGLYSL
SVTVTPSSSL GTQTYICCNV HKPSNTKVDK KVEPKSCDKT HTCPCCPAPE AAGGSPVFLF
PPKPKDTLMI SRTPEVTCVV VDVSHEDEPV KFNNWYVDGVE VHNAKTKPRE EQYASTYRV
SVLTVLHQDW LNGKEYKCKV SNKALPAPIE KTISKAKQGP REPQVCTLPP SRDELTKNQV
SLWCLVKGFT PSDIAVEWES NGQPENNYKT TPPVLDSDGS FFLYSKLTVD KSRWQOQN
SCSVMHEALTH NYHTQKSLSL SPHHHHHHHHH 60
120
180
240
300
360
420
480
450

SEQ ID NO: 404 moltype = AA length = 450
FEATURE Location/Qualifiers
REGION 1..450
source note = An artificially generated sequence
1..450
mol_type = protein
organism = synthetic construct

SEQUENCE: 404
DIVMTQSPLS LPVTPGEPAS ISCRSSQSLV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF
SGVPDRFSGS GSGTDFTLKI SRVEAEDVG YYCGQGTQV YTFGQGTKLE IKSSASTKGP
SVFPLAPSSK STSGGTAALG CLVKDYFPPEP VTVSWNSGAL TSGVHTFPAV LQSSGLYSL
SVTVTPSSSL GTQTYICCNV HKPSNTKVDK KVEPKSCDKT HTCPCCPAPE AAGGSPVFLF
PPKPKDTLMI SRTPEVTCVV VDVSHEDEPV KFNNWYVDGVE VHNAKTKPRE EQYASTYRV
SVLTVLHQDW LNGKEYKCKV SNKALPAPIE KTISKAKQGP REPQVCTLPP SRDELTKNQV
SLWCLVKGFT PSDIAVEWES NGQPENNYKT TPPVLDSDGS FFLYSKLTVD KSRWQOQN
SCSVMHEALTH NYHTQKSLSL SPHHHHHHHHH 60
120
180
240
300
360
420
480
450

SEQ ID NO: 405 moltype = AA length = 222
FEATURE Location/Qualifiers
REGION 1..222
source note = An artificially generated sequence
1..222
mol_type = protein
organism = synthetic construct

SEQUENCE: 405
QVQLVQSGAE VKKPGAVKV SCKASGYTFT DYEMHWIRQP PGQGLEWIGA IDPKTGDTAY
SQKFKGRVTL TADKSTSTAY MELSSLTSBD TAVYYCTRFY SYTYWCGT VTVSSASVAA 60
120

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PSVFIFPPSD	EQLKSGTASV	VCLLNNFYPR	EAKVQWKVDN	ALQSGNSQES	VTEQDSKDST	180
YSLSSTLTL	S KADYEKHKVY	ACEVTHQGLS	SPVTKSFNRG	EC		222
 SEQ ID NO: 406						
FEATURE	moltype = AA length = 222					
REGION	Location/Qualifiers					
1..222						
source	note = An artificially generated sequence					
1..222						
	mol_type = protein					
	organism = synthetic construct					
 SEQUENCE: 406						
QVQLVQSGAE	VKKPGAVKV	SCKASGYTFT	DYEMHWIRQP	PGQGLEWIGA	IDGKTPDTAY	60
SQKFKGRTVL	TADKSTSTAY	MELSSLTSED	TAVYYC TRFY	SYTYWGQGTL	TVTSSASVAA	120
PSVFIFPPSD	EQLKSGTASV	VCLLNNFYPR	EAKVQWKVDN	ALQSGNSQES	VTEQDSKDST	180
YSLSSTLTL	S KADYEKHKVY	ACEVTHQGLS	SPVTKSFNRG	EC		222
 SEQ ID NO: 407						
FEATURE	moltype = AA length = 219					
REGION	Location/Qualifiers					
1..219						
source	note = An artificially generated sequence					
1..219						
	mol_type = protein					
	organism = synthetic construct					
 SEQUENCE: 407						
DVVMQTPTVS	MSVSLGGQVS	ISCRSSQS LV	HNNGN TYVSW	YI QKPSQSPQ	LLIYKVSNRF	60
SGISDRFSGS	GSGTDFTLKI	SRVEPDDLG V	YYCGQGTQDP	YTFGAGTKLE	LKRTVAAPSV	120
FIFPPSDEQL	KSGTASVVCL	LNNFYPREAK	VQWKVDNALQ	SGNSQESVTE	QDSKDSTYSL	180
STSTLTL SKAD	YEKHKVYACE	VTHQGLSSPV	TKSFNRGEC			219
 SEQ ID NO: 408						
FEATURE	moltype = AA length = 219					
REGION	Location/Qualifiers					
1..219						
source	note = An artificially generated sequence					
1..219						
	mol_type = protein					
	organism = synthetic construct					
 SEQUENCE: 408						
DIVMTQSPLS	LPVTPGE PAS	ISCRSSQS LV	HSNRNTYL SW	YQQKPGQAPR	LLIYKVSNRF	60
SGVPDRFSGS	GSGTDFTLKI	SRVEAEDVG V	YYCGQGTQVP	YTFGQGT KLE	IKRTVAAPSV	120
FIFPPSDEQL	KSGTASVVCL	LNNFYPREAK	VQWKVDNALQ	SGNSQESVTE	QDSKDSTYSL	180
STSTLTL SKAD	YEKHKVYACE	VTHQGLSSPV	TKSFNRGEC			219
 SEQ ID NO: 409						
FEATURE	moltype = AA length = 219					
REGION	Location/Qualifiers					
1..219						
source	note = An artificially generated sequence					
1..219						
	mol_type = protein					
	organism = synthetic construct					
 SEQUENCE: 409						
DIVMTQSPLS	LPVTPGE PAS	ISCRSSQS LV	HSNRNTYL HW	YQQKPGQAPR	LLIYKVSNRF	60
SGVPDRFSGS	GSGTDFTLKI	SRVEAEDVG V	YYCGQGTQVP	YTFGQGT KLE	IKRTVAAPSV	120
FIFPPSDEQL	KSGTASVVCL	LNNFYPREAK	VQWKVDNALQ	SGNSQESVTE	QDSKDSTYSL	180
STSTLTL SKAD	YEKHKVYACE	VTHQGLSSPV	TKSFNRGEC			219
 SEQ ID NO: 410						
FEATURE	moltype = AA length = 219					
REGION	Location/Qualifiers					
1..219						
source	note = An artificially generated sequence					
1..219						
	mol_type = protein					
	organism = synthetic construct					
 SEQUENCE: 410						
DIVMTQSPLS	LPVTPGE PAS	ISCRSSQPLV	HSNRNTYL HW	YQQKPGQAPR	LLIYKVSNRF	60
SGVPDRFSGS	GSGTDFTLKI	SRVEAEDVG V	YYCGQGTQVP	YTFGQGT KLE	IKRTVAAPSV	120
FIFPPSDEQL	KSGTASVVCL	LNNFYPREAK	VQWKVDNALQ	SGNSQESVTE	QDSKDSTYSL	180
STSTLTL SKAD	YEKHKVYACE	VTHQGLSSPV	TKSFNRGEC			219
 SEQ ID NO: 411						
FEATURE	moltype = AA length = 219					
REGION	Location/Qualifiers					
1..219						
source	note = An artificially generated sequence					
1..219						
	mol_type = protein					
	organism = synthetic construct					
 SEQUENCE: 411						

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DIVMTQSPS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60	
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IKRTVAAPSV 120	
FIFPPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180	
SSTLTLSKAD YEHKHVYACE VTHQGLSSPV TKSFNRGEC 219	
 SEQ ID NO: 412 moltype = AA length = 219	
FEATURE Location/Qualifiers	
REGION 1..219	
source note = An artificially generated sequence	
1..219 mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 412	
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60	
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCGQGTQVP YTFGQGTKLE IKRTVAAPSV 120	
FIFPPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180	
SSTLTLSKAD YEHKHVYACE VTHQGLSSPV TKSFNRGEC 219	
 SEQ ID NO: 413 moltype = AA length = 219	
FEATURE Location/Qualifiers	
REGION 1..219	
source note = An artificially generated sequence	
1..219 mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 413	
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60	
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120	
FIFPPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180	
SSTLTLSKAD YEHKHVYACE VTHQGLSSPV TKSFNRGEC 219	
 SEQ ID NO: 414 moltype = AA length = 219	
FEATURE Location/Qualifiers	
REGION 1..219	
source note = An artificially generated sequence	
1..219 mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 414	
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60	
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120	
FIFPPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180	
SSTLTLSKAD YEHKHVYACE VTHQGLSSPV TKSFNRGEC 219	
 SEQ ID NO: 415 moltype = AA length = 219	
FEATURE Location/Qualifiers	
REGION 1..219	
source note = An artificially generated sequence	
1..219 mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 415	
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60	
WGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120	
FIFPPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180	
SSTLTLSKAD YEHKHVYACE VTHQGLSSPV TKSFNRGEC 219	
 SEQ ID NO: 416 moltype = AA length = 219	
FEATURE Location/Qualifiers	
REGION 1..219	
source note = An artificially generated sequence	
1..219 mol_type = protein	
organism = synthetic construct	
 SEQUENCE: 416	
DIVMTQSPS LPVTPGEPAS ISCRSSQSPV HSNRNTYIHW YQQKPGQAPR LLIYKVSNRF 60	
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120	
FIFPPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180	
SSTLTLSKAD YEHKHVYACE VTHQGLSSPV TKSFNRGEC 219	
 SEQ ID NO: 417 moltype = AA length = 219	
FEATURE Location/Qualifiers	
REGION 1..219	
source note = An artificially generated sequence	
1..219 mol_type = protein	

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SEQUENCE: 417 organism = synthetic construct
DIVMTQSPLS LPVTPGEPAS ISCRSSQPPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120
FIFPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180
SSTLTL SKAD YEKHKVYACE VTHQGLSSPV TKSFNRGEC 219

SEQ ID NO: 418 moltype = AA length = 219
FEATURE Location/Qualifiers
REGION 1..219
note = An artificially generated sequence
source 1..219
mol_type = protein
organism = synthetic construct

SEQUENCE: 418
DIVMTQSPLS LPVTPGEPAS ISCRPSQSPV HSNRNTYLHW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120
FIFPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180
SSTLTL SKAD YEKHKVYACE VTHQGLSSPV TKSFNRGEC 219

SEQ ID NO: 419 moltype = AA length = 219
FEATURE Location/Qualifiers
REGION 1..219
note = An artificially generated sequence
source 1..219
mol_type = protein
organism = synthetic construct

SEQUENCE: 419
DIVMTQSPLS LPVTPGEPAS ISCRSSQSPV HSNRNTYIOW YQQKPGQAPR LLIYKVSNRF 60
SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCYQGTQVP YTFGQGTKLE IKRTVAAPSV 120
FIFPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180
SSTLTL SKAD YEKHKVYACE VTHQGLSSPV TKSFNRGEC 219

SEQ ID NO: 420 moltype = AA length = 122
FEATURE Location/Qualifiers
REGION 1..122
note = An artificially generated sequence
source 1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 420
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYYYYYGVQ AWGQGTTVT 120
SS 122

SEQ ID NO: 421 moltype = AA length = 122
FEATURE Location/Qualifiers
REGION 1..122
note = An artificially generated sequence
source 1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 421
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYKK 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYYYYYGVQ AWGQGTTVT 120
SS 122

SEQ ID NO: 422 moltype = AA length = 122
FEATURE Location/Qualifiers
REGION 1..122
note = An artificially generated sequence
source 1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 422
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWVRQA PGKGLEWVAQ IKAKSNNYAT 60
YYAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRR VHYGAYYGVD AWGQGTTVT 120
SS 122

SEQ ID NO: 423 moltype = AA length = 122
FEATURE Location/Qualifiers
REGION 1..122
note = An artificially generated sequence
source 1..122
mol_type = protein
organism = synthetic construct

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SEQUENCE: 423
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQA PGKGLEWVAQ IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRR VHYYYYYGVG AWGQGTTVT 120
SS 122

SEQ ID NO: 424      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 424
EVOLVETGGS LVQPGKSLKL TCATSGFTFS NAWMHWRQGS PEKQLEWVAQ IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS IYLQMNSLKE EDTAVYYCRY VHYGAYYGVG AWGQGTSVT 120
SS 122

SEQ ID NO: 425      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 425
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQA PGKGLEWVAL IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYGAYYGVG AWGQGTTVT 120
SS 122

SEQ ID NO: 426      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 426
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQA PGKGLEWVAN IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYGAYYGVG AWGQGTTVT 120
SS 122

SEQ ID NO: 427      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 427
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQA PGKGLEWVAN IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYGAYYGVG AWGQGTTVT 120
SS 122

SEQ ID NO: 428      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 428
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQA PGKGLEWVAQ IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYYAYYGVG AWGQGTTVT 120
SS 122

SEQ ID NO: 429      moltype = AA length = 122
FEATURE          Location/Qualifiers
REGION           1..122
source            note = An artificially generated sequence
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 429
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQA PGKGLEWVAQ IKAKSNYYAT 60
YYAESVKGRF TISRDDSKNS LYLQMNSLKT EDTAVYYCRY VHYYAYYGVQ AWGQGTTVT 120
SS 122

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-continued

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SEQ ID NO: 430      moltype = AA length = 122
FEATURE
REGION
1..122
note = An artificially generated sequence
source
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 430
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQQA PGKGLEWVAQ IKAKSNYYAT 60
YVAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYYYYYGV D AWGQGTTVT 120
SS                                         122

SEQ ID NO: 431      moltype = AA length = 122
FEATURE
REGION
1..122
note = An artificially generated sequence
source
1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 431
QVQLVESGGG VVQPGRLRL SCAASGFTFS NAWMHWRQQA PGKGLEWVAQ IKAKSNYYKK 60
YVAESVKGRF TISRDDSKNS LYLMQNSLKT EDTAVYYCRY VHYYYYYGV D AWGQGTTVT 120
SS                                         122

SEQ ID NO: 432      moltype = AA length = 107
FEATURE
REGION
1..107
note = An artificially generated sequence
source
1..107
mol_type = protein
organism = synthetic construct

SEQUENCE: 432
ASVAAPSVFI FPPSDEQLKS GTASVVCLLN NFYPREAKVQ WKVDNALQSG NSQESVTEQD 60
SKDSTYSLSS TLTLSKADYE KHKVYACEVT HQGLSSPVTK SFNRGEC 107

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1.29. (canceled)

30. An antigen binding molecule comprising an immunoglobulin heavy chain variable region and an immunoglobulin light chain variable region, wherein the antigen binding molecule has T-cell receptor complex-binding activity, wherein the antigen binding molecule comprises a heavy chain variable region and a light chain variable region selected from the group consisting of:

- a) a heavy chain variable region comprising the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 142; and a light chain variable region comprising the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 223;
- b) a heavy chain variable region comprising the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 164; and a light chain variable region comprising the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 223; and,
- c) a heavy chain variable region comprising the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 168; and a light chain variable region comprising the CDR1, CDR2, and CDR3 regions comprised in SEQ ID NO: 223.

31. The antigen binding molecule of claim **30**, further comprising an Fc region having reduced Fc_y receptor binding activity.

32. The antigen binding molecule of claim **30**, wherein the antigen binding molecule is a bispecific antibody.

33. The bispecific antibody of claim **32**, which further comprises a variable domain having binding activity for an antigen expressed by a cancer cell.

34. The bispecific antibody of claim **32**, wherein the bispecific antibody comprises an Fc region comprised of a first heavy chain constant region comprising the amino acid sequence of SEQ ID NO: 60 or 62 and a second heavy chain constant region comprising the amino acid sequence of SEQ ID NO: 61.

35. A nucleic acid encoding the antigen binding molecule of claim **30**.

36. A vector comprising the nucleic acid of claim **35**.

37. A cell comprising the nucleic acid of claim **35**.

38. A method for producing an antigen binding molecule, wherein the method comprises a step of culturing the cell of claim **37**.

39. An antigen binding molecule produced by the method of claim **38**.

40. A pharmaceutical composition comprising the antigen binding molecule of claim **30** and a pharmaceutically acceptable carrier.

41. The antigen binding molecule of claim **30**, wherein the heavy chain variable region and light chain variable region include:

- a) a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 142; and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 223;
- b) a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 164; and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 223; or,

c) a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 168; and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 223.

42. The antigen binding molecule of claim **41**, wherein the antigen binding molecule is a bispecific antibody.

43. The bispecific antibody of claim **42**, which further comprises a variable domain having binding activity for an antigen expressed by a cancer cell.

44. The bispecific antibody of claim **42**, wherein the bispecific antibody comprises an Fc region comprised of a first heavy chain constant region comprising the amino acid sequence of SEQ ID NO: 60 or 62 and a second heavy chain constant region comprising the amino acid sequence of SEQ ID NO: 61.

45. A nucleic acid encoding the antigen binding molecule of claim **41**.

46. A vector comprising the nucleic acid of claim **45**.

47. A cell comprising the nucleic acid of claim **45**.

48. A method for producing an antigen binding molecule, wherein the method comprises a step of culturing the cell of claim **47**.

49. An antigen binding molecule produced by the method of claim **48**.

50. A pharmaceutical composition comprising the antigen binding molecule of claim **41** and a pharmaceutically acceptable carrier.

51. A method of treating cancer that comprises administering the bispecific antibody of claim **32** to a patient in need thereof.

* * * * *