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(12) **United States Patent**
Koide et al.(10) **Patent No.:** US 12,391,770 B2
(45) **Date of Patent:** *Aug. 19, 2025(54) **COMPOSITIONS AND METHODS
COMPRISING ANTIBODIES THAT BIND TO
COVALENT PEPTIDE CONJUGATES**(71) Applicant: **New York University**, New York, NY (US)(72) Inventors: **Shohei Koide**, New York, NY (US); **Benjamin Neel**, New York, NY (US); **Carmine Fedele**, New York, NY (US); **Kai Wen Teng**, Flushing, NY (US); **Akiko Koide**, New York, NY (US); **Takamitsu Hattori**, West New York, NJ (US); **Lorenzo Maso**, New York, NY (US)(73) Assignee: **NEW YORK UNIVERSITY**, New York, NY (US)

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(21) Appl. No.: **18/585,676**(22) Filed: **Feb. 23, 2024**(65) **Prior Publication Data**

US 2024/0327542 A1 Oct. 3, 2024

Related U.S. Application Data

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(60) Provisional application No. 63/253,499, filed on Oct. 7, 2021, provisional application No. 63/154,627, filed on Feb. 26, 2021.

(51) **Int. Cl.**
C07K 16/44 (2006.01)
A61K 47/68 (2017.01)
C07K 16/28 (2006.01)(52) **U.S. Cl.**
CPC **C07K 16/44** (2013.01); **A61K 47/6849** (2017.08); **C07K 16/2833** (2013.01); **C07K 2317/56** (2013.01); **C07K 2317/626** (2013.01)(58) **Field of Classification Search**

None

See application file for complete search history.

(56) **References Cited**

U.S. PATENT DOCUMENTS

8,945,571 B2	2/2015	Mössner et al.
9,447,159 B2	9/2016	Ast et al.
10,344,091 B2 *	7/2019	Koide
11,242,405 B2	2/2022	Scheinberg et al.
2005/0059113 A1	3/2005	Bedian et al.
2019/0256604 A1	8/2019	Koide et al.
2020/0129555 A1	4/2020	Hanada et al.
2020/0399377 A1	12/2020	Weidanz
2022/0227883 A1	7/2022	Holt et al.
2022/0289866 A1	9/2022	Craik et al.
2022/0324998 A1	10/2022	O'Bryan et al.

FOREIGN PATENT DOCUMENTS

EP	3286222 A1	2/2018
EP	2723380 B1	8/2019
EP	4178976 A1	5/2023
EP	4182029 A1	5/2023
WO	01/98324 A1	12/2001
WO	2015160928 A2	10/2015
WO	2020154617 A1	7/2020
WO	2022072760 A1	4/2022
WO	2022241017 A2	11/2022

OTHER PUBLICATIONS

Lloyd et al., Protein Engineering, Design & Selection 22:159-168 (Year: 2009).*
Edwards et al., J Mol Biol. 334(1): 103-118 (Year: 2003).*
Khan and Salunke., J. Immunol 192: 5398-5405 (Year: 2014).*
Poosarla et al., Biotechn. Bioeng. 114(6): 1331-1342 (Year: 2017).*
Rudikoff et al., Proc Natl Acad Sci USA 79: 1979-1983 (Year: 1982).*
Piatesi et al., ChemBio Chem 5: 460-466 (Year: 2004).*
Wu et al., J. Mol. Biol. 294: 151-162 (Year: 1999).*
Lambert, J.M., et al., Antibody-Drug Conjugates (ADCs) for Personalized Treatment of Solid Tumors: A Review, Advances in Therapy, Mar. 30, 2017, vol. 34, pp. 1015-1035.
Teng, K.W., et al., Selective and noncovalent targeting of RAS mutants for inhibition and degradation, Nature Communications, May 11, 2021, vol. 12, No. 2656, 13 pages.

(Continued)

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(57) **ABSTRACT**

Provided are compositions and methods that include binding partners that bind with specificity to target sites on proteins or peptides that comprise a covalently attached molecule. The binding partners are provided as antibodies and antibody derivatives that specifically bind to proteins and peptides that have been covalently modified by attachment of a molecule, such as a drug. The binding partners can bind with specificity to covalently modified peptides when presented in the context of a major histocompatibility complex (MHC). Uses of the compositions and methods for prophylaxis or therapy of disorders are also provided.

8 Claims, 30 Drawing Sheets

Specification includes a Sequence Listing.

(56)

References Cited

OTHER PUBLICATIONS

Visscher, M., et al., Covalent targeting of acquired cysteines in cancer, Current Opinion in Chemical Biology, Nov. 28, 2015, vol. 30, pp. 61-67.

Maserati et al., "Abstract PR04: Anti peptide-HLA (TCR-like) antibodies specific for the KRAS G12V neoantigen," Molecular Cancer Research: American Association for Cancer Research, May 1, 2020.

* cited by examiner

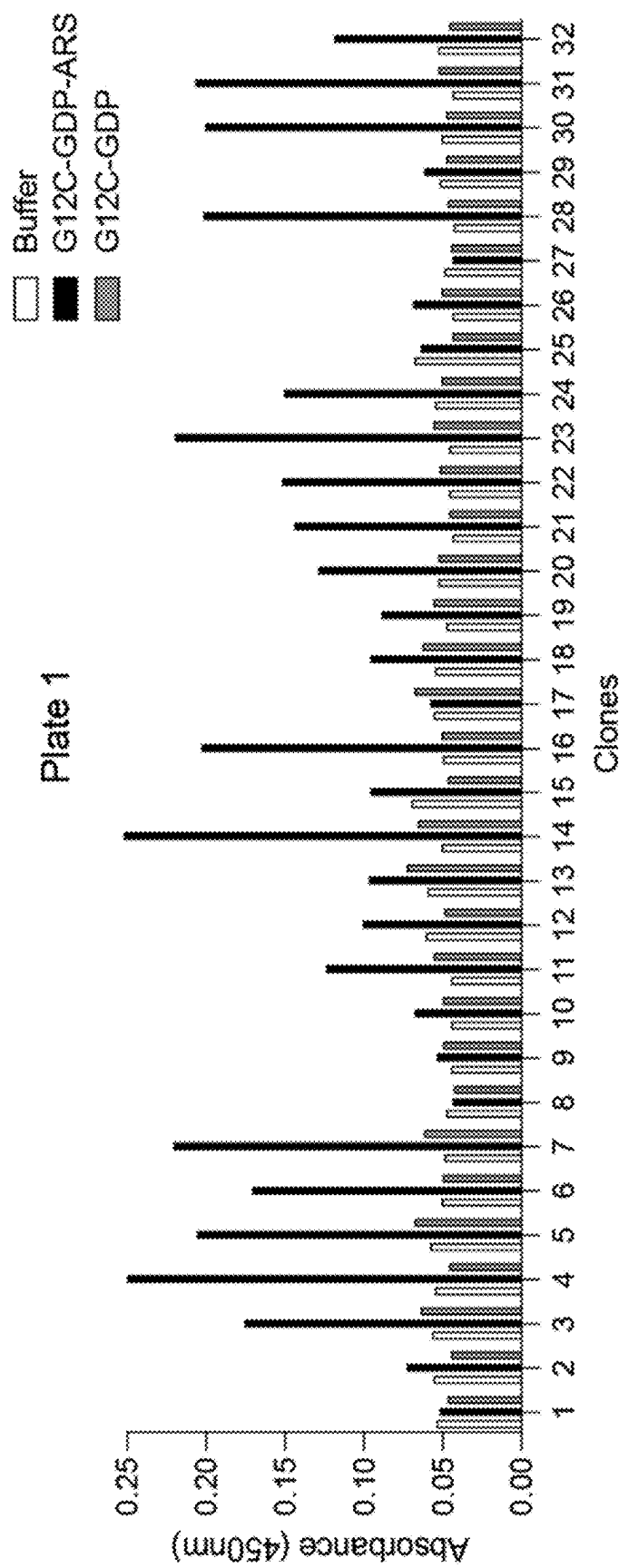


Fig. 1

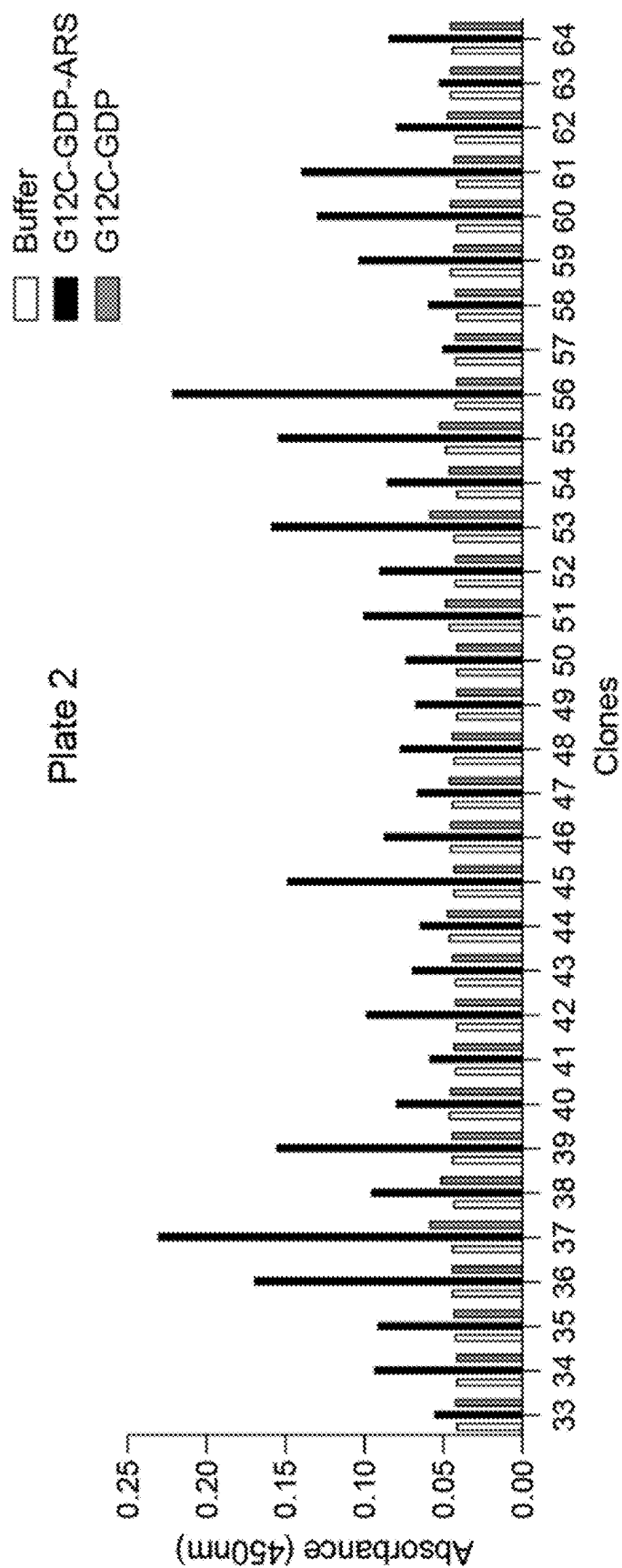


Fig. 1 (continued)

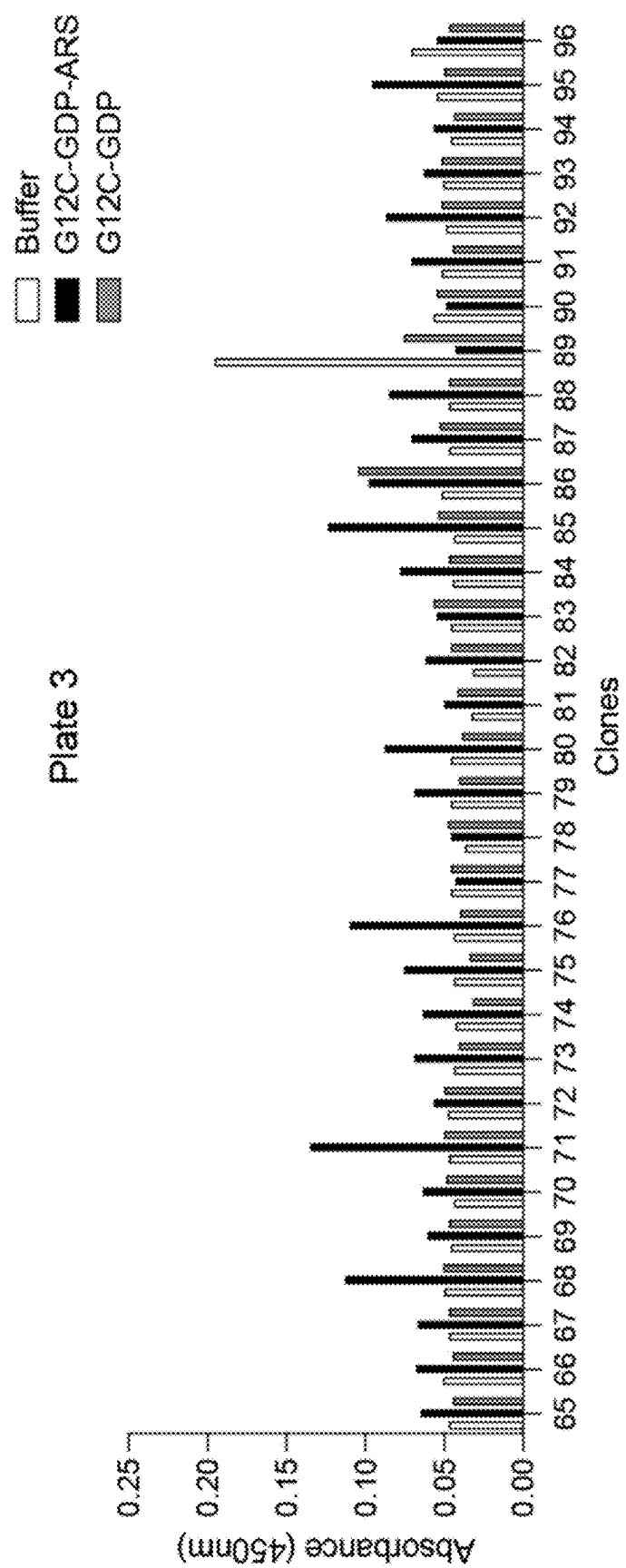


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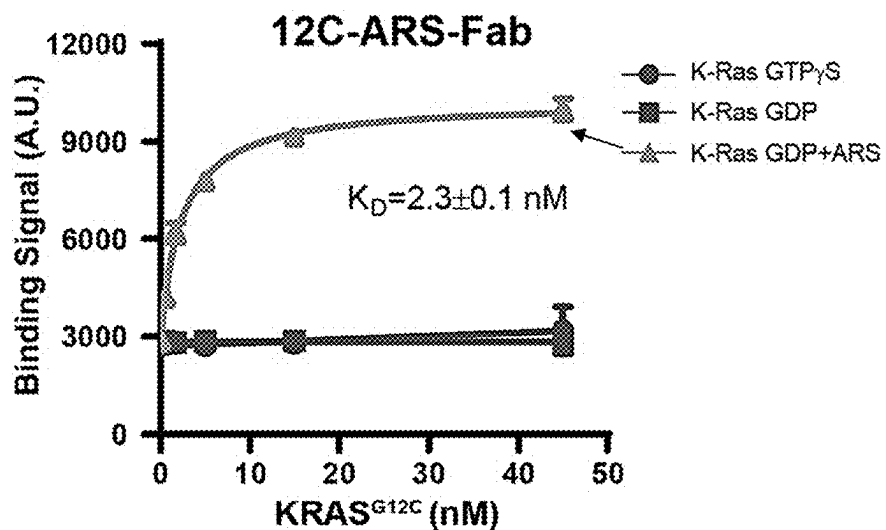


Fig. 2

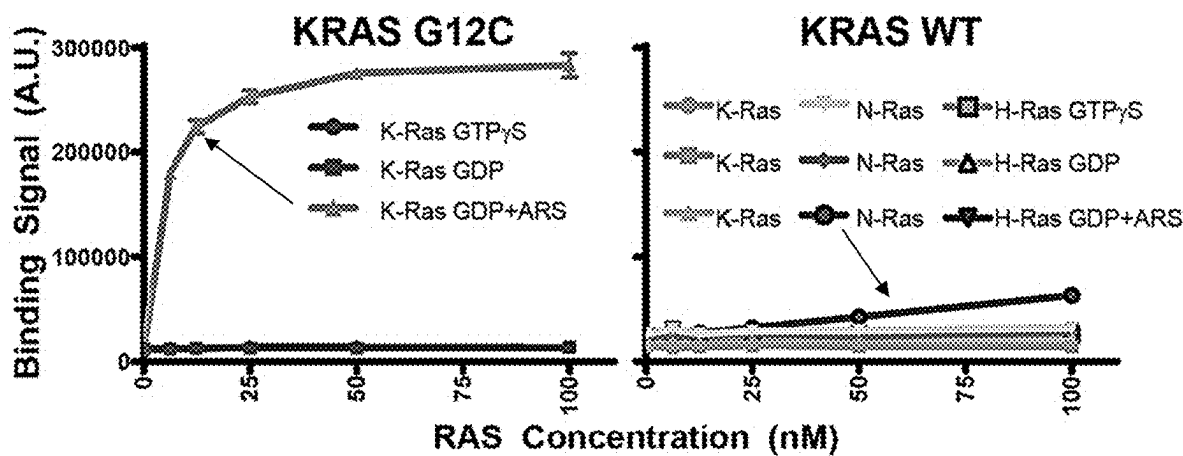


Fig. 3

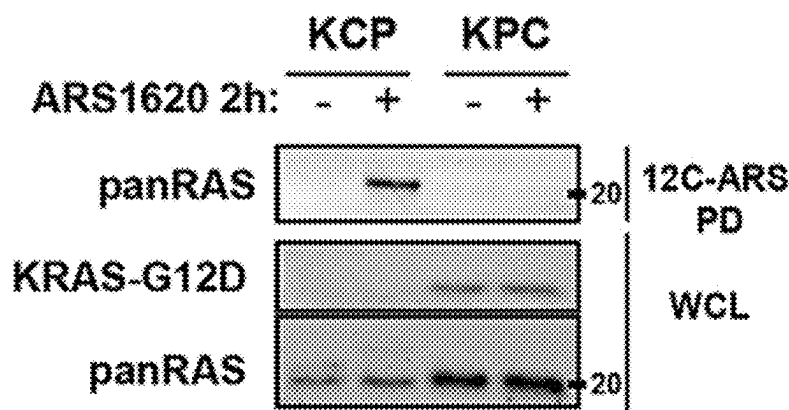
A**RAS-less MEF****B**

Fig. 4

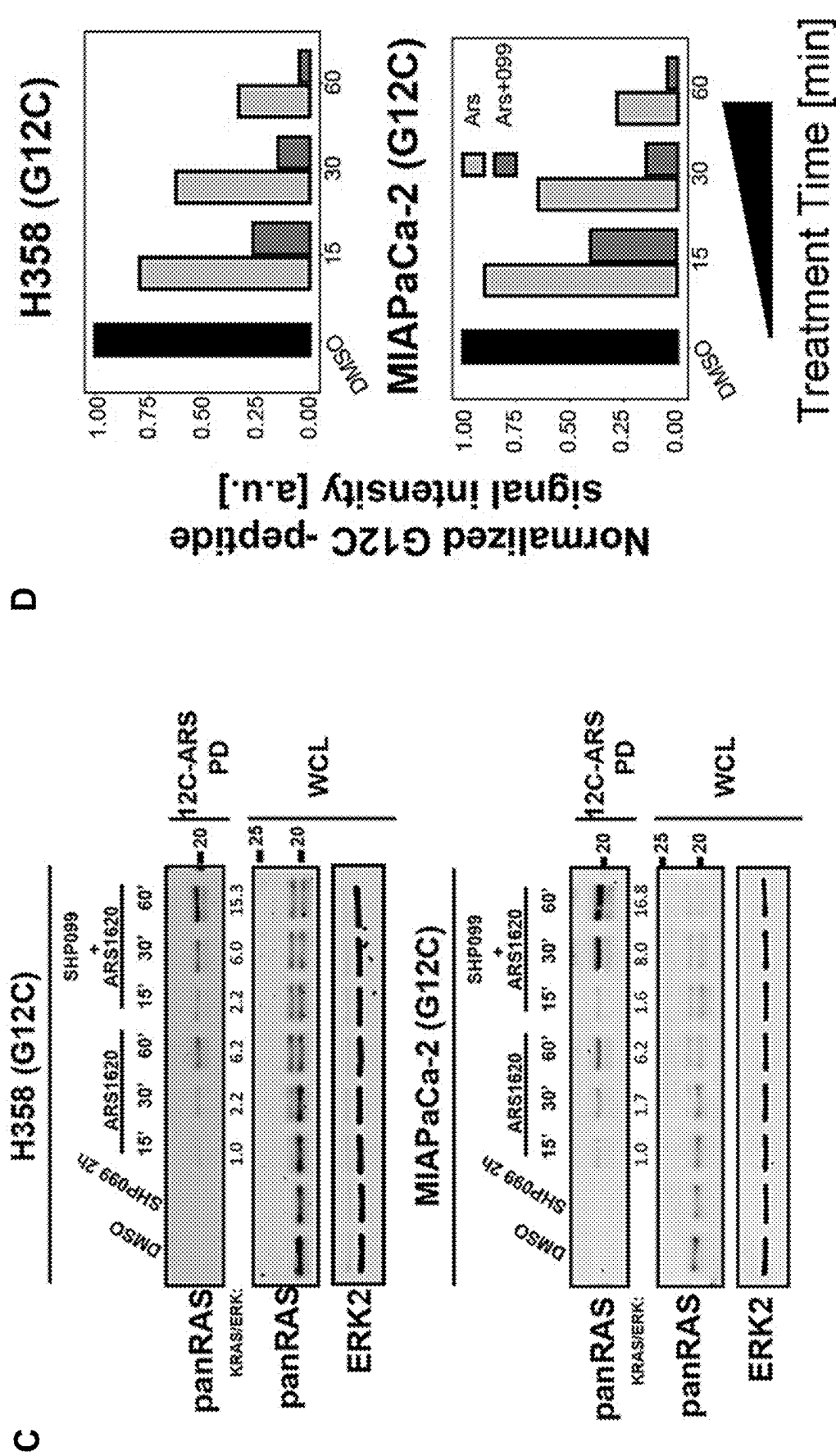


Fig. 4 (continued)

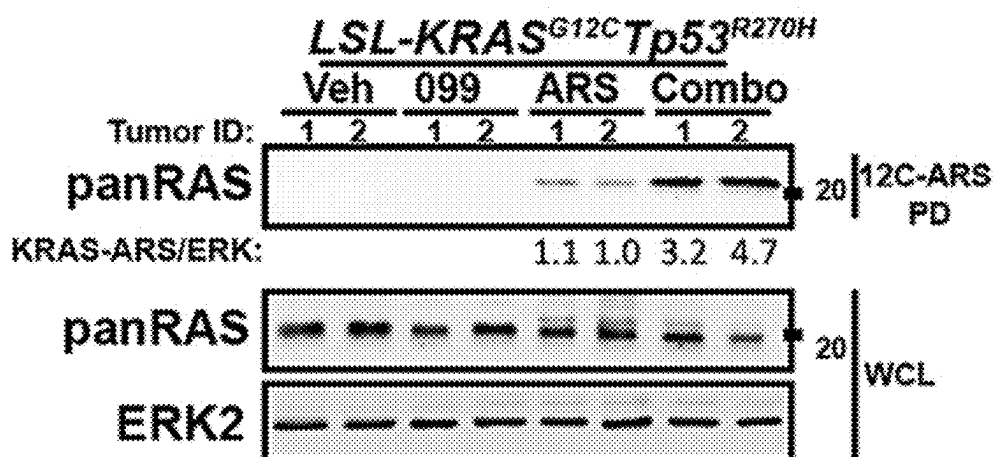
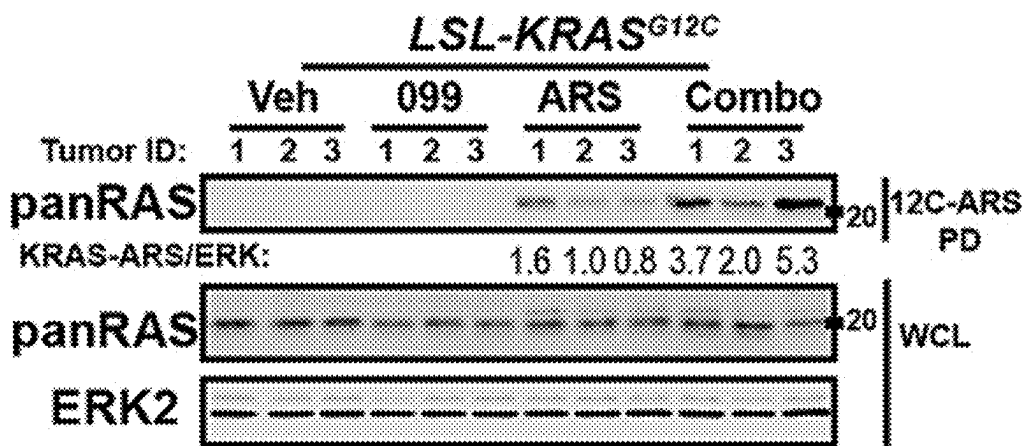
A**B**

Fig. 5

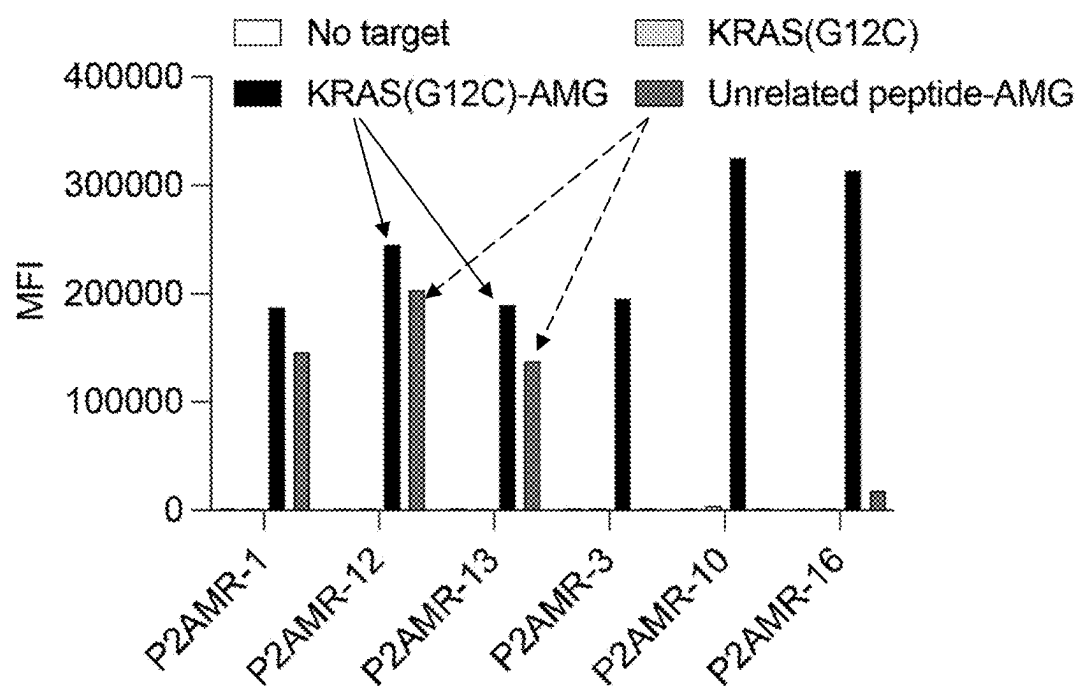


Fig. 6

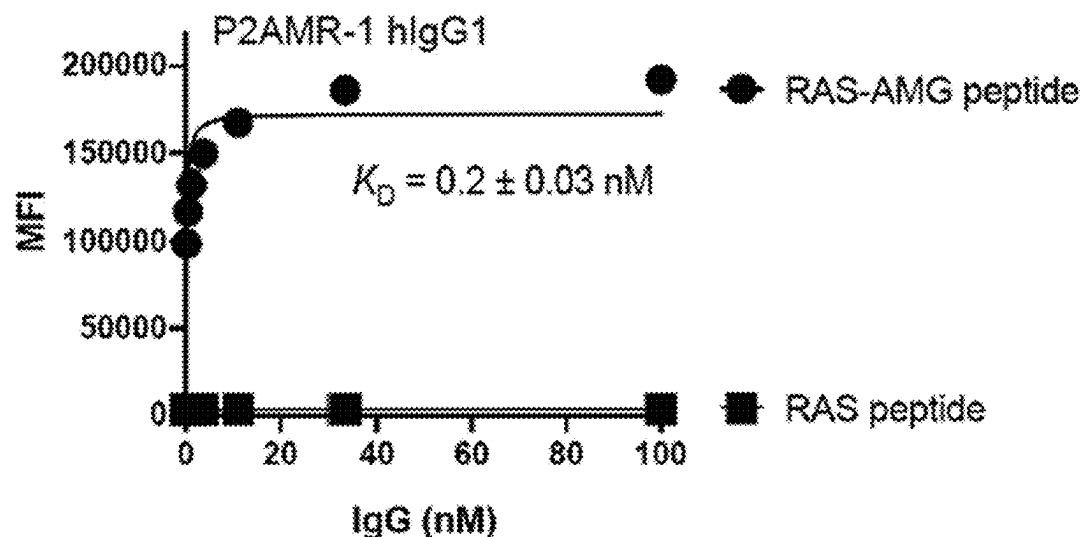


Fig. 7

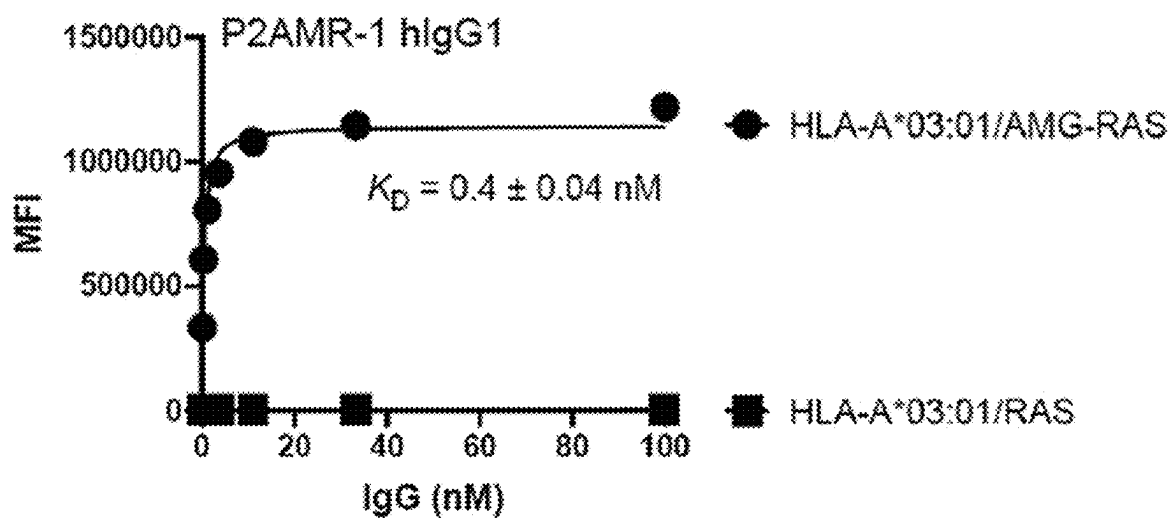


Fig. 8

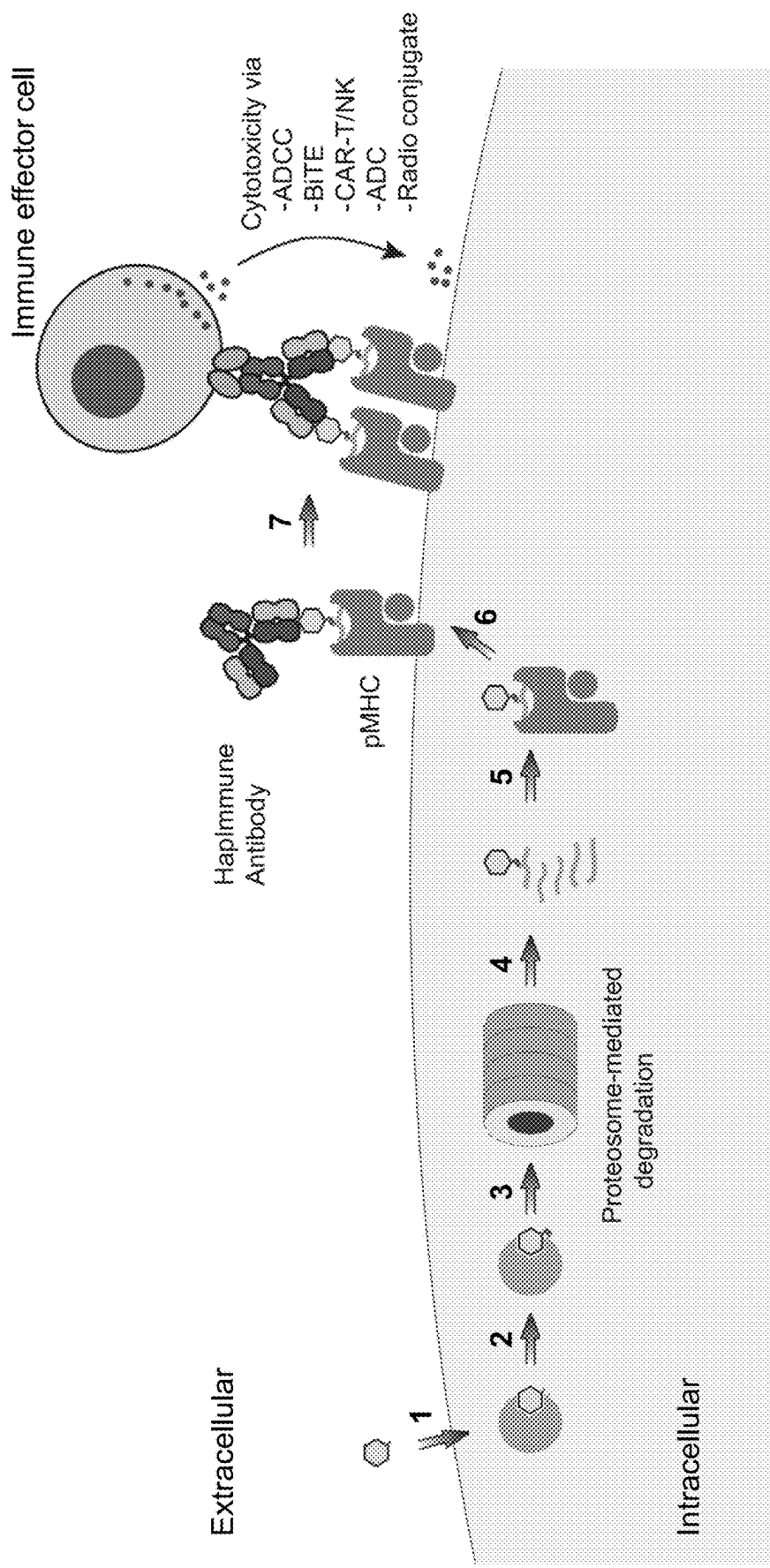


Fig. 9

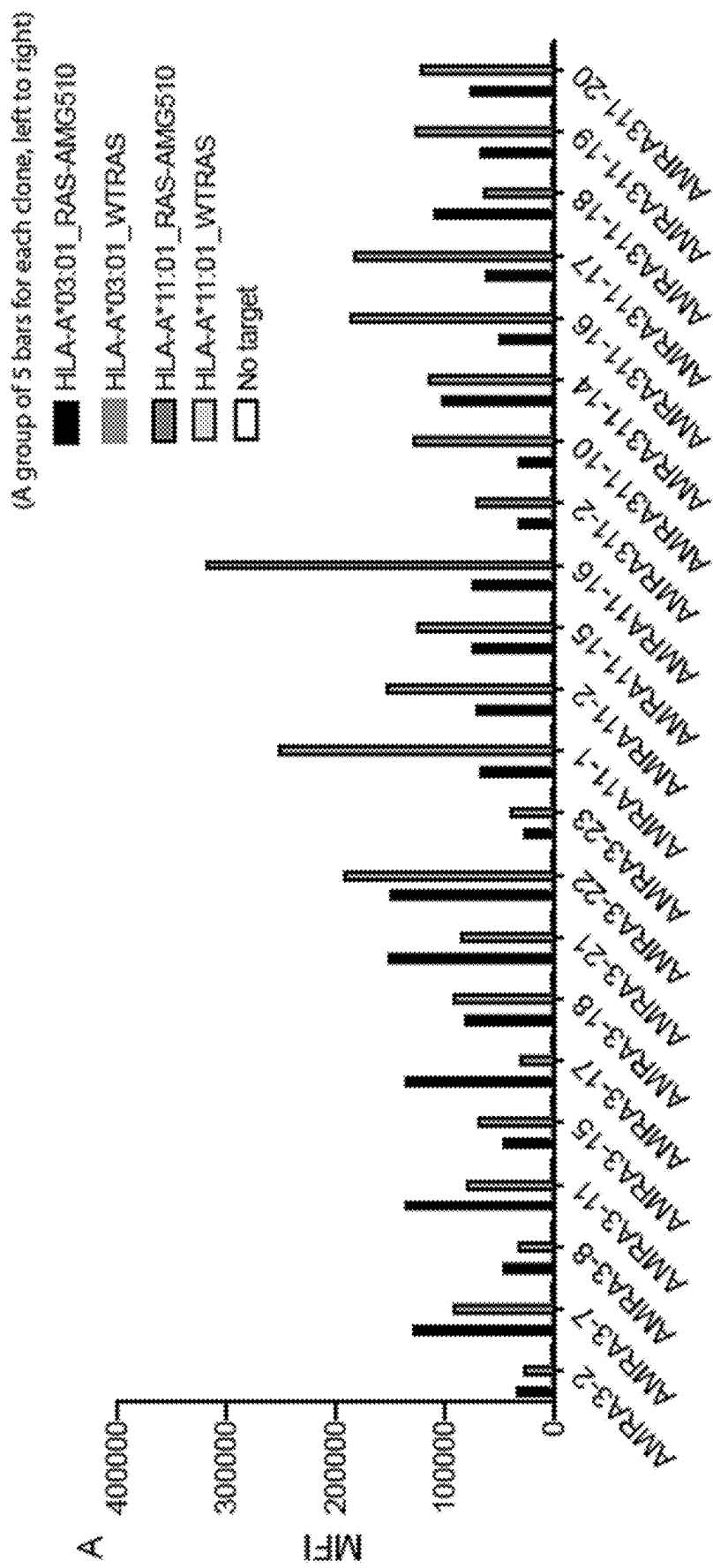


Fig. 10

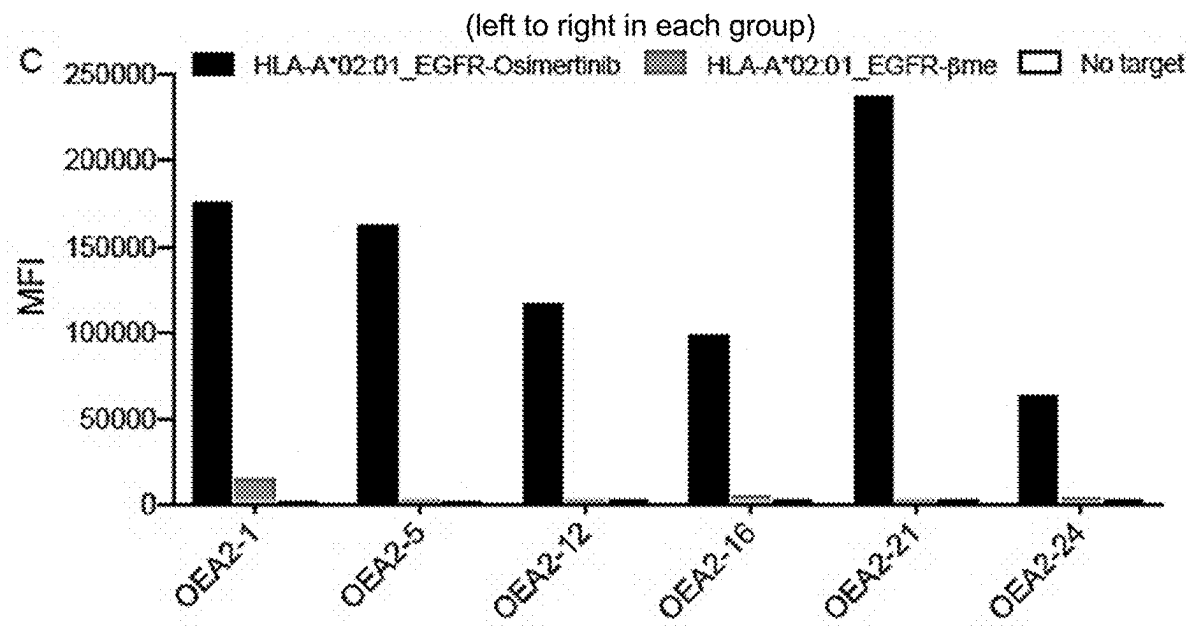
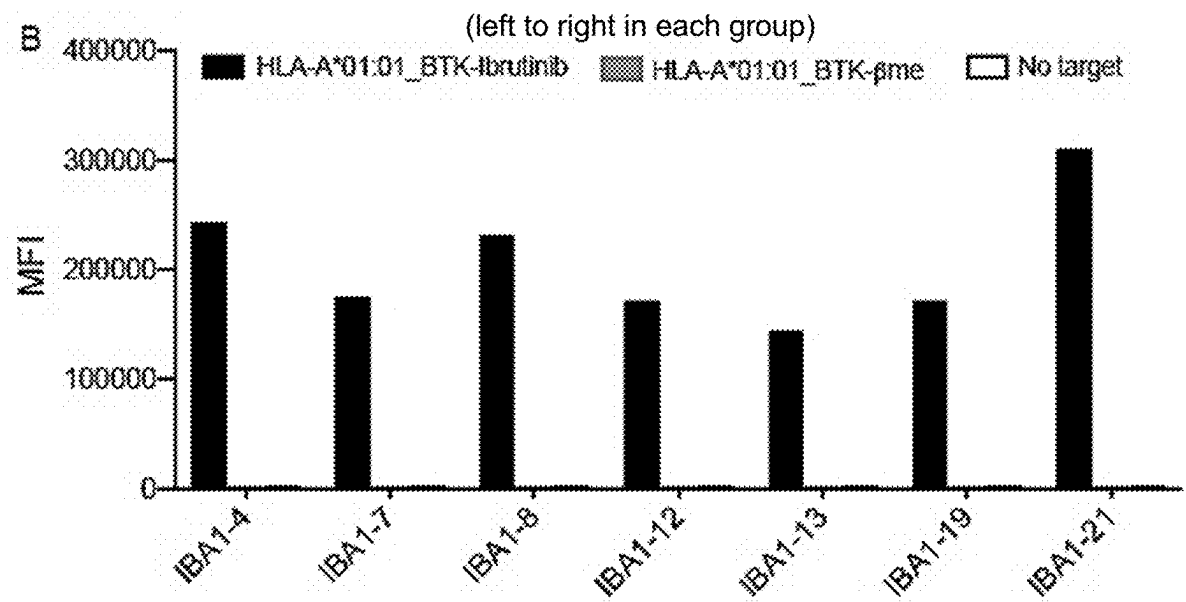


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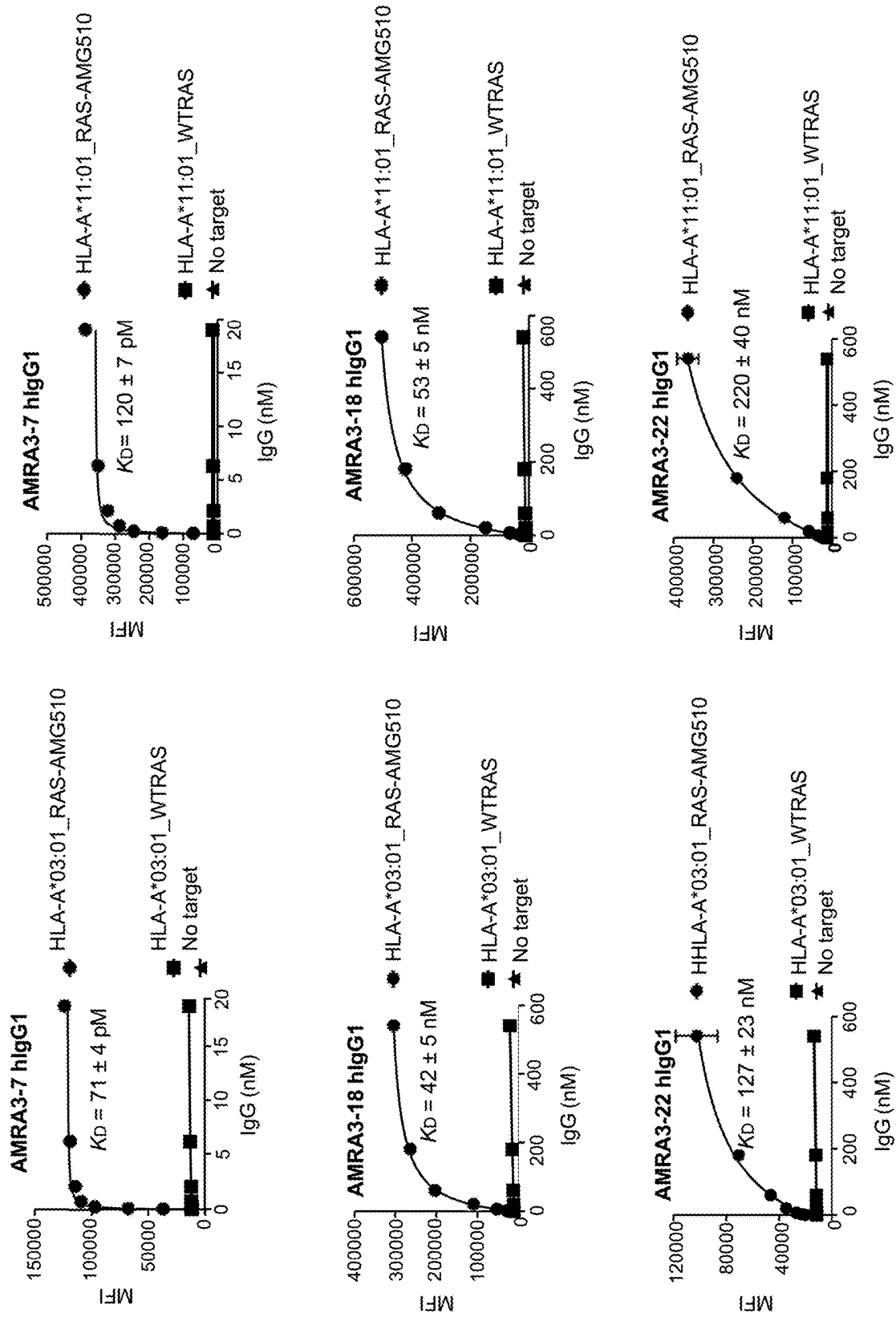


Fig. 11

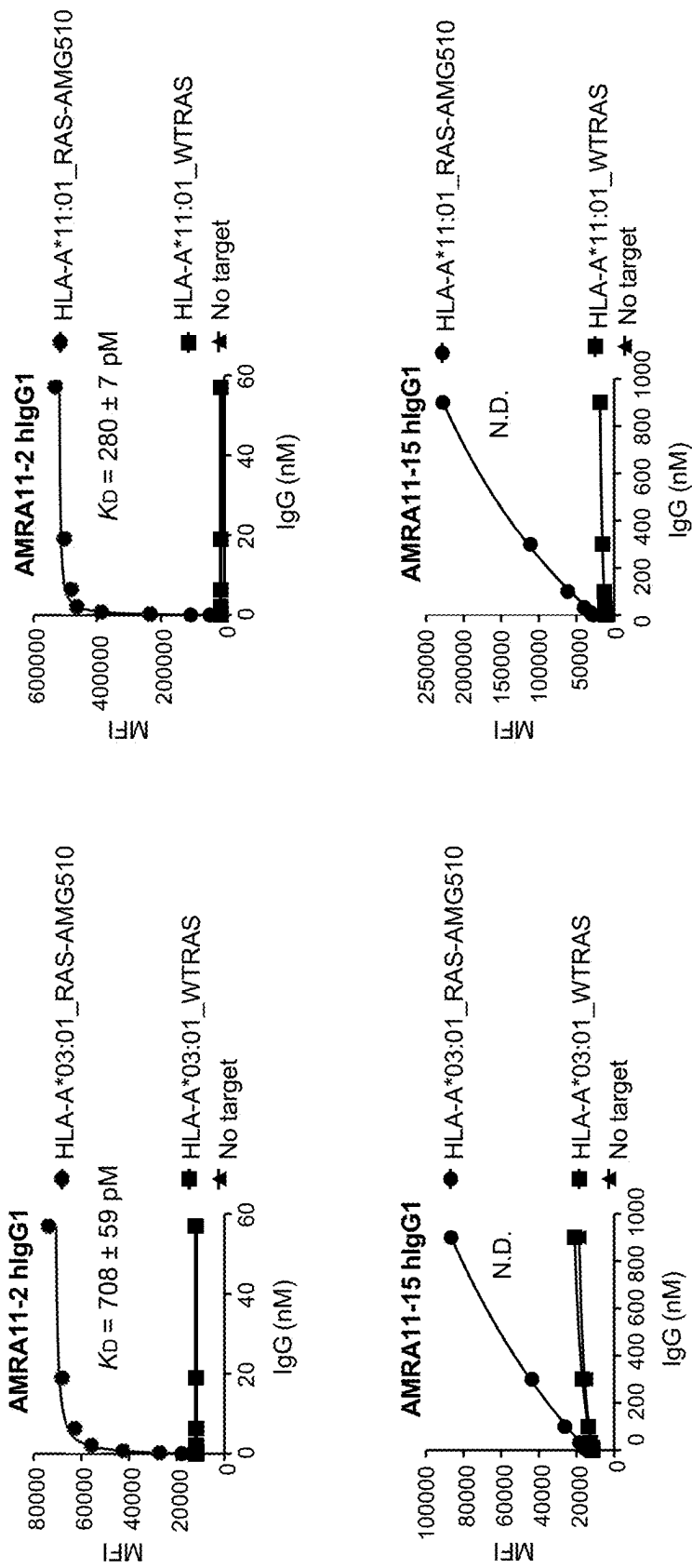


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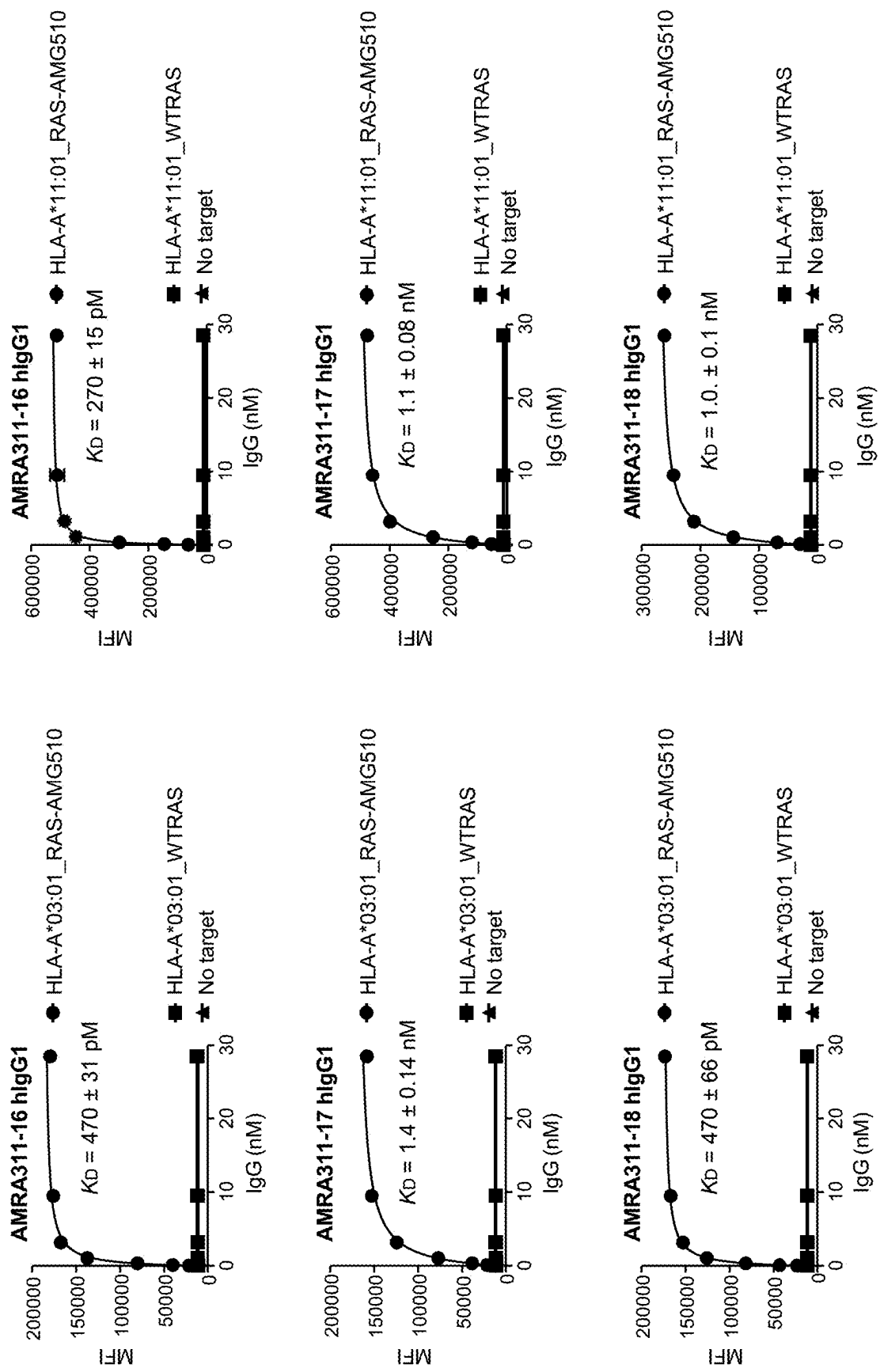


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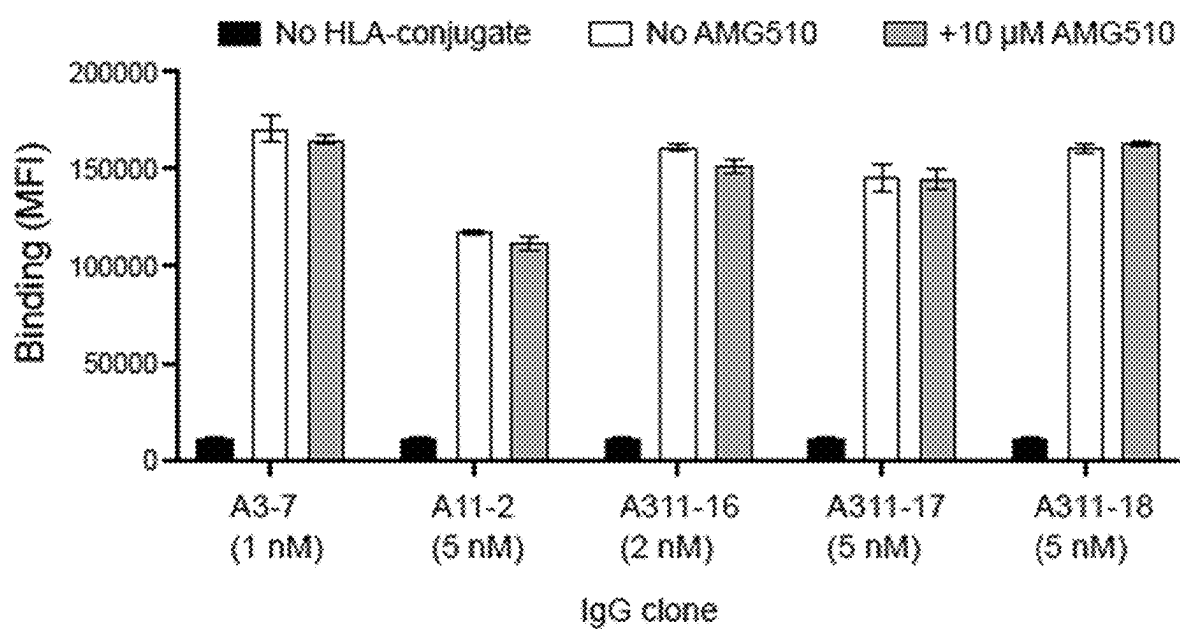


Fig. 12

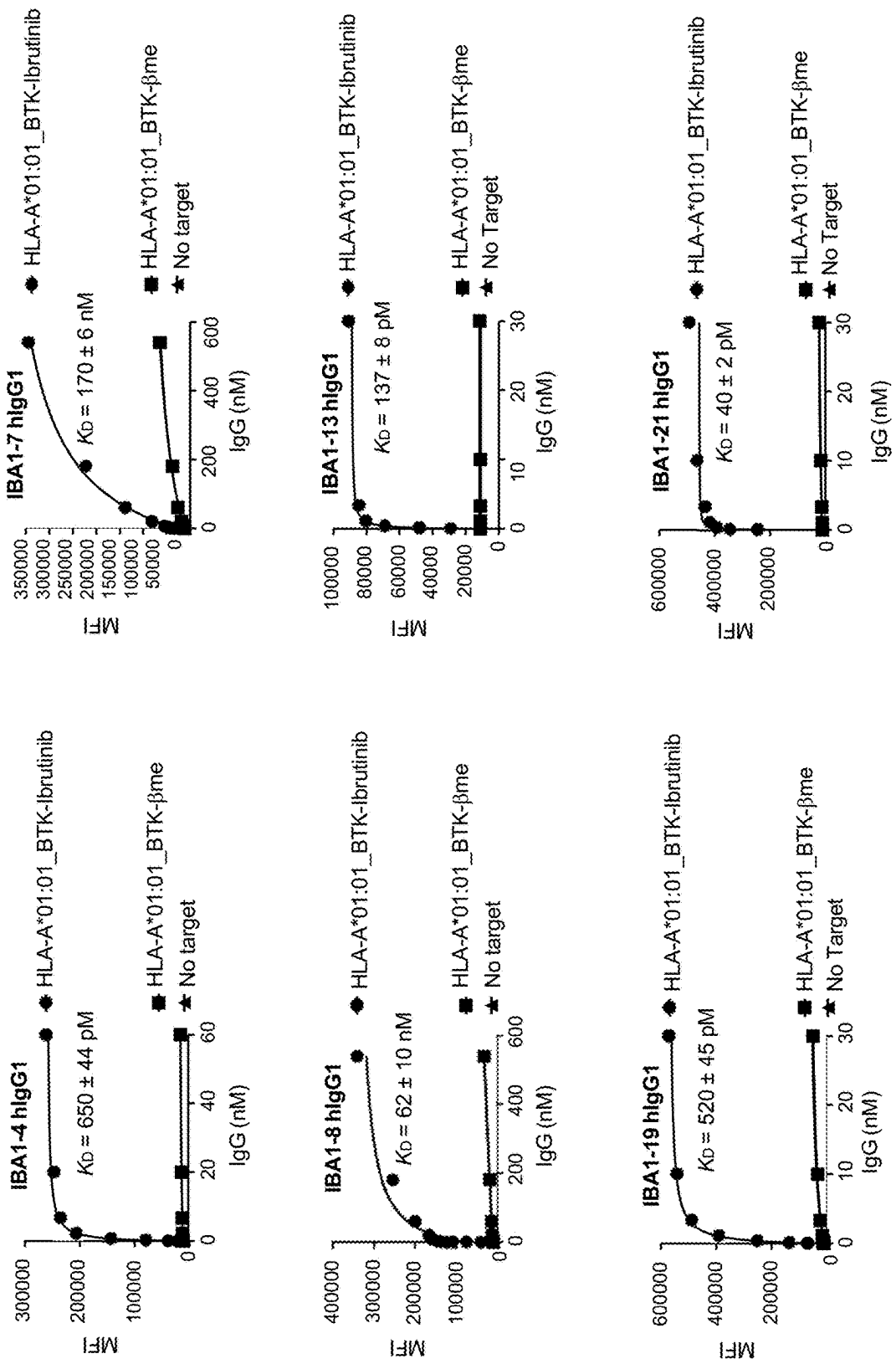


Fig. 13

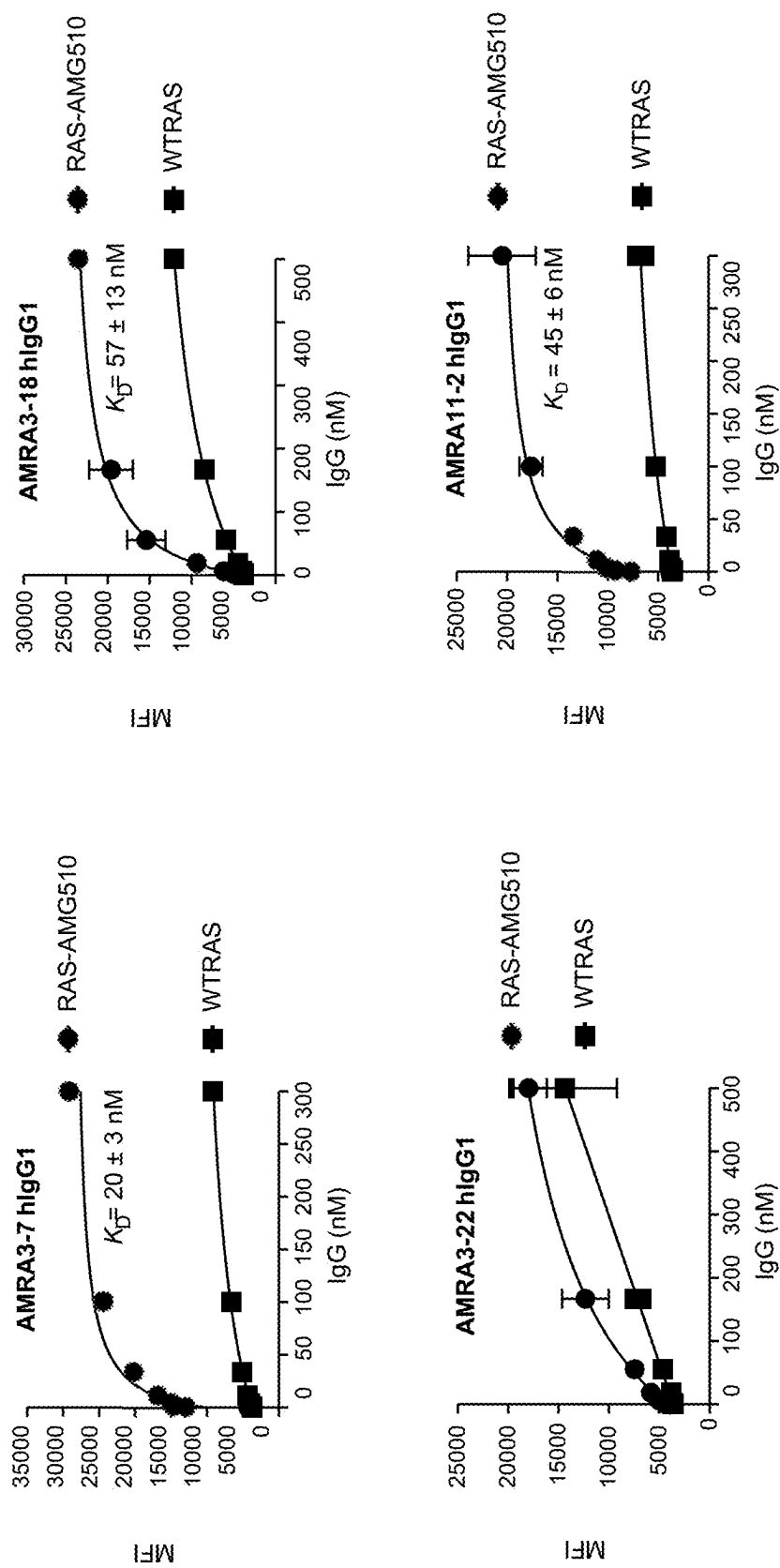


Fig. 14

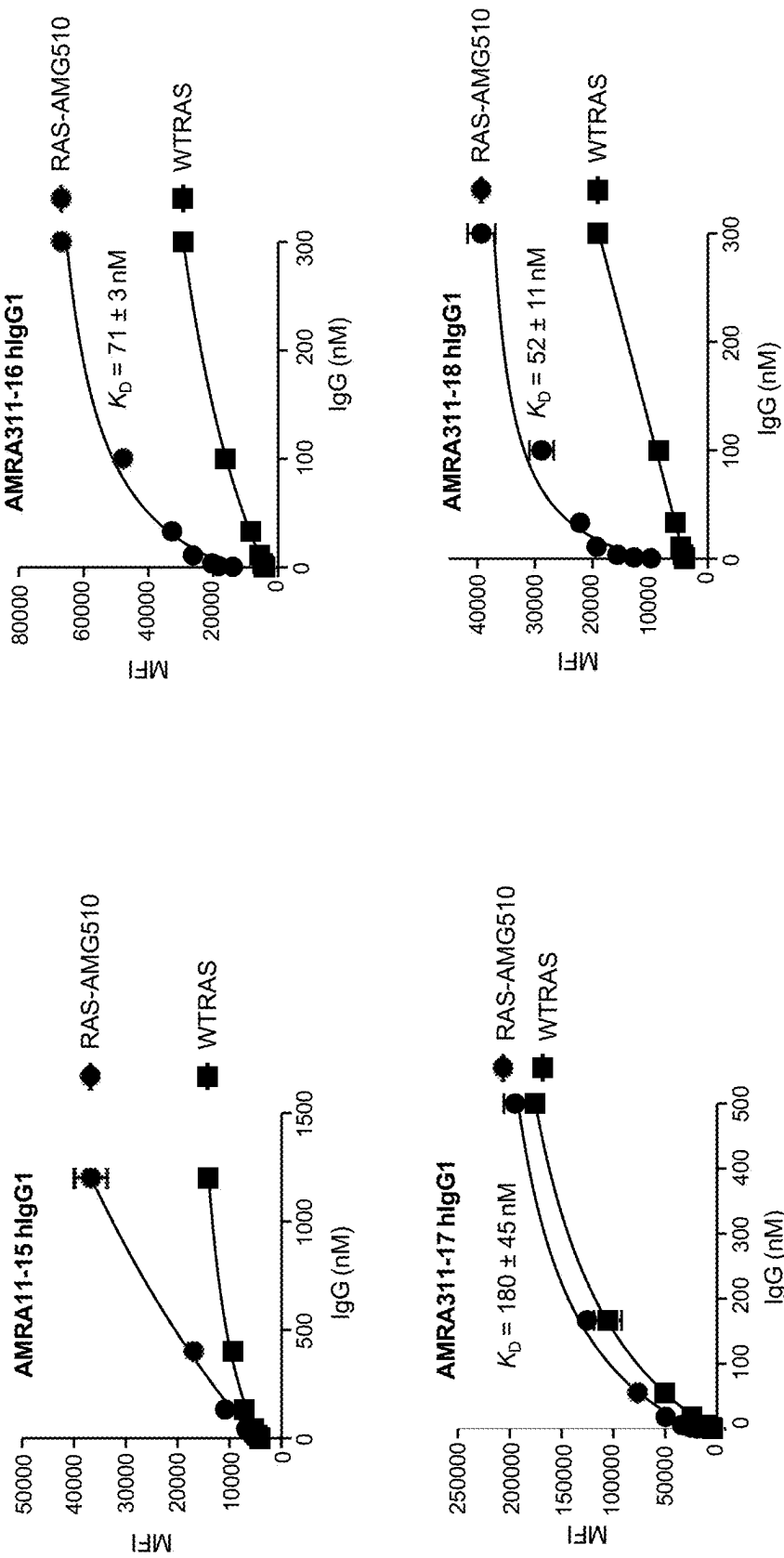


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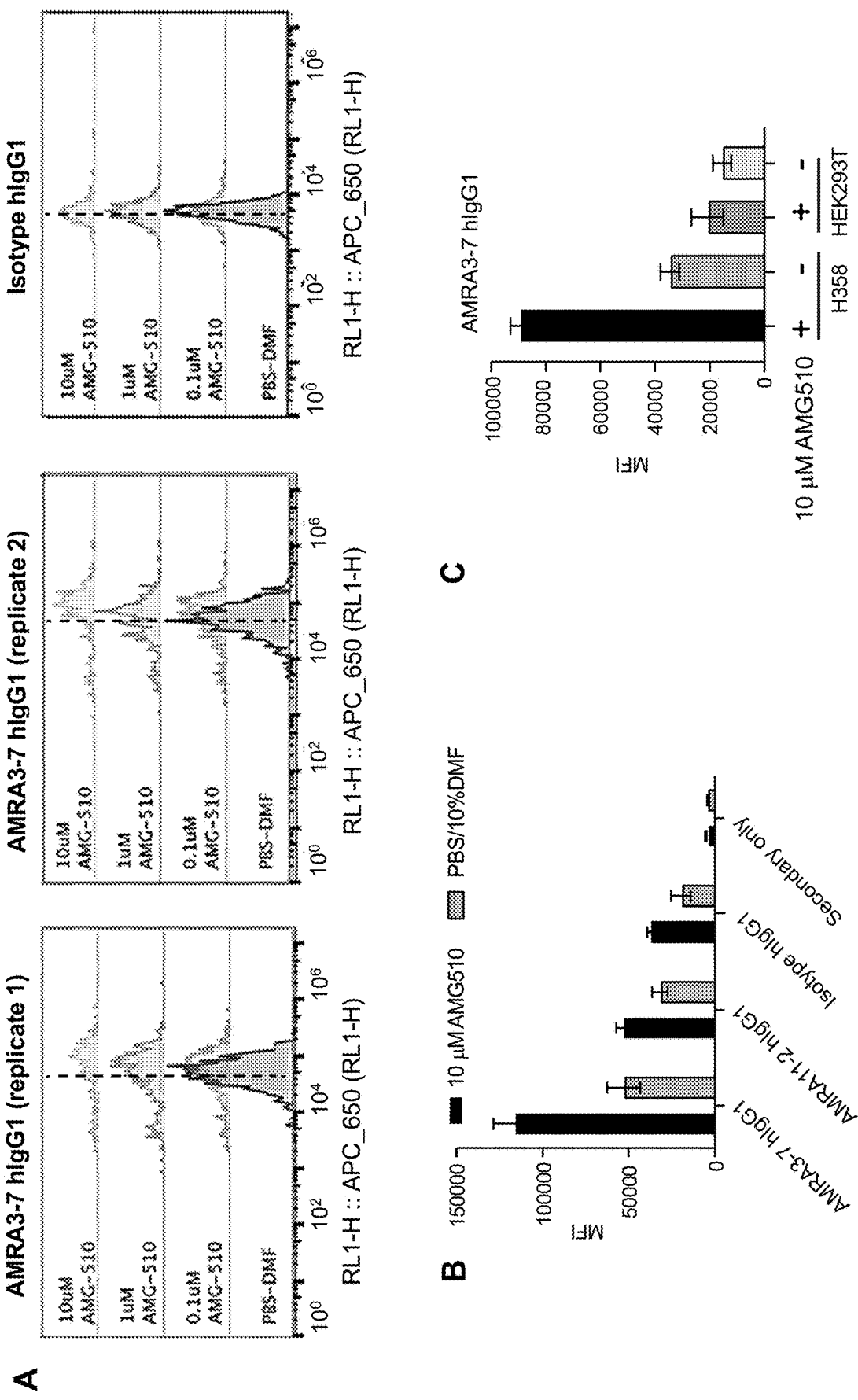


Fig. 15

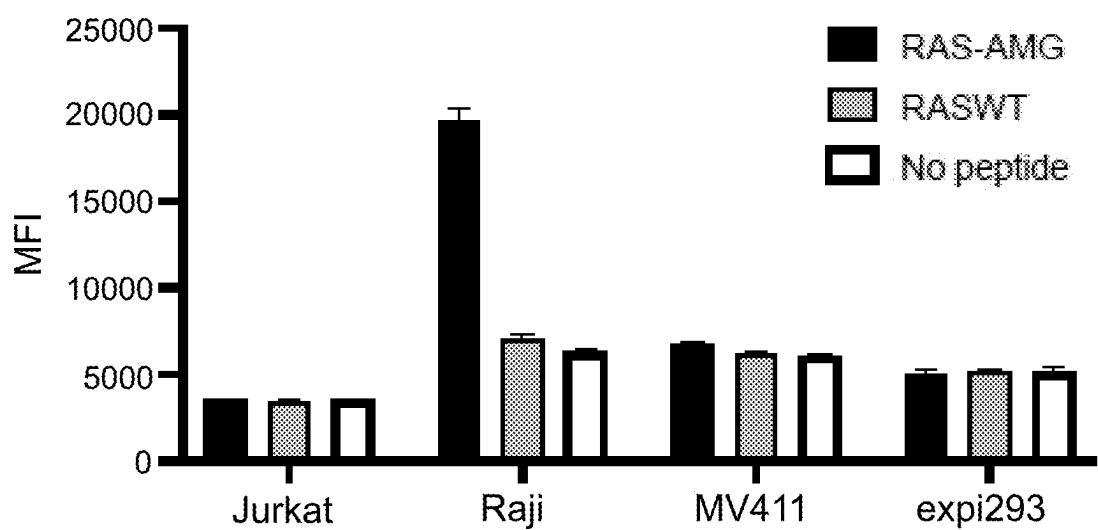
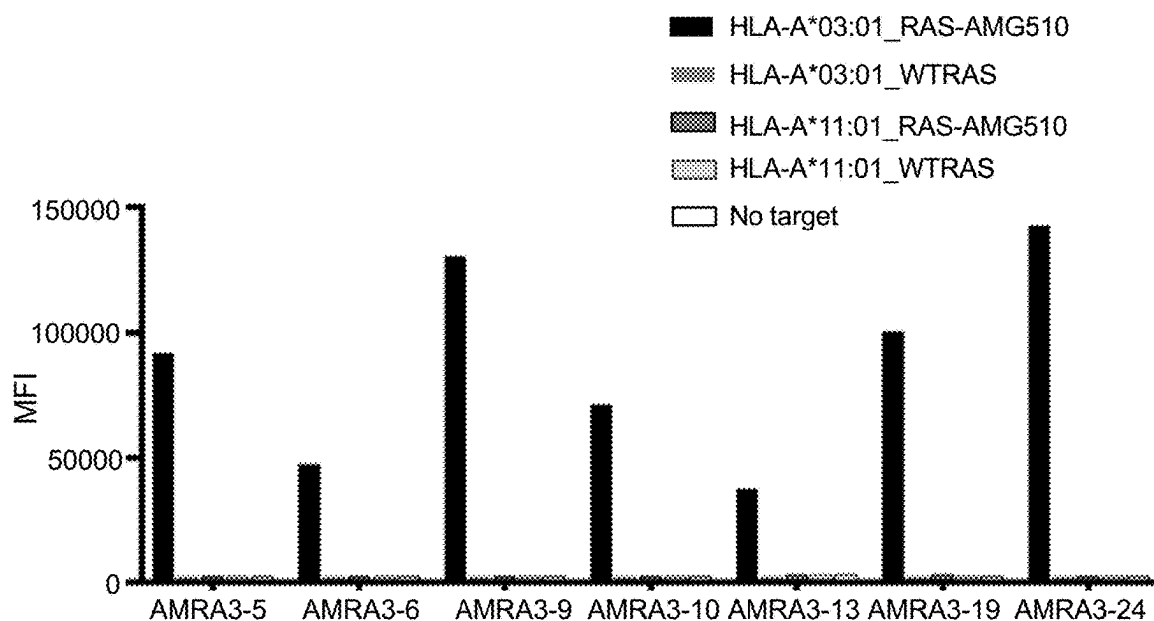


Fig. 16

A

(A group of 5 bars for each clone, left to right)

**B**

(A group of 5 bars for each clone, left to right)

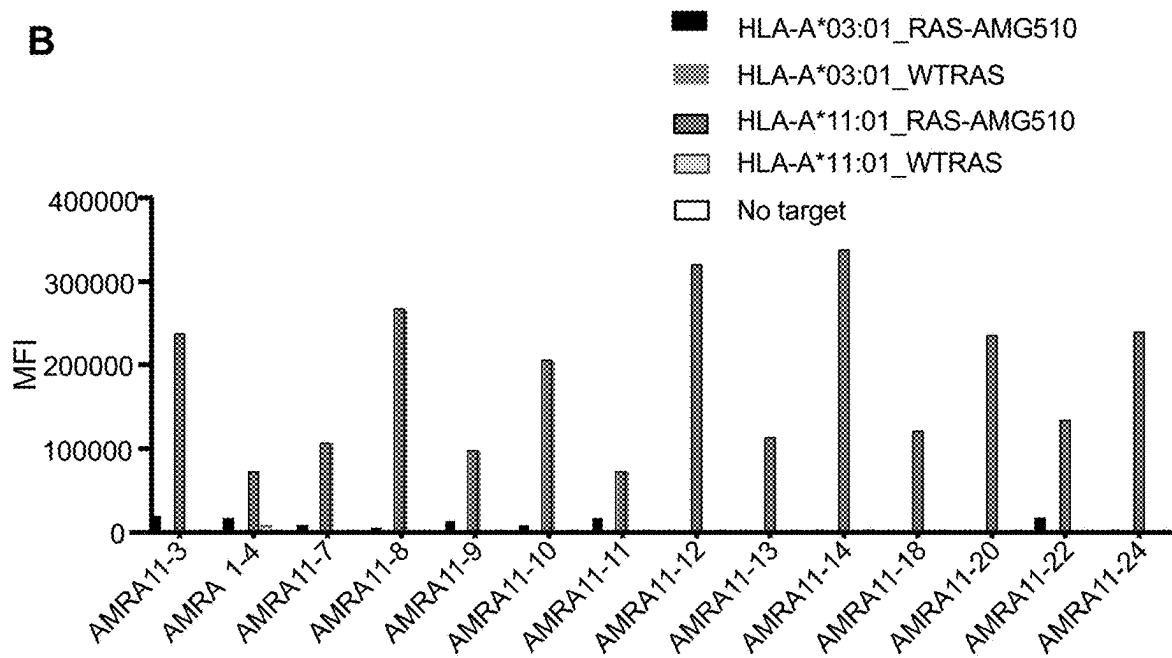


Fig. 17

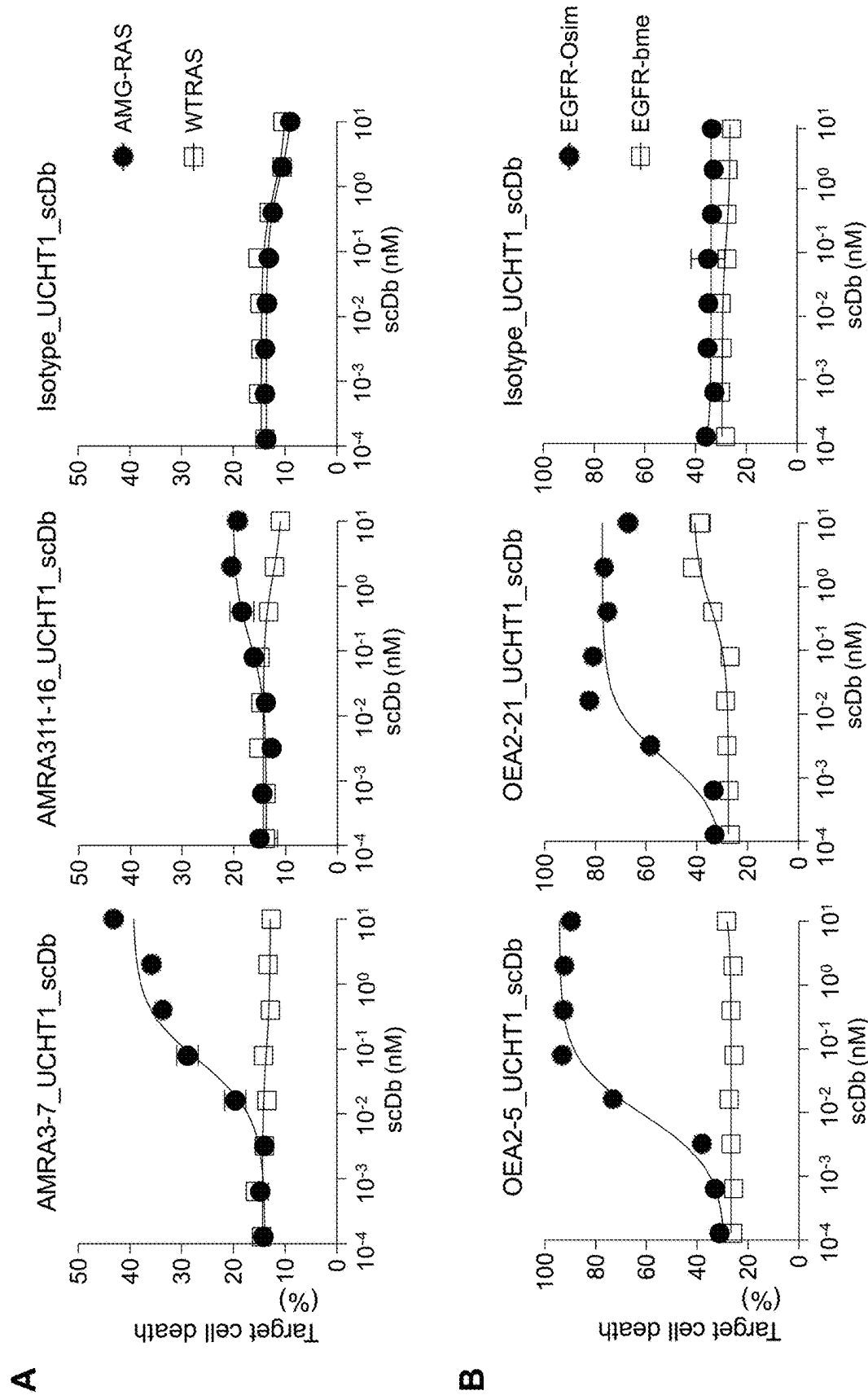


Fig. 18

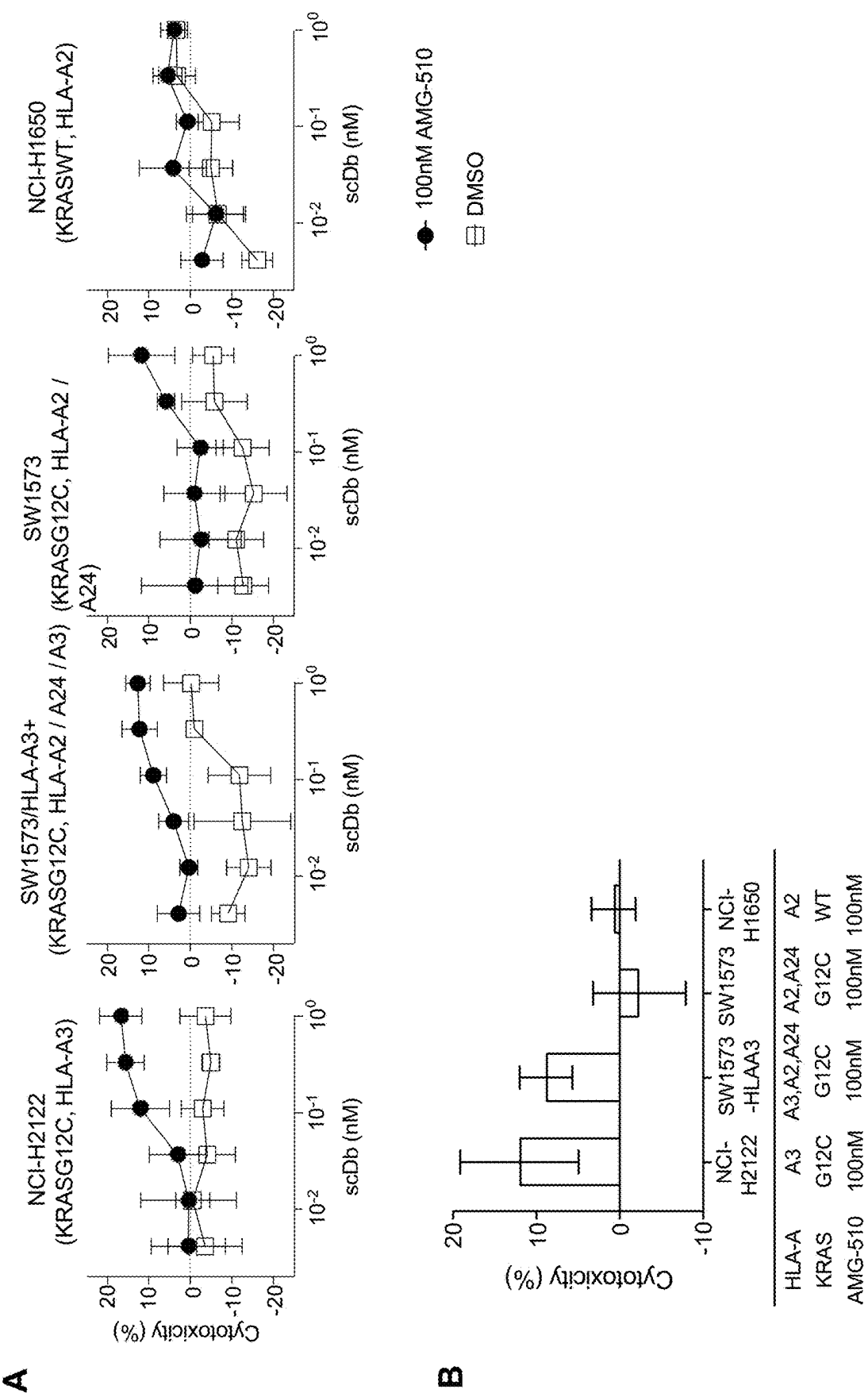


Fig. 19

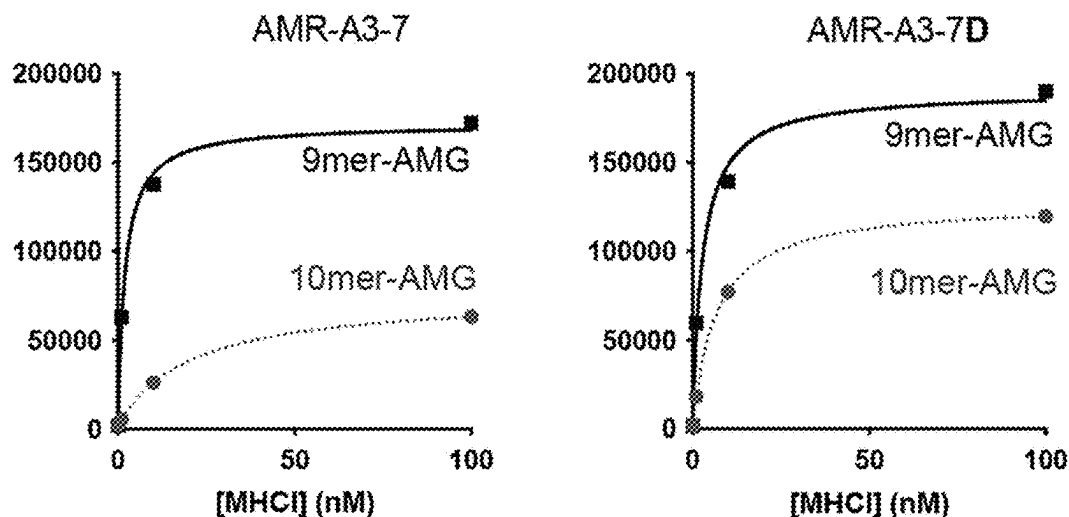


Fig. 20

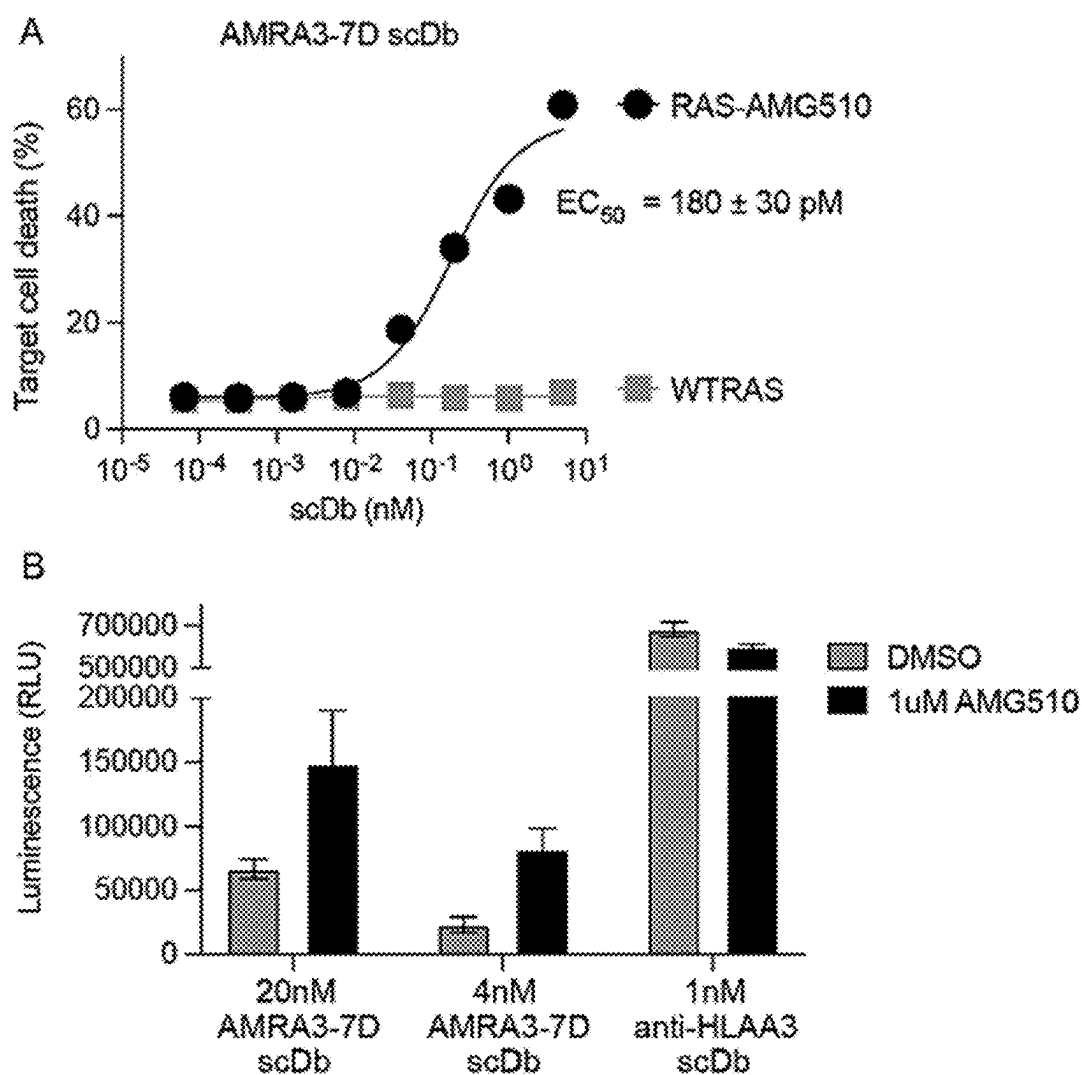


Fig. 21

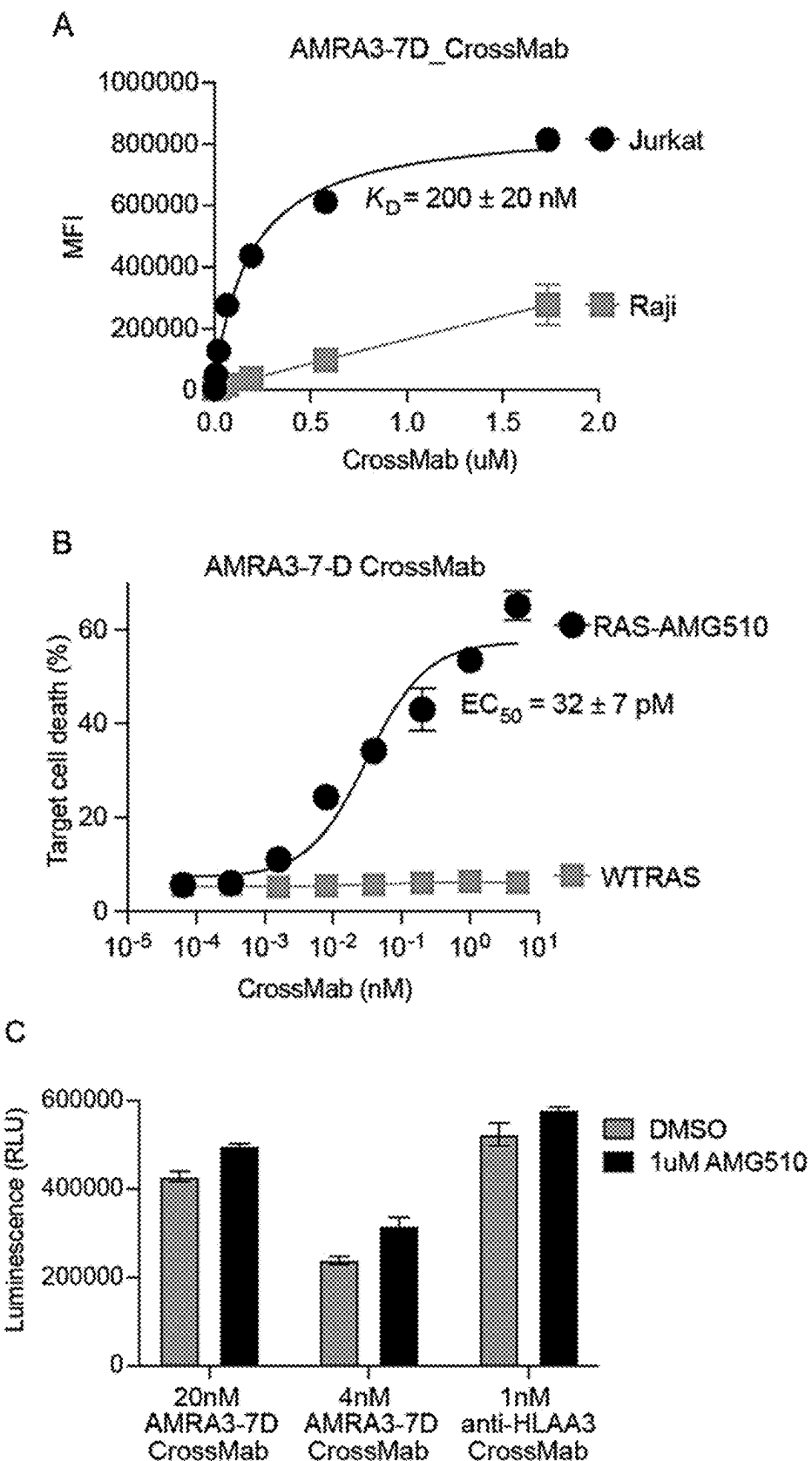


Fig. 22

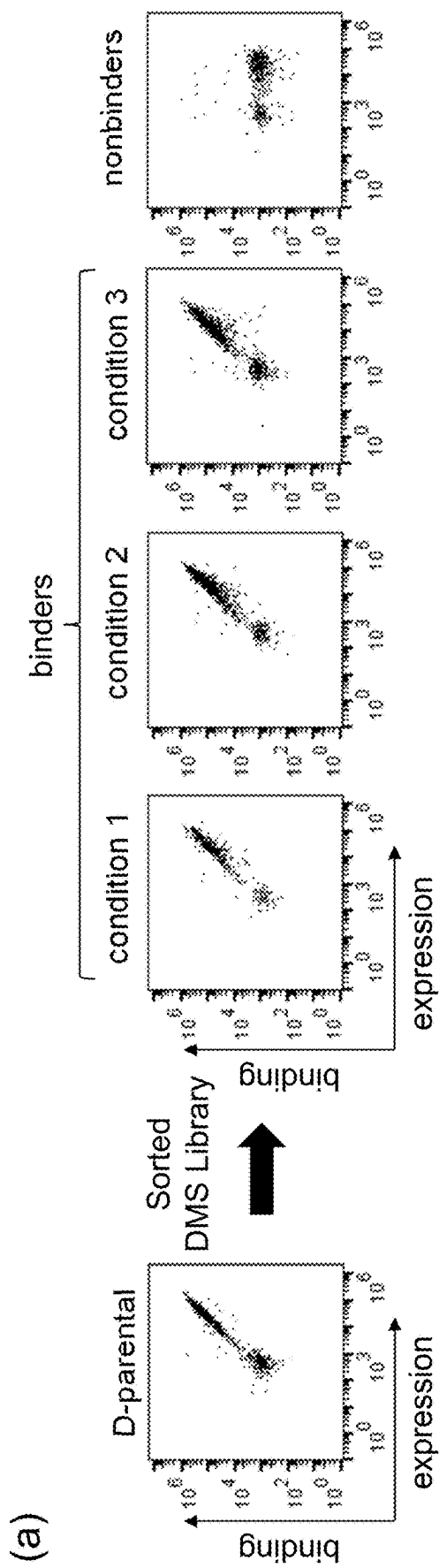


Fig. 23

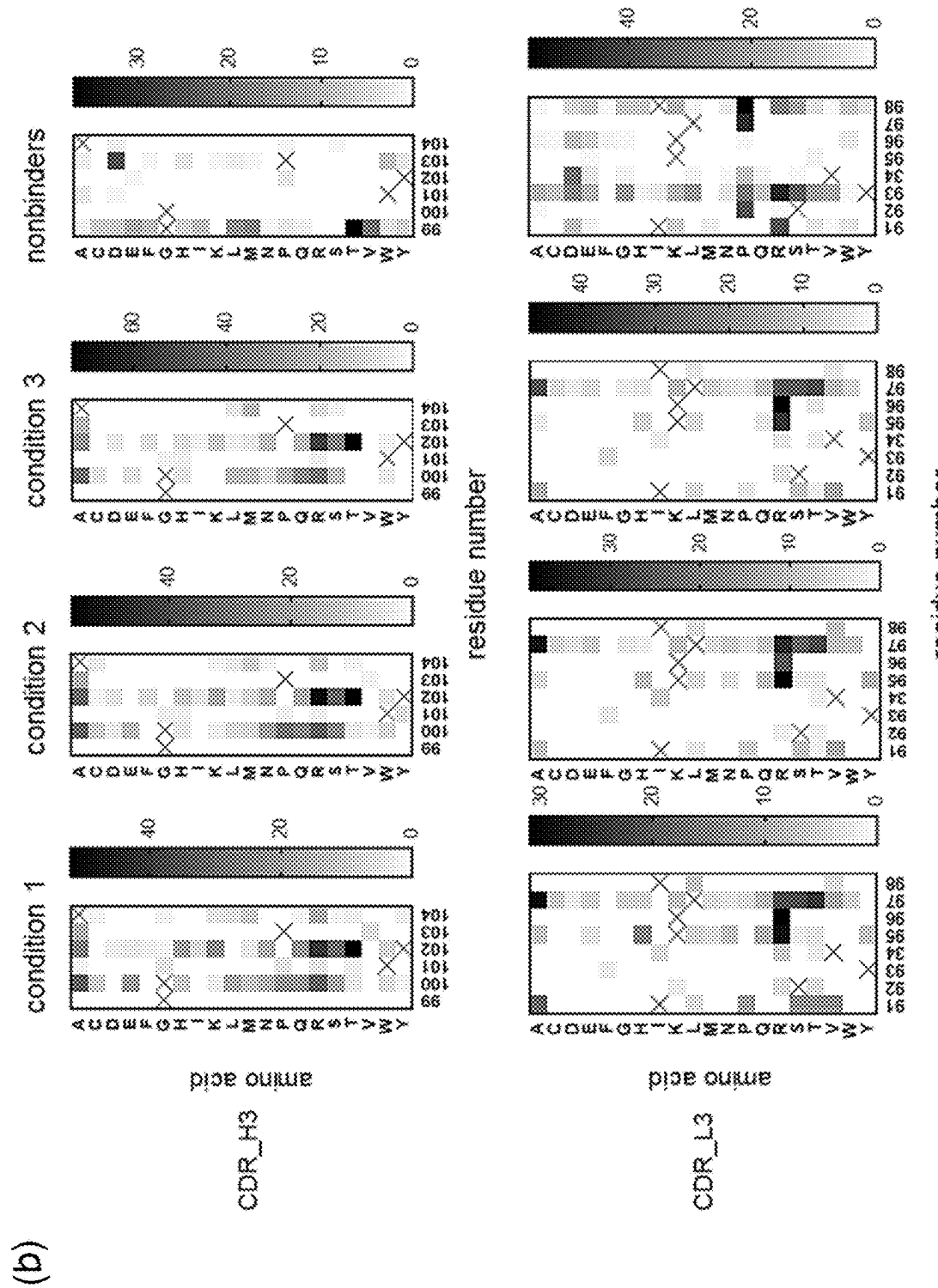


Fig. 23 (continued)

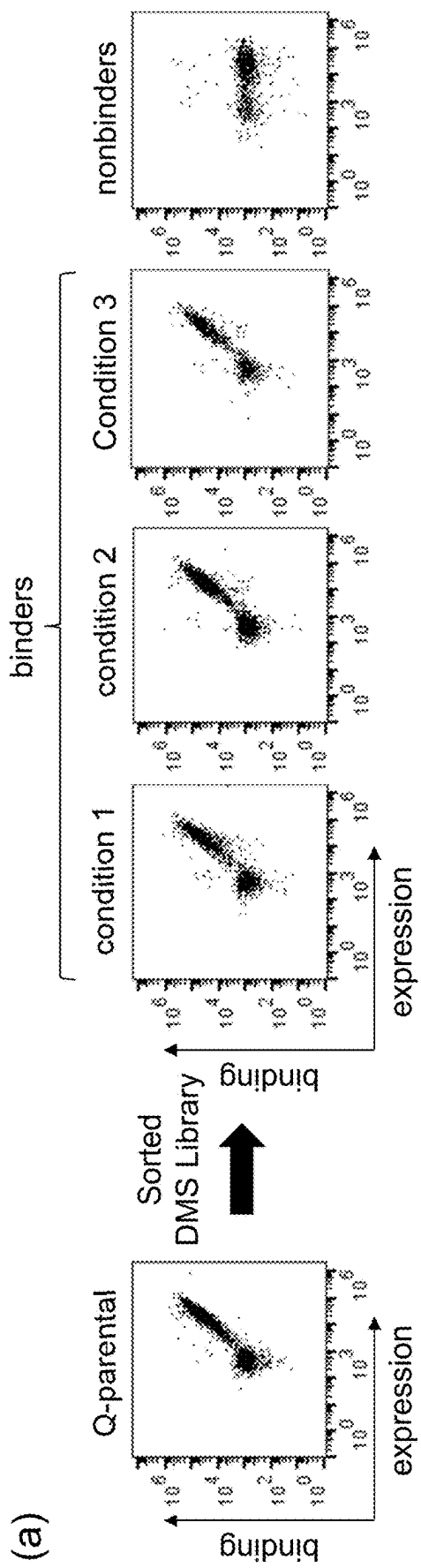


Fig. 24

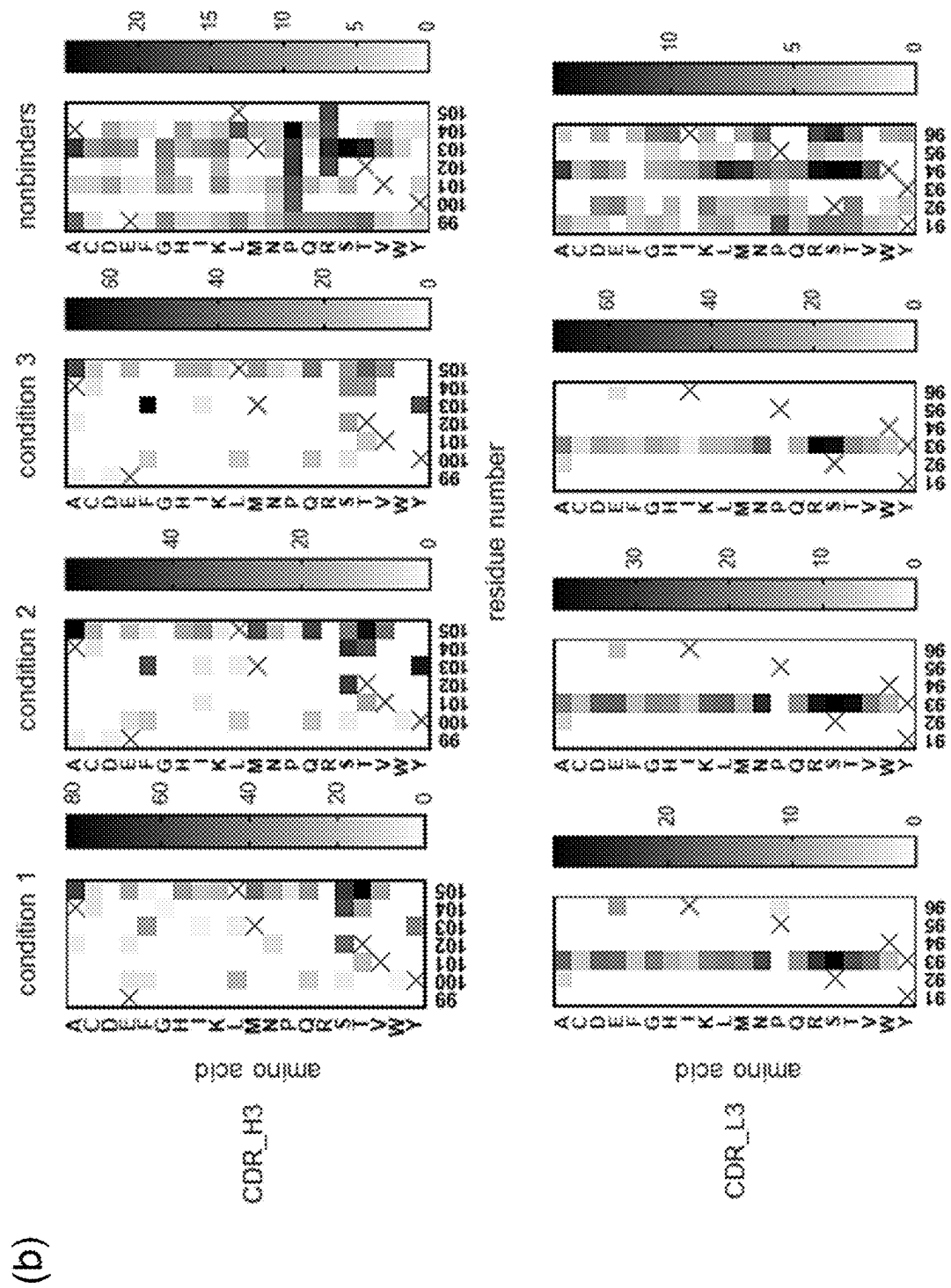


Fig. 24 (continued)

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**COMPOSITIONS AND METHODS
COMPRISING ANTIBODIES THAT BIND TO
COVALENT PEPTIDE CONJUGATES**

**CROSS REFERENCE TO RELATED
APPLICATIONS**

This application is a continuation of U.S. patent application Ser. No. 18/547,623, filed Aug. 23, 2023, which is a National Stage entry of International Patent Application No. PCT/US2022/018171, filed Feb. 28, 2022, which claims the benefit of U.S. Provisional Patent Application No. 63/253,499, filed Oct. 7, 2021, and U.S. Provisional Patent Application No. 63/154,627, filed Feb. 26, 2021, the entire disclosures of each of which are incorporated herein by reference.

**STATEMENT REGARDING FEDERALLY
SPONSORED RESEARCH**

This invention was made with government support under grant nos. CA194864, CA267362 and CA049152 awarded by the National Institutes of Health. The government has certain rights in the invention.

SEQUENCE LISTING

The instant application contains a Sequence Listing, which has been submitted electronically in XML format and is hereby incorporated by reference in its entirety. Said XML copy, created on Jan. 2, 2025, is named 64443-701_301_SL.xml, and is 449,032 bytes in size.

BACKGROUND

There is an ongoing and unmet need for agents that can bind to targets that include drugs that are covalently bound to proteins or peptides. In particular, there is a need to improve the efficacy of targeted therapy and also to increase tumor immunogenicity and the efficacy of immune therapy against cancer driven by intracellular oncogenes. The disclosure is pertinent to these needs.

BRIEF SUMMARY

The present disclosure provides compositions and methods that include binding partners that bind with specificity to target sites on proteins or peptides that comprise a covalently attached molecule. It is believed that this is the first disclosure of binding partners with this binding function. The disclosure illustrates this approach using binding partners in the form of numerous antibodies and antibody derivatives that specifically bind to proteins and peptides that have been covalently modified by attachment of a molecule, wherein the molecules are illustrated by a variety of drugs. Further, the disclosure demonstrates binding partners that bind with specificity to peptides that have been covalently modified by attachment of a small molecule drug are specific for the described covalently modified peptides when presented in the context of a human leukocyte antigen (HLA), wherein HLA is a representative example of a major histocompatibility complex (MHC). Thus, binding partners that are specific for peptide-drug conjugates in an HLA complex are demonstrated. The disclosure includes polynucleotides encoding the described binding partners and cells that are modified to express the binding partners. The disclosure

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includes diagnostic, prophylactic and therapeutic approaches using the binding partners.

BRIEF DESCRIPTION OF THE FIGURES

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FIG. 1. Graphs depicting results of phage enzyme-linked immunosorbent assay (ELISA) of phage-displayed antibody clones. Binding to KRAS^{G12C}-GDP and KRAS^{G12C}-GDP-ARS-1620 conjugate was determined. For each clone, the bars in the graph are, from left to right, Buffer, G12C-GDP-ARS, and G12C-GDP.

FIG. 2. Graph showing 12C-ARS Fab59 binding to KRAS^{G12C} in GTPyS- or GDP-bound nucleotide state with or without ARS-1620 was characterized using the bead binding assay.

FIG. 3. Graphs showing that 12C-ARS-Fab59 specifically binds KRAS^{G12C}-GDP conjugated to ARS-1620.

FIG. 4. Shown in (A) is data obtained using 12C-ARS-Fab59 to measure ARS-1620/KRAS^{G12C} adducts by pull-down assays from lysates prepared from cell lines. Shown in (B) are immunoblots of whole cell lysates and 12C-ARS Fab-pull-downs (PD) from RAS-less MEFs reconstituted with the indicated KRAS mutants (4A) and KCP (Kras^{G12C}; Tp53^{R172H}; Pdx-Cre) mouse pancreas cancer cells (4B), treated in the presence or absence of ARS-1620. Shown in (C) are whole cell lysates and 12C-ARS Fab pull-downs (PD) from H358 and MIAPaCa-2 cells, treated as indicated. Shown in (D) is ARS-adduct formation in samples from C, quantified by LC/MS-MS assay. ARS-1620 and SHP099 concentrations were 10 µM in all panels.

FIG. 5. Data showing that 12C-ARS-Fab59 can be used to measure the engagement of ARS-1620 to mutant KRAS by pull-down assay with lysates prepared from animal tissues.

35 A, B, anti-pan RAS and anti-ERK2 (loading control) immunoblots of lysates and 12C-ARS Fab pull-downs (PD) from LSL-KRAS^{G12C}-Tp53^{R270H} (A) and LSL-KRAS^{G12C} (B) tumors after 3 days of oral gavage with ARS-1620 (200 mg/kg/d) alone or with the SHP2 inhibitor SHP099 (75 mg/kg/d) are shown.

40 FIG. 6. Binding of antibody clones to AMG510 conjugated to the KRAS(G12C) peptide and the poly-Ser peptide. For each antibody clone, the bars are from left to right are No target, KRAS(G12C), KRAS(G12C)-AMG, and Unrelated peptide-AMG. Signals for the two negative controls, no target and the KRAS(G12C) peptide without a conjugated drug, were too low to be visible in the graph. The antibody clones were displayed on the yeast cell surface, and binding of the targets conjugated to fluorescently labeled streptavidin was detected using flow cytometry.

45 FIG. 7. Binding of the P2AMR-1 clone in the human IgG1 format to AMG510 conjugated to the KRAS(G12C) peptide (circles) and the same peptide without drug conjugation (squares). The peptide was immobilized to streptavidin-coated beads, and the antibody bound to the beads were detected with a fluorescently labeled secondary antibody. The apparent K_D value is shown. Data shown here are from triplicate measurements. Error bars are within the size of the symbols.

50 FIG. 8. Recognition of AMG510 presented on class I MHC molecule. The AMG510-RAS (G12C) conjugate (circles) and unconjugated peptide (squares) were loaded onto HLA-A*03:01 and immobilized on streptavidin-coated beads. Antibody binding was detected using the same method as in FIG. 2. The apparent K_D value is shown. Data shown here are from triplicate measurements. Error bars are within the size of the symbols.

FIG. 9. Cartoon representation of the disclosed concept referred to as HaplImmune.

FIG. 10. Data representing development of antibodies that bind MHC/peptide-drug conjugate complexes. (A) Multiplex bead-binding assay (MBBA) of phages displaying different antibody clones. (B) MBBA assay of phages displaying different antibody clones to: HLA-A*01:01 in complex with the BTK peptide conjugated with Ibrutinib. (C) MBBA assay of phages displaying different antibody clones to: HLA-A*02:01 in complex with the EGFR peptide conjugated with Osimertinib.

FIG. 11. Binding titration graphs using the multiplex bead-binding assay (MBBA) of purified antibodies targeted to the KRAS(G12C)-AMG510 conjugate. Clone names are shown over each graph. Antigen nomenclature is described in FIG. 10.

FIG. 12. Graph showing that binding of antibodies to the AMG510-peptide conjugate in complex with an HLA was not affected by the presence of the free drug, AMG510.

FIG. 13. Graphs of binding titrations using the multiplex bead-binding assay (MBBA) of purified antibodies targeted to the BTK-Ibrutinib conjugate. Clone names are shown over each graph. Antigen nomenclature is described in FIG. 10.

FIG. 14. Graphs of binding titrations of purified antibodies to the KRAS(G12C)-AMG510 conjugate presented by endogenous HLA molecules on the cell surface. Raji cells were first incubated with the KRAS (G12C)-AMG510 conjugate or the KRAS(wild type) peptide, and excess conjugate and peptide were washed away. The antibody levels detected using a fluorescently labeled secondary antibody are shown as a function of IgG concentration used for staining. Apparent dissociation constant (K_D) values were determined using nonlinear least-squared fitting of a 1:1 binding function. Data shown here are from triplicate measurements.

FIG. 15. Antibody binding to a KRAS(G12C)-expressing cell line pretreated with AMG510. (A) Flow cytometry histograms. (B) Quantification of the median fluorescence intensity of H358 cells treated with or without AMG510. (C) Quantification of the median fluorescence intensity of H358 cells and HEK293T cells (a negative control) treated with or without AMG510 and stained with the AMRA3-7 antibody.

FIG. 16. Graph showing binding of P2AMR-1 IgG to cells preincubated with the KRAS(G12C) peptide-AMG510 conjugate, KRAS(wild type) peptide, or no peptide.

FIG. 17. Graphs showing binding of purified antibodies in the IgG format to the indicated drug-peptide/HLA complexes as measured using the multiplex bead binding assay (MBBA). (A) Antibody clones identified with AMG510 conjugated to KRAS(G12C) peptide in complex with HLA-A*03:01 as the antigen. (B) Antibody clones identified with AMG510 conjugated to KRAS(G12C) peptide in complex with HLA-A*11:01 as the antigen.

FIG. 18. Graphs showing the cytotoxic effect of single-chain Diabodies (scDbs) on cells pulsed with an exogenous peptide-drug conjugate. (A) Raji cells were first pulsed with AMG510 conjugated to a peptide corresponding to a fragment of KRAS(G12C) or a control peptide corresponding to KRAS(wild type). The pulsed cells were co-cultured with human T cells (Effector:Target=3:1) in the presence of single-chain Diabodies (scDbs) at the indicated concentrations. After incubation, dead cells were stained and detected by flow cytometry. Data shown here are from triplicate measurements. Error bars indicate the s.d. Where error bars are not visible, the errors are smaller than the symbols. (B) Equivalent experiments using T2 cells and Osimertinib

conjugated with an EGFR peptide. As a negative control, peptide conjugated with beta-mercaptoethanol was used.

FIG. 19. Graphs showing specific cytotoxic effect of AMRA3-7_UCHT1 scDb on drug-treated lung cancer cell lines. (A) Lung cancer cell lines were treated with 100 nM AMG510 for 24 hr, then co-cultured with human T-cells (E:T=5:1) in the presence of AMRA3-7_UCHT1 scDb. After incubation, cell viability was measured. The scDb antibody showed a dose-dependent cytotoxic effect only on the AMG510-treated cells with the cognate KRAS mutation (G12C) and HLA (HLA-A3). (B) Cytotoxic effect of scDb at 0.1 nM concentration. Data shown here are from quadruplicate measurements.

FIG. 20. Binding titration curves of AMR-A3-7 and AMR-A3-7D displayed on the yeast cell surface. Binding to HLA-A*03:01 presenting AMG510 conjugated to the Cys residue in the 9mer and 10mer RAS (G12C) peptides, VVGACGVGK (SEQ ID NO: 1) and VVVGACGVGK (SEQ ID NO: 2), respectively, is shown.

FIG. 21. Graphs showing cell killing effects of AMRA3-7D scDb. (A) Dose-dependent cell killing effect of AMRA3-7D scDb tested with Raji cells that were first pulsed with AMG510 conjugated to a peptide corresponding to a fragment of KRAS(G12C) or a control peptide corresponding to KRAS(wild type). (B) Cell killing effect of AMRA3-7D scDb tested against H2122 non-small cell lung cancer cell line treated with AMG510 or DMSO only (negative control).

FIG. 22. Cell binding and killing effects of AMRA3-7D CrossMab. (A) Binding of AMRA3-7D to Jurkat cells, which express CD3 and to Raji cells, which do not express CD3. (B) Dose-dependent cell killing effect of AMRA3-7D CrossMab tested with Raji cells that were first pulsed with AMG510 conjugated to a peptide corresponding to a fragment of KRAS(G12C) or a control peptide, corresponding to KRAS(wild type). (C) Cell killing effect of AMRA3-7D CrossMab tested with H2122 non-small cell lung cancer cell line treated with AMG510 or DMSO only (negative control).

FIG. 23. Deep mutational scanning of AMR-A3-7D. (a) Representative flow cytometry profiles of yeast cells displaying AMRA3-7D and its deep mutational scanning library populations. (b) The prevalence of mutations at each position in the sorted subsets of the deep mutational scanning library is shown in a heat map format. FIG. 23 discloses “GGWYPA” as SEQ ID NO: 155 and “ISYVKKL” as SEQ ID NO: 153.

FIG. 24. Deep mutational scanning of OEA2-5. (a) Representative flow cytometry profiles of yeast cells displaying OEA2-5 in the single-chain Fv format and its deep mutational scanning library populations. (b) The prevalence of mutations at each position in the sorted subsets of the deep mutational scanning library is shown in a heat map format. FIG. 24 discloses “EYVTMAL” SEQ ID NO: 159 and “YSYWPI” as SEQ ID NO: 157.

DETAILED DESCRIPTION

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs.

Every numerical range given throughout this specification includes its upper and lower values, as well as every narrower numerical range that falls within it, as if such narrower numerical ranges were all expressly written herein.

As used in the specification and the appended claims, the singular forms “a” “and” and “the” include plural referents unless the context clearly dictates otherwise. Ranges may be expressed herein as from “about” one particular value, and/or to “about” another particular value. When such a range is expressed, another embodiment includes from the one particular value and/or to the other particular value. Similarly, when values are expressed as approximations, by the use of the antecedent “about” it will be understood that the particular value forms another embodiment. The term “about” in relation to a numerical value is optional and means for example +/-10%.

This disclosure includes every amino acid sequence described herein and all nucleotide sequences encoding the amino acid sequences. Every antibody sequence and antigen binding fragments of them are included. Polynucleotide and amino acid sequences having from 80-99% similarity, inclusive, and including and all numbers and ranges of numbers there between, with the sequences provided here are included in the invention. All of the amino acid sequences described herein can include amino acid substitutions, such as conservative substitutions, that do not adversely affect the function of the protein that comprises the amino acid sequences. In this regard, the disclosure provides alternative residues for certain positions in described binding partner as described below. In certain examples, the alternative residues were identified by deep mutational scanning, which demonstrates binding functionality for each binding partner that contains the described amino acid change(s). The disclosure includes each binding partner with each alternative residue substituted for the original residue alone and in any combination with the described alternative residues. Thus, any binding partner described herein may have any single described residue change or a combination of described changes. Representative changes for particular antibodies are described in Table A, Table B, Table C, and Table D. The changes may be in CDR1, CDR2, CDR3, and combinations thereof. The changes can also include amino acid insertions. The disclosure includes each amino acid sequence that is encompassed by the description of alternative amino acids by reference to a specific sequence identifier and those described in the aforementioned Tables.

As described above, the present disclosure provides antibodies and antigen binding fragments thereof (collectively “binding partners” and each individually a “binding partner”). The term “antibody” includes each binding partner format herein. The binding partners bind with specificity to a protein or fragment thereof, or a peptide provided in peptide form, that comprises a covalently attached molecule. The covalently attached molecule forms a peptide conjugate. A “peptide conjugate” as used herein means any protein or peptide that has been modified so that it is covalently conjugated to another molecule. The peptide conjugate is considered to be a novel antigen, i.e., a neoantigen. The other molecule that is covalently conjugated to the protein or peptide to form the peptide conjugate is not particularly limited, with the proviso that the other molecule is not an additional amino acid that is added to the described peptide conjugates. In embodiments, the molecule that is covalently conjugated to the protein or peptide has or had biological activity before conjugation, or it may be biologically inert before conjugation. In embodiments, the molecule is a drug, including but not necessarily limited to small molecule drugs. Representative and non-limiting examples of drugs that covalently attach to a peptide or protein to form a peptide conjugate are described below. Peptide conjugates include but are not limited to covalently modified full length

proteins and fragments thereof. Peptide conjugates include fragments of full length proteins that include a covalent modification and are produced, for example, by intracellular processing. In certain embodiments, a full length protein may be covalently modified within a cell and subsequently processed such that a peptide conjugate that is a fragment of the full length protein is produced. As described further below, the produced peptide conjugate may be displayed on a cell surface. The cell surface display of the peptide conjugate may be any form of cell surface display, including but not limited to by way of any receptor having an extracellular segment, or it may be displayed by way of any type of major histocompatibility complex (MHC) or human leukocyte antigen (HLA). Non-limiting examples of HLA types that display peptide conjugates, and to which the described binding partners bind with specificity, are described further below.

In embodiments, the binding partners preferentially bind to the protein or peptide or a complex comprising the protein or peptide when covalently bound to the peptide conjugate, relative to the same protein or peptide that is not bound to the drug. Accordingly, binding partners described herein either do not detectably bind, or bind with a lower affinity, to the same protein or fragment thereof in the absence of the covalently attached molecule. In embodiments, the binding partners bind to the protein or peptide comprising the covalently attached drug with an affinity that is 10-10,000 fold, including all numbers and ranges of numbers from 10-10,000, greater than the affinity for the protein or peptide that does not comprise the covalently bound molecule. In this regard, and without intending to be bound by any particular theory, it is considered that the presence of the covalently bound molecule contributes to the presence of an epitope to which the binding partners bind with specificity.

In embodiments, the molecule that is covalently bound to form the peptide conjugate is a drug and may be any targeted covalent inhibitor (TCI), but an inhibition property is not necessarily required. In embodiments, the molecule reacts with a specific residue within the target protein. In embodiments, the molecule reacts at least in part with a segment of the protein or peptide that comprises a nucleophilic, or an electrophilic, residue. In embodiments, the segment of the protein or peptide to which the molecule reacts comprises any of Cys, Lys, Tyr, and His. In embodiments, the molecule reacts at least in part with a segment of the protein or peptide that comprises a wild type Cys, or a mutation of a residue to a Cys, and thus may be covalently attached by a so-called sulfur tether. In embodiments, the drug is any drug described in Ghosh A K, Samanta I, Mondal A, Liu W R. Covalent Inhibition in Drug Discovery. ChemMedChem. 2019; 14(9): 889-906. doi:10.1002/cmdc.201900107, or in De Cesco, et al., European Journal of Medicinal Chemistry 138 (2017) 96e114, or in Bauer, RA, Drug Discovery Today, Volume 20, Number 9, September 2015, from which the disclosures of compounds that covalently modify protein targets is incorporated herein by reference.

In non-limiting embodiments, any of said Cys, Lys, Tyr, and His amino acids are present in the protein or peptide to which the molecule binds because the wild type protein has been mutated to include one or a combination of the described residues. In non-limiting embodiments, the molecule binds to a protein or peptide that is correlated with a condition, such as a cancer. In embodiments, the target (e.g., the protein or peptide to which the molecule covalently binds) is a receptor, including but not necessarily limited to any receptor having a catalytically active segment. In embodiments, the drug binds to an enzyme that is not

necessarily a receptor, including but not limited to any kinase. In embodiments, a protein target comprises a receptor with one or more activating mutations, which promote ligand-independent enzyme activity.

In embodiments, the molecule targets and thus covalently binds to an amino acid sequence present within any of the following proteins and/or variants thereof, which may or may not comprise a mutation, such as a mutation that is related to a particular condition, including but not limited to any type of cancer. In embodiments, the protein is any protein described in Visscher M, et al., Covalent targeting of acquired cysteines in cancer. *Curr Opin Chem Biol.* 2016; 30:61-67. doi: 10.1016/j.cbpa.2015.11.004, from which the description is incorporated herein by reference. Visscher et al. also teaches methods for identifying disease-associated mutated genes that introduces a Cys residue suitable for covalent modification. In embodiments, the protein is KRAS, Bruton's tyrosine kinase (BTK), any member of the epidermal growth factor receptor (EGFR) family, also referred to as the ERBB family, including but not limited to EGFR (ERBB1), HER2/NEU (ERBB2), HER3 (ERBB3), and HER4 (ERBB4); a fibroblast growth factor receptor (FGFR); the receptor kinase known in the art as MET, BRAF, a cyclin-dependent kinase (CDK); Acetyl Choline Esterase (ACHE); TP53, IDH1, GNAS, FBXW7, CTNNB1, DNMT3A, any cathepsin, including cathepsin B, C, F, H, K, L, O, S, V, W and X; any caspase; any protein involved in obesity, such as Pancreatic lipase and METAP2, or any Cancer Testis Antigen. In embodiments, the drug targets and therefore covalently binds to any viral protein, including but not limited to a polymerase, including any viral DNA polymerase, RNA polymerase, reverse transcriptase, or RNA-dependent RNA polymerase, or a viral protein that is required, for example, viral cell entry, or a protein encoded by any a transposable element. In embodiments, the drug targets EGFR and may be selected from PD168393, PF00299804 (dacomitinib), EKB569 (pelitinib), afatinib, WZ4002, osimertinib (formerly known as AZD9291), PF-06459988, nazartinib, naquitinib, olmutinib, avitinib, and rociletinib, neratinib, pyrotinib, poziotinib, and derivatives thereof. In embodiments, the drug targets Bruton's tyrosine kinase (BTK), and may be selected from ibrutinib, acalabrutinib, zanubrutinib, CHMFL-BTK-11, ONO/GS-405, PRN1008, and CC-292. In embodiments, the drug targets any p90 ribosomal S6 kinase (RSK), and may be selected from fluoromethylketone (FMK) and dimethyl fumarate. In embodiments, the drug targets any FGFR, and may be selected from FIIN-1, FIIN-2, FIIN-3, BGJ398, AZD4547, PRN1371, FGF401.

In a non-limiting embodiment, the binding partner binds with specificity to a site comprised by a neoantigen that includes a covalently linked small molecule drug or other covalently linked molecule as a component of an antigen in a specific MHC context.

In embodiments, the molecule that becomes covalently bound to form the peptide conjugate targets any RAS oncogene protein product, including but not necessarily limited to HRAS, NRAS, KRAS4A, and KRAS4B. The amino acid sequences of RAS proteins are known in the art, and residue numbering is identical for the relevant part of all RAS isotypes that are discussed in this disclosure for which the amino acid sequence is available from, for example, UniProt P01116, from which the amino acid sequence is incorporated herein as of the effective filing date of this application or patent. The G12 position is numbered according to the known amino acid sequence, regardless of whether

or not the G12 is the twelfth amino acid in an express RAS peptide sequence of this disclosure.

In one embodiment, the molecule covalently binds to a KRAS protein or peptide that comprises a mutation. In embodiments, the mutation is at least one of KRAS residues 12, 13, or 61. Reference to any drug herein includes its name in capitalized and un-capitalized form.

In embodiments, the drug targets a KRAS protein comprising a KRAS G12C mutation. In non-limiting embodiments, the drug that targets a KRAS protein is selected from 2E07, 6H05, SML-8-73-1, MRTX849, JNJ74699157, LY3499446, ARS-853, ARS-1620, MRTX1275, AMG510, or derivatives thereof. In an embodiment the drug comprises a proteolysis targeting chimera (PROTAC) derivative of a covalent drug, a non-limiting description of which is available in doi: 10.1021/acscentsci.Oc00411, from which the description of PROTACs is incorporated herein by reference. In embodiments, the PROTAC is LC-1 or LC-2. In embodiments, the disclosure relates to an autophagy-mediated degrader, referred to as an AUTAC, as described in doi.org/10.1080/15548627.2020.1718362, from which the description of AUTACs is incorporated herein by reference.

A non-limiting example of a binding partner that binds to KRAS(G12C)-AMG510 is referred to herein as AMRA3-7D. The disclosure includes all derivatives of AMRA3-7D that are described herein, including the alternative residues described below by way of deep mutational analysis, and in the forms of an scDb and a CrossMab, for which representative amino acid sequences are provided. The amino acid sequence of the light chain (V_L) and heavy chain (V_H) of AMRA3-7D are:

V_L :
(SEQ ID NO: 3)
DIQMTQSPSSLSASVGDRVTITCRASQSSVSSAVAWYQQKPGKAPKLLI

$SASSLYSGVPSRFSGRSGTDFTLTISSLQPEDFATYYCQQISYVKKLI$

$TFGQGTKEIKRTV$
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 168, respectively, in order of appearance)

V_H :
(SEQ ID NO: 4)
EVQLVESGGGLVQPGGSLRLSCAASGFTFSDYSIHWVRQAPGKGLEWVA

$SISSSSGSTSYADSVKGRTISADTSKNTAYLQMNSLRAEDTAVYYCAR$

$GGWYPAMDYWGQGTLVTVSS$
(Bolded CDR sequences are disclosed as SEQ ID NOS 169, 170, and 171, respectively, in order of appearance)

A non-limiting example of a binding partner that binds to an Epidermal Growth Factor receptor (EGFR)-osimertinib conjugate is referred to herein as OEA2-5. The disclosure includes all derivatives of OEA2-5 that are described herein, including the alternative residues described below by way of deep mutational analysis, and in the form of an scDb, for which representative amino acid sequences are provided.

The amino acid sequences of the light chain (V_L) and heavy chain (V_H) of OEA2-5 are:

V_L :

(SEQ ID NO: 5)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISSLQPEDFATYYCQQYSYWPITF

GQGKTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 172, respectively, in order of appearance)

V_H :

(SEQ ID NO: 6)
EVQLVESGGGLVQPGGSLRLSCAASGFTISSLSSYIHWWVRQAPGKGLEWVA

YISPSYGSTSYADSVKGRTFISADTSKNTAYLQMNSLRAEDTAVYYCAR

EYVTMADYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 173, 174, and 175, respectively, in order of appearance)

In embodiments, the binding partner binds to a protein in its native form, with the exception that the drug or other molecule is covalently attached to it. "Native form" means the intact protein that retains its biological function before covalent attachment of the drug or other molecule. In embodiments, the native form or the protein is its form before being fragmented such as by intracellular processing. In embodiments, the binding proteins therefore bind to full length polypeptides that are covalently attached to the drug or other molecule and wherein the covalently bound drug or other molecule at least in part permits the preferential binding of the binding partners. In general, a polypeptide, which is used interchangeably herein with the term "protein," comprises more than 50 contiguous amino acids. In embodiments, a binding partner binds with specificity to an intact protein that is covalently attached to a drug or other molecule. In other embodiments, the binding partners bind with specificity to a peptide comprising the covalently bound molecule. In embodiments, the binding partner binds with specificity to a peptide having a specific amino acid sequence and is covalently conjugated to another molecule, such as a drug. In embodiments, the binding partner binds preferentially to a peptide covalently bound to a molecule such as a drug, where the sequence of the peptide is not relevant. This preferential binding is relative to binding to the same peptide that is not conjugated to the drug. In embodiments, the binding partner binds preferentially to a peptide comprising a KRAS(G12) mutation, or to a variant thereof, wherein the variant is at least 50% similar to the KRAS(G12)-containing peptide. This preferential binding is relative to binding to a KRAS(G12)-containing peptide, or the variant thereof, respectively, that is not covalently conjugated to the drug or other molecule.

In embodiments, the described binding partners bind with specificity to peptide conjugates that are of suitable length to be presented in a major histocompatibility complex (MHC), referred to as human leukocyte antigen (HLA) in humans, or to MHC or its equivalent complex in non-human animals, including but not limited to non-human mammals.

In general, the peptide conjugate comprises fewer than 50 contiguous amino acids. In embodiments, peptide conjugates which comprise the described epitope may therefore be from 2-49 amino acids in length. In embodiments, the peptide to which the drug or other molecule is covalently attached, and which attached drug may be comprised by the

epitope, comprises from 4-12 contiguous amino acids, which may or may not be derived from a longer protein during the processing of a protein, such as an antigen processed for presentation by an MHC molecule. In embodiments, the drug is conjugated to a peptide that comprises, or consists of 7-30 amino acids. In embodiments, the drug or other molecule is conjugated to a peptide that comprises, or consists of, 7, 8, 9, 10, 11, 12, 13, 14, or 15 amino acids, and which may be presented in an MHC Class I context. In

10 embodiments, the drug or other molecule is conjugated to a peptide that is 9-30 amino acids, inclusive, and including all numbers and ranges of numbers there between, and which may be presented in an MHC Class II context. In embodiments, the drug or other molecule is conjugated to a peptide comprises at least 7 amino acids.

In embodiments, a binding partner binds with specificity to a peptide conjugate that is covalently conjugated to a drug or other molecule independent of MHC presentation. In embodiments, non-limiting examples of which are described 20 below in Example 3, the binding partner binds with specificity to the peptide conjugate only when the peptide conjugate is presented by an MHC molecule. In embodiments, the binding partner can bind with specificity to a peptide conjugate in both an MHC-independent and an MHC-presentation context. In embodiments, the MHC-peptide conjugate complex comprises an antigen to which a described binding partner binds with specificity.

In embodiments, the binding partners accordingly can bind to cells via any MHC that can present peptide conjugates. In embodiments, the HLA is expressed by cells that are restricted to Class I, Class II, or Class III MHC presentation. In embodiments, the binding partners can bind to cells that express Class I MHC that presents the peptide conjugate. Those skilled in the art will recognize that Class 35 I MHC includes, among other components, a polymorphic α chain and $\beta 2$ microglobulin, wherein the peptide conjugate binds to the polymorphic chain.

In embodiments, the cells are antigen presenting cells (APCs). In embodiments, the cells are so-called professional 40 antigen presenting cells, and thus may include but are not limited to macrophages and dendritic cells, which display Class II MHC. Those skilled in the art will recognize that Class II MHC includes, among other components, MHC polymorphic α and β chains, and the displayed peptide conjugate binds to both chains. In other embodiments, Class 45 II MHC may be displayed with the peptide conjugate by other cell types, such as cancer/tumor cells, and thus the disclosure provides for direct recognition of such cells using the described binding partners, without requirement for a professional APC.

In embodiments, the peptide conjugate is displayed by a non-classical MHC complex, which may include CD1d, MR1, MHC-E, -F, -G and/or other emerging family members that will be recognized by those skilled in the art.

55 The disclosure includes binding partners that bind with specificity to a peptide conjugate displayed only by a specific MHC type, and thus provides binding partners that discriminate between MHC types. Representative examples of such binding partners are described herein at least by way of FIG. 17.

In embodiments, a binding partner of this disclosure can bind with specificity to a peptide conjugate comprising a covalently conjugated drug or other molecule that is displayed by more than one specific MHC type. In embodiments, a binding partner of this disclosure can bind with specificity to a peptide conjugate comprising a covalently conjugated drug only in a specific MHC context. In embodi-

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ments, the peptide conjugate is displayed by an MHC class I type selected from HLA-A, -B, -C, and combinations thereof. In certain aspects, the peptide conjugate is displayed in the context of any MHC class I that is A*02/B*35/C*04. In embodiments, the peptide conjugate is displayed by any of MHC of class II that is DR*01/DR*04/DR*07/DP*04. In embodiments, the HLA comprises A*01:01, A*02:01, A*03:01, A*11:01, A*24:02, A*26:01, B*07:02, B*08:01, B*27:05, B*39:01, B*40:01, B*58:01, or B*15:01. Specific examples of antibodies include antibodies that bind to KRAS(G12C)-AMG510 conjugate presented on HLA-A*03:01 and HLA-A*11:01, BTK-Ibrutinib conjugate presented on HLA-A*01:01, and EGFR-Osimertinib conjugate presented on HLA-A*02:01. In non-limiting embodiments, the disclosure provides scDbs that are specific for a particular drug that is covalently bound to a described peptide that is present on a specific HLA, or the same drug that is covalently bound to a described peptide that is present on two different HLAs. Representative scDbs are described in Example 4. Data obtained using the scDbs are presented via FIGS. 18 and 19. Data obtained using CrossMab formats are provided in Example 5 and its accompanying figures.

In embodiments, the peptide conjugate is displayed by cells that participate in, or can be the targets of, cell-mediated immune responses. In embodiments the peptide conjugate that is displayed in any suitable MHC context is comprised by a cell that is recognized by a leukocyte, including but not necessarily limited to a T cell or a natural killer (NK) cell. In embodiments, the T cell is a CD4+ T cell, a CD8+ T cell, a double positive CD4+/CD8+ T cell, a CD4+/CD8+ double negative T cell, or a γδ T cell. Thus, and as described further below, the disclosure provides binding partners that are configured to interact with both the presented peptide conjugate and cells that participate in cell-mediated immune responses. In embodiments, certain described binding partners are capable of binding to a complex of 1) a specific MHC and 2) a specific peptide conjugate. In embodiments, certain described binding partners are capable of being bound to a specific peptide conjugate presented by at least two different MHCs.

In embodiments, any binding partner of this disclosure comprises at least one chain that comprises a complementary determining region (CDR) that is CDR1, CDR2, or CDR3 from any heavy or light chain amino acid sequence described herein. In certain examples in the present specification, the CDRs are shown in bold font. The amino acid sequences of the CDR sequences are separately encompassed by this disclosure by way of their positions in the described heavy and light chain amino acid sequences. The disclosure includes binding partners that comprise a described heavy chain CDR1, CDR2, and CDR3. The disclosure also includes binding partners that comprise a described light chain CDR1, CDR2, and CDR3. The disclosure also includes binding partners that comprise a described heavy chain CDR1, CDR2, and CDR3 and a described light chain CDR1, CDR2, and CDR3. For amino acid sequences of this disclosure that include amino acids that comprise purification or protein production tags, such as HIS tags and/or AVI-tags, the disclosure includes the proviso that the sequences of the described tags may be excluded from the amino acid sequences. Amino acids between the described tags may also be excluded.

Binding partners of this disclosure can be provided as intact immunoglobulins or as fragments of immunoglobulins, including but not necessarily limited to antigen-binding (Fab) fragments, Fab' fragments, (Fab')₂ fragments, Fd (N-terminal part of the heavy chain) fragments, Fv frag-

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ments (two variable domains), diabodies (Dbs), dAb fragments, single domain fragments or single monomeric variable antibody domains, single-chain Diabodies (scDbs), isolated complementary determining regions (CDRs), single-chain variable fragment (scFv), and other antibody fragments that retain antigen binding function. In embodiments, one or more binding partners are provided as a component of a Bi-specific T-cell engager (BiTE), bispecific killer cell engager (BiKE), CrossMab (e.g., a binding partner containing four different chains; immunoglobulin crossover (also known as Fab domain exchange or CrossMab format) technology (see e.g., WO2009/080253; Schaefer et al., Proc. Natl. Acad. Sci. USA, 108:11187-11192 (2011).), or a chimeric antigen receptor (CAR), such as for producing chimeric antigen receptor T cells (e.g., CAR T cells) and CAR natural killer (NK) cells, and killer macrophages. The disclosure includes binding partners that include the described heavy and light chain variable regions.

In embodiments, the binding partners are multivalent. In embodiments, a tri-specific binding partner is provided. In embodiments, cells express at least a segment of one or more binding partners in the form of a CAR. In an embodiment, a binding partner of this disclosure may be provided as a complex with a polynucleotide, such as an RNA polynucleotide, to form an aptamer. In embodiments, a multi-valent binding partner includes one binding component, such as a paratope, that confers specificity to a particular target on a desired cell type, such as any cancer cell marker. In embodiments, a tri-specific leukocyte engager is provided. In embodiments, the binding partners may be part of a molecule that is activated only in the presence of a protease or other enzyme present in a tumor microenvironment, such embodiments being pertinent to, for instance, a probody, examples of which are known in the art, for example in doi: 10.1126/scitranslmed.3006682, doi: 10.1038/s41467-020-16838-w, and doi: 10.1038/s41587-019-0135-x, from which the descriptions of probodies, and protease activation, are incorporated herein by reference. In an embodiment, the disclosure provides a universal hapten that can be grafted onto inhibitors.

In embodiments, a CAR of this disclosure comprises scFv that comprises heavy and light chains as described herein. As is known in the art for previously described CARs, the scFv is present in a contiguous polypeptide that further comprises a CD3zeta chain and a costimulatory domain. In embodiments, the costimulatory domain comprises a 4-1BB costimulatory domain or a CD28 costimulatory domain. A CAR may also contain a co-receptor hinge sequence, such as a CD8 a co-receptor hinge sequence.

In embodiments, binding partners of this disclosure may comprise a constant region, e.g., an Fc region. Any isotype of constant region can be included. Binding partners that comprise a constant region may be particularly adapted for antibody-dependent cell mediated cytotoxicity (ADCC) and thus may function to kill targeted cells by cell-mediated responses by any of a variety of effector cells. Similarly, a constant region may be particularly adapted for enhancing complement-mediated responses.

In embodiments, a binding partner of this disclosure may be modified such that it is present in a fusion protein. In embodiments, an antigen binding segment of a binding partner may be present in a fusion protein, and/or the constant region may be a component of a fusion protein. In embodiments, a fusion protein comprises amino acids from at least two different proteins. Fusion proteins can be produced using any of a wide variety of standard molecular biology approaches, including but not necessarily limited to

expression from any suitable expression vector. In embodiments, a binding partner described herein may be present in a fusion protein with a detectable protein, such as green fluorescent protein (GFP), enhanced GFP (eGFP), mCherry, and the like. In embodiments, as an alternative to an expression vector, an mRNA or chemically modified mRNA encoding any binding partner described herein can be delivered to cells such that the binding partner is translated by the cells.

In embodiments, binding partners described herein are used to carry drugs or toxins, and thus the binding partners may be provided as immunotoxins, or in the form of antibody-drug conjugates (ADCs).

In embodiments, agents useful in the generation of immunotoxins include enzymatically active toxins and enzymatically active fragments thereof. Suitable enzymatically active toxins include but are not limited to diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from *Pseudomonas aeruginosa*), ricin A chain, abrin A chain, modeccin A chain, alpha sarcin, *Aleurites fordii* proteins, dianthin proteins, *Phytolaca americana* proteins (PAPI, PAPII, and PAP-S), *Momordica charantia* inhibitor, curcin, crotin, *Sapaponaria officinalis* inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin and the tricothecenes. These can be provided as components of fusion proteins or can be covalently attached to the binding partner by any suitable conjugation approach.

The binding partner may be connected to a chemotherapeutic agent by using any suitable linker to form an antibody drug conjugate (ADC). In embodiments, the linker comprises a disulfide, a hydrazine, or a thioether. The chemotherapeutic agent may be reversibly or irreversibly attached to the binding partner.

Cleavable linkers may be particularly useful for killing bystander cells. In embodiments, a protease recognition site may be included to liberate the chemotherapeutic agent from the binding partner by operation of a protease that recognizes and cleaves at the protease recognition site. The ADC may therefore be considered to contain a prodrug.

In embodiments, binding partners of this disclosure may comprise linking sequences. As a non-limiting example, an ScFv may comprise a linker that links segments comprising paratopes to one another. Suitable amino acid linkers may be mainly composed of relatively small, neutral amino acids, such as glycine, serine, and alanine, and can include multiple copies of a sequence enriched in glycine and serine. In specific and non-limiting embodiments, the linker comprises 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, and 20 amino acids. In an example, the linker may be the glycine-serine-alanine linker G₄SA₃ (SEQ ID NO: 11) or a glycine-serine linker (G₂S)₄ linker (SEQ ID NO: 12). In embodiments, a binding partner may include a cellular localization signal, or a secretion signal. In embodiments, binding partner may comprise a transmembrane domain, and thus may be trafficked to, and anchored in a cell membrane. For secretion, any suitable secretion signal can be used, and many are known in the art.

In embodiments, the binding partners can be part of an ADC and therefore the binding partners comprise a drug. The drug can include, but is not necessarily limited to, any suitable chemotherapeutic agent. In embodiments, the ADC comprises a binding partner and a chemotherapeutic agent that is an anti-microtubule agent, an alkylating agent, or a DNA minor groove binding agent. In embodiments, the chemotherapeutic agent comprises a maytansinoid, a dolastatin, an auristatin drug analog, or a cryptophycin. In embodiments, the chemotherapeutic agent is a duocarmycin

derivative, or an antibiotic, such as an enediyne antibiotic, or pyrolobenodiazepine (PBD), including dimers thereof. In embodiments, the chemotherapeutic agent is an enzyme inhibitor, such as a topoisomerase or polymerase inhibitor.

In embodiments, the chemotherapeutic agent comprises doxorubicin, or a metal-containing compound, such as a platinum-containing compound, non-limiting examples of which include cisplatin, carboplatin or oxaliplatin. In embodiments, the ADC comprises a binding partner described herein, and any drug that is described in Barf and Kaptein, dx.doi.org/10.1021/jm3003203, J. Med. Chem. 2012, 55, 6243-6262, or in Wilson et al., dx.doi.org/10.1021/jm400224q, J. Med. Chem. 2013, 56, 7463-7476, or Lambert and Morris, Adv Ther (2017) 34:1015-1035, from which the descriptions of drugs for use as components as ADCs is incorporated herein by reference. In embodiments, the binding partner is conjugated to or otherwise includes a cytokine, including but not necessarily limited to an interleukin, including but not limited to IL-2 and IL-12, or an interferon (IFN), to thereby provide a cytokine conjugate.

For production of binding partners, any suitable expression system may be used. In general, polynucleotides encoding binding partners are used to express the binding partners in any suitable cell system, non-limiting embodiments of which include NS0 murine myeloma cells, human cell lines, and Chinese hamster ovary (CHO) cells. In embodiments, the disclosure provides a polynucleotide that can selectively hybridize to a polynucleotide encoding any CDR or combination of CDRs described herein. In embodiments, the polynucleotide selectively hybridizes to a polynucleotide encoding a heavy chain CDR1, CDR2, and CDR3 of any described binding partner. In embodiments, the polynucleotide selectively hybridizes to a polynucleotide encoding a light chain CDR1, CDR2, and CDR3 of any described binding partner. In embodiments, the polynucleotide selectively hybridizes to a polynucleotide encoding CDR1, CDR2, and CDR3 of a heavy and light chain of any described binding partner.

In embodiments, a binding partner described herein may be a component of a fusion protein. In embodiments, such as for a binding partner that is produced as a fusion protein, a peptide linker may be used. In embodiments, the peptide linker comprises any self-cleaving signal. In embodiments, the self-cleaving signal may be present in the same open reading frame (ORF) as the ORF that encodes the binding partner. A self-cleaving amino acid sequence is typically about 18-22 amino acids long. Any suitable sequence can be used, non-limiting examples of which include: T2A (EGRGSSLTCGVEDENPGP (SEQ ID NO: 7)); P2A (ATNFSLKKQAGDVENPGP (SEQ ID NO: 8)); E2A (QCTNTYALKLAGDVESNPGP (SEQ ID NO: 9)) and F2A (VKQTLNFDLKLADGVESNPGP (SEQ ID NO: 10)).

To the extent any segment of a protein comprising a binding partner described herein was a component of a library, including but not necessarily limited to a phage display library or a yeast surface display library, the disclosure includes the proviso that the binding partner may be free of any segment of the library that comprises a bacteriophage or yeast amino acid sequence, including but not limited to phage coat protein or a yeast host protein, including but not limited to Aga2. Thus, in certain embodiments, the binding partner may be present in a fusion protein, but the fusion protein does not comprise bacteriophage coat protein. In embodiments, any binding partner described herein may be free of any of pIII phage coat protein, or any part of M1, fd filamentous phage, T4, T7, or λ phage protein.

In embodiments, a binding partner of this disclosure comprises a detectable label, which may be used for diagnostic or therapeutic purposes. For example, a detectable label can be used for localization of the binding partner for pathology and/or in vivo imaging approaches. In embodiments, a binding partner is conjugated to any of a variety of radioactive agents, including but not limited to a highly radioactive atom, such as In111, At211, I131, I125, Y90, Re186, Re188, Sm153, Bi212, P32, Pb212, and radioactive isotopes of Lu. In particular embodiments, such as for imaging, the binding partner may be conjugated to a radioactive atom for scintigraphic approaches, for example Tc99m (metastable technetium-99), 1123, or a spin label for nuclear magnetic resonance (NMR) imaging (also known as magnetic resonance imaging, or "MRI"), such as 1123, 1131, 1124, F19, C13, N15, O17 or Gadlinium (III) or Manganese (II). In embodiments, the radioactive agent is suitable for use in CAT scan or PET imaging. In embodiments, Indium 111, Technetium99 or Iodine131 can be used for planar scans or single photon emission computed tomography (SPECT). Positron emitting labels such as Fluorine19 Iodine 123 and Iodine 124 can be used in positron emission tomography. Paramagnetic ions such as Gadlinium (III) or Manganese (II) can be used in magnetic resonance imaging MRI. In embodiments, the described radioactive isotopes that are attached to a described binding partner can also be used in therapeutic approaches. In embodiments, radioactive agents or isotopes include alpha-emitting radionuclides. In embodiments, radioactive agents or isotopes include beta-emitting radionuclides. In some embodiments, the present disclosure provides an antibody of the present technology conjugated to a diagnostic or therapeutic agent. The diagnostic agent may comprise a radioactive or non-radioactive label, a contrast agent (such as for magnetic resonance imaging, computed tomography or ultrasound), and the radioactive label can be a gamma-, beta-, alpha-, Auger electron-, or positron-emitting isotope. A diagnostic agent is a molecule which is administered conjugated to an antibody moiety, i.e., antibody or antibody fragment, or subfragment, and is useful in diagnosing or detecting a disease by locating the cells containing the antigen.

Any binding partner described herein may be fully or partially humanized. Techniques for humanization of antibodies are known in the art and can be adapted for use in the present disclosure. In embodiments, humanization may be performed, for example, by CDR-grafting. In embodiments, for humanization or to otherwise improve a characteristic of the binding partners, one or more amino acids in a variable region can be changed. In embodiments, one or more amino acids in a framework region can be changed.

The disclosure includes binding partners for use in diagnostic and therapeutic approaches. For therapeutic approaches, in certain embodiments, binding partners may be delivered as mRNA or DNA polynucleotides that encode the binding partners. It is considered that administering a DNA or RNA encoding any binding partner described herein is also a method of delivering such binding partners to an individual or one or more cells. Methods of delivering DNA and RNAs encoding proteins are known in the art and can be adapted to deliver the binding partners, given the benefit of the present disclosure. In embodiments, one or more expression vectors are used and comprise viral vectors. Thus, in embodiments, a viral expression vector is used. Viral expression vectors may be used as naked polynucleotides, or may comprise any of viral particles, including but not limited to defective interfering particles or other replication defective viral constructs, and virus-like particles. In embodiments,

the expression vector comprises a modified viral polynucleotide, such as from an adenovirus, a herpesvirus, or a retrovirus. In embodiments, a retroviral vector adapted from a murine Moloney leukemia virus (MLV) or a lentiviral vector may be used, such as a lentiviral vector adapted from human immunodeficiency virus type 1 (HIV-1).

In an embodiment, an oncolytic viral vector is used. Oncolytic viruses (OVs), including vaccinia (OVV), mediate anticancer effects by both direct oncolysis and stimulation of innate immune responses through production of damage-associated molecular patterns (DAMPs) and the presence of virus-derived pathogen-associated molecular patterns (PAMPs), leading to increased type I interferon production. Additionally, OVV-mediated oncolysis may facilitate the direct acquisition of tumor-derived antigens by host antigen-presenting cells within the tumor microenvironment, thereby leading to improved T cell priming as well as coordination of the effector phase of antitumor immune responses. In alternative embodiments, a recombinant adeno-associated virus (AAV) vector may be used. In certain embodiments, the expression vector is a self-complementary adeno-associated virus (scAAV).

Pharmaceutical formulations containing binding partners are included in the disclosure and can be prepared by mixing them with one or more pharmaceutically acceptable carriers. Pharmaceutically acceptable carriers include solvents, dispersion media, isotonic agents, and the like. The carrier can be liquid, semi-solid, e.g. pastes, or solid carriers. Examples of carriers include water, saline solutions or other buffers (such as phosphate, citrate buffers), oil, alcohol, proteins (such as serum albumin, gelatin), carbohydrates (such as monosaccharides, disaccharides, and other carbohydrates including glucose, sucrose, trehalose, mannose, mannitol, sorbitol or dextrins), gel, lipids, liposomes, resins, porous matrices, binders, fillers, coatings, stabilizers, preservatives, liposomes, antioxidants, chelating agents such as EDTA, salt forming counter-ions such as sodium; non-ionic surfactants such as TWEEN, PLURONICS or polyethylene glycol (PEG), or combinations thereof. In embodiments, a liposomal formulation comprising one or more binding partners is provided. Liposomal formulations include but are not limited to liposomal nanoparticles.

In embodiments, an effective amount of one or more binding partners is administered to an individual in need thereof. In embodiments, an effective amount is an amount that reduces one or more signs or symptoms of a disease and/or reduces the severity of the disease. An effective amount may also inhibit or prevent the onset of a disease or a disease relapse. A precise dosage can be selected by the individual physician in view of the patient to be treated. Dosage and administration can be adjusted to provide sufficient levels of binding partner to maintain the desired effect. Additional factors that may be taken into account include the severity and type of the disease state, age, weight, and gender of the patient, desired duration of treatment, method of administration, time and frequency of administration, drug combination(s), reaction sensitivities, and/or tolerance/response to therapy.

Binding partners and pharmaceutical compositions comprising the binding partners can be administered to an individual in need thereof using any suitable route, examples of which include intravenous, intramuscular, intraperitoneal, intracerebrospinal, subcutaneous, intra-articular, intrasynovial, oral, topical, or inhalation routes, depending on the particular condition being treated. The compositions may be administered parenterally or enterically. The compositions may be introduced as a single administration or as multiple

administrations or may be introduced in a continuous manner over a period of time. For example, the administration(s) can be a pre-specified number of administrations or daily, weekly, or monthly administrations, which may be continuous or intermittent, as may be therapeutically indicated.

In embodiments, the individual in need of a composition of this disclosure has been diagnosed with or is suspected of having cancer. In embodiments, the cancer is a solid tumor or a hematologic malignancy. In embodiments, the cancer is renal cell carcinoma, breast cancer, prostate cancer, pancreatic cancer, lung cancer, liver cancer, ovarian cancer, cervical cancer, colon cancer, esophageal cancer, glioma, glioblastoma or another brain cancer, stomach cancer, bladder cancer, testicular cancer, head and neck cancer, melanoma or another skin cancer, any sarcoma, including but not limited to fibrosarcoma, angiosarcoma, osteosarcoma, and rhabdomyosarcoma, and any blood cancer, including all types of leukemia, lymphoma, and myeloma. In embodiments, the individual is in need of treatment for any pre-neoplastic disorder, including myelodysplastic syndromes or myeloproliferative neoplasms. In embodiments, a described binding partner is used prophylactically for any of the described types of cancer.

In embodiments, administering one or more binding partners, including but not necessarily in a pharmaceutical formulation, to an individual in need thereof, exhibits an improved activity relative to a control. In an embodiment, the control comprises different antibodies, a different form of the same antibodies/binding partner, or antibodies/binding partners that are delivered without adding additional agents. In embodiments, a binding partner described herein provides for improved antibody dependent cell cytotoxicity (ADCC), or for internalization (such as for an ADC), relative to a control. In embodiments, a control protein or peptide does not comprise the covalently linked molecule. The control peptide may comprise the same sequence as the experimental peptide, or if the experimental peptide comprises a mutation the control peptide may comprise the wild type sequence.

A composition of this disclosure, such as a pharmaceutical formulation, can contain only one, or more than one binding partner, and thus combinations of different binding partners are included. Likewise, one or more binding partners can be combined with any other therapeutic agent, non-limiting examples of which include conventional chemotherapeutic agents, and modulators of T-cell costimulatory molecules, often referred to as immune checkpoint inhibitors. T-cell costimulatory molecules are known in the art (PMID 30115704), including, but not limited to, CTLA4, PD-1, PD-L1, LAG3, TIM3, TIGIT, VISTA, B7-1, B7-2, PD-L2, LSECTin, Galectin-9, CEACAM-1, CD155, CD112, CD28, ICOS, ICOSL, OX40, OX40L, GITR, GITRL, 4-1BB, 4-1BBL, CD40, CD40L, CD27, and CD70. Thus, the disclosure includes combination therapy using one or more described binding partners and any of modulators of T-cell costimulatory molecules, including but not limited to CTLA-4 inhibitors, PD-1 inhibitors and PD-L1 inhibitors. As non-limiting examples, anti-PD-1 agents include Pembrolizumab and Nivolumab. Anti-PD-L1 examples include Avelumab and Atezolizumab. An anti-CTLA-4 example is Ipilimumab. The binding partners may also be combined with any form of adoptive immunotherapy.

In embodiments, the disclosure comprises administering to an individual in need thereof one or more binding partners and at least one additional agent to provide an additive effect, or a greater than additive effect such as a synergistic result. In embodiments, the described effect comprises inhibi-

bition of cancer growth, inhibition of metastasis, or other beneficial effect. An additive effect or synergistic effect may also be achieved by using a combination of at least two described binding partners.

Various techniques have been developed for the production of binding partners and are included in the scope of this disclosure. In embodiments, the binding partners are produced by host cells by way of recombinant expression vectors. The present disclosure includes all polynucleotide sequences encoding the amino acid sequences described herein, expression vectors comprising such polynucleotide sequences, and in vitro cell cultures comprising such expression vectors. In embodiments, the cell cultures include prokaryotic cells or eukaryotic cells. In embodiments, the cell cultures are mammalian cells. In embodiments, the cells are CHO cells. In embodiments, the cells are HEK293 cells and their derivatives. Kits comprising the binding partners, and/or cell cultures expressing the binding partners, are provided by this disclosure. In general, the kits comprise one or more sealed containers that contain the binding partners, or cells expressing them. Instructions for using the binding partners for therapeutic and/or diagnostic purposes can be included in the kits.

Cells that are modified to express any described binding partner include but are not necessarily limited CD4+ T cells, CD8+ T cells, Natural Killer T cells, $\gamma\delta$ T cells, and cells that are progenitors of T cells, such as hematopoietic stem cells or other lymphoid progenitor cells, such as immature thymocytes (double-negative CD4-CD8-) cells, or double-positive thymocytes (CD4+CD8+). In embodiments, the progenitor cells comprise markers, such as CD34, CD117 (c-kit) and CD90 (Thy-1). In embodiments, the modified cells comprise macrophages. The described modified cells may be used therapeutically or prophylactically.

In embodiments, the disclosure provides for generation of a binding partner. This approach comprises providing a plurality of distinct binding partners, exposing the plurality of distinct (e.g., different) binding partners to one or a diversity of peptide conjugates, and selecting binding partners that bind with specificity to the peptide conjugates that contain the covalently conjugated drug or other molecule, but do not bind to the protein or peptide that does not comprise the covalently conjugated drug or other molecule. As described above, this approach can be performed on a manner that either does, or does not, require the amino acid sequence of the protein or peptide to be part of the antigenic determinant. The described approach can be used to select binding partners that are specific for presentation of a peptide conjugate as a component of any MHC complex.

In embodiments, binding partners described herein and as otherwise will be apparent by those skilled in the art, can be used to determine whether or not a particular drug or other molecule forms a covalent interaction with a protein or peptide. Thus, the disclosure provides for exposing protein or peptide substrates to drug candidates and using the binding partners described herein or as identified as described herein to determine whether or not the drug forms a covalent interaction with the pertinent substrate. This determination can be made based on whether or not the binding partner binds to the protein or peptide that has been covalently attached to the drug. This approach can be used in lieu of currently available techniques, such as mass spectroscopy and the like.

In embodiments, binding partners of this disclosure may be used in any immunological diagnostic test, including but not limited to the imaging approaches described above. In embodiments, one or more binding partners described herein

can be used as a component in any form of, for example, enzyme-linked immunosorbent assay (ELISA) assay, including but not limited to a direct ELISA, a sandwich ELISA, a competitive ELISA, and a reverse ELISA. In embodiments, one or more binding partners described herein can also be incorporated into an immunodiagnostic device, such as a microfluidic device, a lateral flow device, and the like. The binding partners may also be used in, for example, Western blots and immunoprecipitation assays.

The following Examples are intended to illustrate but not limit the disclosure. In embodiments, antibodies described in Example 3 have different properties relative to those described in Example 1. Other differences between binding partners will be apparent from the Examples and their accompanying figures. The different properties include, but are not necessarily limited to, specificity for a drug conjugate displayed in the context of a specific MHC type. Thus, binding partners may exhibit different binding partners when a peptide conjugates is in a particular MHC complex.

Example 1

This Example provides a description of the identification and characterization of binding partners that bind with specificity to ARS-1620, which forms a covalent interaction with KRAS^{G12C}.

In particular, FIG. 1 demonstrates phage ELISA of phage-displayed antibody clones.

Binding to KRAS^{G12C}-GDP and KRAS^{G12C}-GDP-ARS-1620 conjugate was determined. Buffer denotes binding signal to the wells that did not contain KRAS^{G12C}. From these candidates, four different antibodies were identified. Among these, 12C-ARS-Fab59 showed high affinity binding to KRAS^{G12C}-GDP covalently bound to ARS-1620. The results are presented in FIG. 2, which shows 12C-ARS Fab59 binding to KRAS^{G12C} in the GTP γ S- or GDP-bound nucleotide state with or without ARS-1620, as characterized by the bead binding assay (PMID: 33358997). FIG. 3 also demonstrates that 12C-ARS-Fab59 specifically binds KRAS^{G12C}-GDP conjugated to ARS-1620. In particular, 12C-ARS-Fab binding to KRAS^{G12C} (left) or WT RAS isoforms (right) in the GTP γ S- or GDP-bound nucleotide state with or without ARS-1620 conjugation, is shown as indicated.

FIG. 4 demonstrates the use of 12C-ARS-Fab59 to measure ARS-1620/KRAS^{G12C} adducts by pull-down assays from lysates prepared from cell lines. To produce the data shown in FIG. 4, immunoblots were performed on whole cell lysates and 12C-ARS Fab-pull-downs (PD) from RAS-less MEFs reconstituted with the indicated KRAS mutants (4A) and from KCP (Kras^{G12C}; Tp53^{R172H}; Pdx-Cre) mouse pancreas cancer cells (4B), treated in the presence or absence of ARS-1620. FIG. 4C shows whole cell lysates and 12C-ARS Fab pull-downs (PD) from H358 and MIAPaCa-2 cells, treated as indicated, which were subjected to SDS-PAGE and immunoblotting with anti-pan RAS and anti-ERK2 antibodies, the latter as a loading control. FIG. 4D shows ARS-adduct formation in samples from 4C, quantified by LC/MS-MS assay. ARS-1620 and SHP099 concentrations were 10 μ M in all panels.

FIG. 5 shows that 12C-ARS-Fab59 can be used to measure the engagement of ARS-1620 to mutant KRAS by pull-down assay with lysates prepared from animal tissues. In particular, in FIGS. 5A-B, anti-pan RAS and anti-ERK2 (loading control) immunoblots of lysates and 12C-ARS Fab pull-downs (PD) from LSL-KRAS^{G12C};Tp53^{R270H} (A) and LSL-KRAS^{G12C} (B) tumors after 3 days of oral gavage with

ARS-1620 (200 mg/kg/d) alone or with the SHP2 inhibitor SHP099 (75 mg/kg/d) are shown.

To produce the foregoing results, the following materials and methods were used.

RAS Nucleotide Exchange and Generation of ARS-1620-Conjugated RAS

Purified RAS (1-174) proteins containing a 6xHIS-tag (SEQ ID NO: 13) and an AVI-tag (1), used in the binding experiments and phage display selections, were prepared by diluting stock protein (typically containing 20-100 μ M RAS) 25-fold with 20 mM Tris-Cl buffer pH 7.5 containing 5 mM EDTA, 0.1 mM DTT, and 1 mM (final concentration) of nucleotide (GDP or GTP γ S). For generating ARS-bound RAS, ARS-1620 (final concentration: 100 μ M) was added during the nucleotide exchange reaction of RAS along with GDP. Samples were incubated at 30° C. for 30 minutes. MgCl₂ was then added to a final concentration of 20 mM to quench the nucleotide exchange reaction, and the solution was incubated on ice for at least 5 minutes prior to use.

Selection of Phage-Displayed Antibody Fragments Against ARS-Bound KRAS^{G12C}

General procedures for the development of Fabs against purified protein targets have been described (2). Four rounds of phage display library selection were performed, with biotinylated KRAS(G12C)-GDP+ARS-1620 at 100 nM, 100 nM, 50 nM, and 20 nM in the first, second, third and fourth rounds, respectively. The first round recovered clones that bound to KRAS^{G12C}-GDP+ARS-1620; the second round recovered clones that bound to KRAS^{G12C}-GDP+ARS-1620, previously pre-cleared with KRAS^{G12C}-GDP; the third round recovered clones that bound to KRAS^{G12C}-GDP+ARS-1620, previously pre-cleared with KRAS^{G12C}-GTP. The final round recovered clones that bound to KRAS^{G12C}-GDP+ARS-1620, previously pre-cleared with KRAS^{G12C}-GDP. Phage captured on beads were eluted in 100 μ l of 0.1 M Gly-HCl (pH 2.1) and immediately neutralized with 35 μ l of 1M Tris-Cl (pH 8). Recovered clones were analyzed by phage ELISA and DNA sequencing, as described (2).

Bead Binding Assays

General methods have been previously described (3). Fifty microliters (50 μ l) of M280 streptavidin beads (Thermo Fisher) were incubated with 100 μ l of biotinylated 12C-ARS Fab, at 30 nM or 4 nM. Ligand-free streptavidin on the beads was then blocked by adding excess biotin. Beads were washed with supplemented TBST (50 mM Tris pH7.5, 150 mM NaCl, 20 mM MgCl₂, 0.1 mM DTT, 0.05% Tween-20) and dispensed into wells of a 96-well U bottom plate (Greiner). Beads were then incubated at 1:1 ratio with purified RAS proteins diluted in supplemented TBS (50 mM Tris pH7.5, 150 mM NaCl, 20 mM MgCl₂, 0.1 mM DTT) at 2 \times the concentration stated for the titration curve for 30 minutes at room temperature. Beads containing bound Fab and RAS were transferred to the wells of a 96-well filter plate (Millipore, MSHVN4550) and washed twice with supplemented TBST before incubating with Neutravidin-Dylight 650 (Thermo Fisher Scientific) for 30 minutes at 4° C. The beads were washed twice with supplemented TBST before resuspension in supplemented TBS for flow cytometry using an iQue screener (Sartorius). The median signal intensity in the Dylight650 channel for the 75-95th percent-

-continued

 V_H :

(SEQ ID NO: 21)
**EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSIHWWVRQAPGKGLEWVA
 SISSSSGTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
 SPYVYYWYMVGFDYWGQGTLVTVSS**
 (Bolted CDR sequences are disclosed as SEQ ID NOS 185, 170, and 186, respectively, in order of appearance)

This reference listing pertains to Example 1.

1. Spencer-Smith R, Koide A, Zhou Y, Eguchi R R, Sha F, Gajwani P, Santana D, Gupta A, Jacobs M, Herrero-Garcia E, Cobbert J, Lavoie H, Smith M, Rajakulendran T, Dowdell E, Okur M N, Dementieva I, Sicheri F, Therrien M, Hancock J F, Ikura M, Koide S, O'Bryan J P. Inhibition of RAS function through targeting an allosteric regulatory site. *Nat Chem Biol.* 2017; 13 (1): 62-8. doi: 10.1038/nchembio.2231. PubMed PMID: 27820802; PMCID: 5193369.
2. Fellouse F A, Esaki K, Birtalan S, Raptis D, Cancasci V J, Koide A, Jhurani P, Vasser M, Wiesmann C, Kossiakoff A A, Koide S, Sidhu SS. High-throughput generation of synthetic antibodies from highly functional minimalist phage-displayed libraries. *J Mol Biol.* 2007; 373 (4): 924-40. Epub 2007 Sep. 11. doi: 10.1016/j.jmb.2007.08.005. PubMed PMID: 17825836.
3. Nishikori S, Hattori T, Fuchs S M, Yasui N, Wojcik J, Koide A, Strahl B D, Koide S. Broad ranges of affinity and specificity of anti-histone antibodies revealed by a quantitative Peptide immunoprecipitation assay. *J Mol Biol.* 2012; 424 (5): 391-9. Epub 2012 Oct. 9. doi: 10.1016/j.jmb.2012.09.022. PubMed PMID: 23041298; PMCID: 3502729.
4. Patricelli M P, Janes M R, Li L S, Hansen R, Peters U, Kessler L V, Chen Y, Kucharski J M, Feng J, Ely T, Chen J H, Firdaus S J, Babbar A, Ren P, Liu Y. Selective Inhibition of Oncogenic KRAS Output with Small Molecules Targeting the Inactive State. *Cancer Discov.* 2016; 6 (3): 316-29. Epub 2016 Jan. 8. doi: 10.1158/2159-8290.CD-15-1105. PubMed PMID: 26739882. 5. doi.org/10.1084/jem.20201414

Example 2

This Example provides a description of binding partners that bind with specificity to AMG510 that is covalently linked to peptides.

To produce the results described in this Example, some methods as described in Example 1 were adapted. For this Example, AMG510 (purchased from Selleckchem) was conjugated to a peptide corresponding to KRAS(G12C) residues 4-18:

(SEQ ID NO: 22)
 H2N-YKLVVVGACGVGKSA (dPEG4) (K-long chain Biotin)-amide

and a poly-Ser peptide containing a central Cys:

(SEQ ID NO: 23)
 H2N-SSSSCSSSSW (K-long chain Biotin)-amide.

A human single-chain Fv yeast-display library was sorted using these peptides as targets by using established methods (1-3). After rounds of library sorting, individual clones were screened. We developed three antibodies that bound to AMG510 conjugated to both KRAS(G12C) and poly-Ser

peptide (FIG. 6). Consequently, these antibodies recognize predominantly the AMG510 moiety but not the peptide moiety of the conjugates. Additionally, we developed other clones that are selective to AMG510 conjugated to the KRAS(G12C) peptide.

One such clone, P2AMR-1 was then produced in the format of human IgG1 and further characterized. It bound to AMG510 conjugated to the KRAS(G12C) peptide with high apparent affinity in a bead binding assay (FIG. 7)(4). The antibody clone also bound tightly to AMG510 conjugated a shorter KRAS(G12C) peptide, VVGACGVGK (SEQ ID NO: 1), in the context of HLA-A*03:01 (BioLegend Flex-T) (FIG. 8).

P2AMR-1 detected AMG510 conjugated to KRAS (G12C) peptide that had been added to Raji cells, which are known to express HLA-A*03:01. By contrast, P2AMR-1 did not detect KRAS(wild type) peptide loaded in the same manner (FIG. 16). In addition, P2AMR-1 did not bind to AMG510 conjugated to the KRAS(G12C) peptide added to cells that are not known to express HLA-A*03:01, e.g., MV4-11 and Expi293 cells (FIG. 16).

These results demonstrate that the presently provided antibodies, which represent binding partners of this disclosure, recognize the AMG510 moiety in a manner agnostic of the conjugation partner, and they suggest that our antibodies and their derivatives can be used to identify cells that present AMG510-KRAS (G12C) peptide conjugate on MHC molecules on the cell surface.

More generally, these results suggest methods for targeting any cells that harbor intracellular targets that form covalent adducts with small molecule ligands.

This example demonstrates the following non-limiting binding partners, restricted to AMG510 covalent modifications of the described substrates. CDR residues (Kabat scheme) are shown in bold.

P2AMR-1

 V_L :

(SEQ ID NO: 24)
 QSVLIQPRSVSGSPGQSVTISCT**GTSSDVGGNYVS**WYQQHPGKAPKLM

IYDVSKRPSGVPDFRGSKSGNTASLTVSGLQAEDEADYYCGSYADTD

IVFGTGTKLTVL

(Bolted CDR sequences are disclosed as SEQ ID NOS 187, 188, and 189, respectively, in order of appearance)

 V_H :

(SEQ ID NO: 25)
 QVQLVQSEPEVKPGSSVKLSCKASGGTFSTD**AITWVRQAPGQGLEYMG**

GIIPLLDSVDYAQRFQGRVTVSADKSTGTAYMEVRSLGSEDTAKYYCAK

WSSVDTGLDYGQGTLTVVSS

(Bolted CDR sequences are disclosed as SEQ ID NOS 190, 191, and 192, respectively, in order of appearance)

P2AMR-12 (this clone has only the heavy chain)

 V_H :

(SEQ ID NO: 26)
 QVQLQESGPGLVKPSETLSLTCTVSGDSII**NDPHYWGIRQSPGKGLEW**

IGSTSHSGHTYFNPSLKSRSVMSIDVAKNQFSLNVRVSFTAADTAVYYCA

RMRRYYSGTYPVYYFDYWGQGTLTVSS

(Bolted CDR sequences are disclosed as SEQ ID NOS 193, 194, and 195, respectively, in order of appearance)

25

-continued

P2AMR-13

V_L:(SEQ ID NO: 27)
SSVLTQPPSASGTPGQRVTISCGSSSNIGSNFWSWYQQLPGTAKLII

SSNNQRPSGVPDFSGSKSDTSASLAISGLQSEDEADYYCAAWDDSLNG

PVFGGGTQLTVAL

(Bolted CDR sequences are disclosed as SEQ ID NOS 196, 197, and 198, respectively, in order of appearance)

V_H:(SEQ ID NO: 28)
QVQLVQSEAEVKPGSSVKVSKASGGTFSRYGVSVWRQAPGQGLEWMG

GIIPMFGTANYAQKFQGRVTITADESTSTAYMELRSLRSEDTAVYYCAR

GDNSAYSDAFNIWGQGTMVTVVSS

(Bolted CDR sequences are disclosed as SEQ ID NOS 199, 200, and 201, respectively, in order of appearance)

This reference listing pertains to Example 2.

1. Chao G, Lau W L, Hackel B J, Sazinsky S L, Lippow S M, Wittrup K D. Isolating and engineering human antibodies using yeast surface display. Nat Protoc. 2006; 1 (2): 755-68. Epub 2007 Apr. 5. doi: 10.1038/nprot.2006.94. PubMed PMID: 17406305.
2. Feldhaus M J, Siegel R W, Opresko L K, Coleman J R, Feldhaus J M, Yeung Y A, Cochran J R, Heinzelman P, Colby D, Swers J, Graff C, Wiley H S, Wittrup K D. Flow-cytometric isolation of human antibodies from a nonimmune *Saccharomyces cerevisiae* surface display library. Nat Biotechnol. 2003; 21 (2): 163-70. Epub 2003 Jan. 22. doi: 10.1038/nbt785nbt785 [pii]. PubMed PMID: 12536217.
3. Hattori T, Taft J M, Swist K M, Luo H, Witt H, Slattery M, Koide A, Ruthenburg A J, Krajewski K, Strahl B D, White K P, Farnham P J, Zhao Y, Koide S. Recombinant antibodies to histone post-translational modifications. Nat Methods. 2013; 10 (10): 992-5. doi: 10.1038/nmeth.2605. PubMed PMID: 23955773; PMCID: 3828030.
4. Nishikori S, Hattori T, Fuchs S M, Yasui N, Wojcik J, Koide A, Strahl B D, Koide S. Broad ranges of affinity and specificity of anti-histone antibodies revealed by a quantitative Peptide immunoprecipitation assay. J Mol Biol. 2012; 424 (5): 391-9. Epub 2012 Oct. 9. doi: 10.1016/j.jmb.2012.09.022. PubMed PMID: 23041298; PMCID: 3502729.

Example 3

This Example describes antibodies that bind to peptide-drug conjugates, but only in the context of specific MHC display of the described peptide-drug conjugates. The antibodies were produced as follows.

Antigen Preparation

KRAS(G12C) peptides ((H2N-VVGACGVGK-OH (SEQ ID NO: 1) and H2N-VVGACGVGK-OH (SEQ ID NO: 2)) were reacted with AMG510 (Selleckchem) and loaded onto Flex-T HLA-A*03:01 and Flex-T HLA-A*11:01 (produced by Biolegend), or onto HLA-A*03:01 and HLA-A*11:01 produced in house. KRAS(WT) peptide ((H2N-VVGAGGVGK-OH (SEQ ID NO: 29)) was loaded onto the HLA molecules in the same manner. EGFR peptide (H2N-QLMPFGCLL-OH (SEQ ID NO: 30)) was reacted with Osimertinib (Selleckchem) and loaded onto Flex-T

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HLA-A*02:01 or HLA-A*02:01 produced in house. As a control, the same peptide was reacted with beta-mercaptoethanol and loaded onto the HLA molecule. BTK peptide (H2N-YMANGLNY-OH (SEQ ID NO: 31)) was reacted with Ibrutinib (Selleckchem) and loaded onto Flex-T HLA-A*01:01 or HLA-A*01:01 produced in house. As a control, the same peptide was reacted with beta-mercaptoethanol and loaded onto the HLA molecule. The peptide-loaded HLA mixtures prepared with Flex-T HLA proteins were used without further purification. The peptide-loaded HLA mixtures prepared with HLA samples prepared in house were further purified using size-exclusion chromatography with a Superdex S200 column.

15 Antibody Phage-Display Library Sorting

Sorting of an antibody phage-display library was performed as described previously⁽¹⁾. Briefly, a phage-display library was first sorted with all four antigens at 100 nM in the first round, followed by sorting with a single antigen at 100, 50, and 20 nM in the second, third, and fourth rounds, respectively. To enrich for clones with the desired specificity, counterselection was performed using KRAS(WT) peptide-loaded MHC molecules or beta-mercaptoethanol-treated peptide-loaded MHC molecules in the second, third, and fourth rounds.

Binding of individual phage clones were tested using the multiplex bead binding assay⁽²⁾.

30 Antibody Yeast-Display Library Sorting and Clone Characterization

Display of antibody clones in the form of single-chain Fv (scFv) on the yeast surface, library sorting using fluorescence-activated cell sorting, and characterization of individual clones were performed essentially as described previously (Hattori et al. PMID 23955773; Cao et al. PMID 17406305).

40 Deep Mutational Scanning

Deep mutational scanning was performed following general procedures published previously (PMID 32841599). A yeast-display library, in the scFv format, contained variants in which a single position was diversified with the NNK codon. A yeast display library was subjected to FACS using an antigen of interest to enrich a pool of clones that bound the antigen and a pool of clones that did not bind the antigen. The DNA sequences of the enriched pools were determined, and amino acid substitutions were deduced.

50 Antibody Production

The genes encoding selected antibody clones were transferred from the phage-display vector to IgG expression vectors (pFUSEss-CHIg-hG1 and pFUSE2ss-CHIg-hK, InvivoGen), and IgG proteins were produced using the ExpiCHO cell line (Thermo Fisher) and purified using a Protein Capture Devices with Protein A (GORE).

60 Data presented in this Example relates to FIGS. 9-15, 17, which provide the following information:

FIG. 9 provides a cartoon representation of a concept of the disclosure referred to as HapImmune. The numbers 1-7 denote relevant steps. 1. A covalent inhibitor is administered, and it enters the cell harboring the target protein. 2. The inhibitor binds the target and forms a covalent bond with the target. 3 and 4. As a part of natural protein turnover

(or induced protein degradation in the case of a PROTAC), the target-drug conjugate is degraded by the proteosome system. As a result, peptides with the conjugated drug are produced. 5. A peptide conjugate is incorporated into a compatible MHC molecule. 6. The MHC/peptide-drug conjugate complex translocates to the surface of the cell. A HapImmune antibody recognizes the complex. 7. The surface bound antibody recruits an immune effector cell, such as an NK cell, which in turn initiates cell killing activities. Multiple modalities are envisioned for effecting cell killing activities, including ADCC, ADCP, CDC, BITE, CAR-T, CAR-NK, ADC, and radioisotope conjugate, but they are not explicitly depicted here.

FIG. 10 shows data from development of antibodies that bind MHC/peptide-drug conjugate complexes. (A) Multiplex bead-binding assay (MBBA)⁽¹⁾ of phages displaying different antibody clones. For each phage clone, binding to a total of five antigens presented on beads was tested: HLA-A*03:01 in complex with the KRAS(G12C) peptide conjugated with AMG510 (denoted as HLA-A*03:01_RAS-AMG510 in the figure); HLA-A*03:01 in complex with the KRAS(wild type) peptide (denoted as HLA-A*03:01_WTRAS); HLA-A*11:01 in complex with the KRAS (G12C) peptide conjugated with AMG510 (denoted as HLA-A*11:01_RAS-AMG510); HLA-A*11:01 in complex with the KRAS(wild type) peptide (denoted as HLA-A*11:01_WTRAS); beads presenting no antigen (denoted as No target). (B) MBBA assay of phages displaying different antibody clones to: HLA-A*01:01 in complex with the BTK peptide conjugated with Ibrutinib (denoted as HLA-A*01:01_BTK-Ibrutinib in the figure); HLA-A*01:01 in complex with the BTK peptide conjugated with beta-mercaptoethanol (denoted as HLA-A*01:01_BTK-βme); beads presenting no antigen (denoted as No target). (C) MBBA assay of phages displaying different antibody clones to: HLA-A*02:01 in complex with the EGFR peptide conjugated with Osimertinib (denoted as HLA-A*02:01_EGFR-Osimertinib in the figure); HLA-A*02:01 in complex with the EGFR peptide conjugated with beta-mercaptoethanol (denoted as HLA-A*02:01_EGFR-βme); beads presenting no antigen (denoted as No target).

FIG. 11 shows results from binding titration using the multiplex bead-binding assay (MBBA) of purified antibodies targeted to the KRAS(G12C)-AMG510 conjugate. Clone names are shown over each graph. Antigen nomenclature is described in FIG. 10. The left column shows binding data with HLA-A*03:01 complexes, whereas the right column shows data with HLA-A*11:01 complexes. Apparent dissociation constant (K_D) values were determined using nonlinear least-squared fitting of a 1:1 binding function. The data for the wild-type RAS peptide complexes and for the no target were all close to the baseline and overlap, and thus their apparent K_D values were not determined. Data shown here are from triplicate measurements. Error bars are within the size of the symbols.

FIG. 12 demonstrates that binding of antibodies to the drug-peptide conjugate in complex with an MHC was not affected by the presence of the free drug. MBBA binding signals of select "AMR" series antibodies to HLA-A*03:01 in complex with the KRAS(G12C) peptide conjugated with AMG510 in the absence (the white bars) and presence (the gray bars) of 10 μM free AMG510 are shown. The antibody concentrations were adjusted to give sub-saturating signals and are shown in parentheses. Data shown here are from triplicate measurements.

FIG. 13 shows results from binding titrations using the multiplex bead-binding assay (MBBA) of purified antibod-

ies targeted to the BTK-Ibrutinib conjugate. Clone names are shown over each graph. Antigen nomenclature is described in FIG. 10. Apparent dissociation constant (K_D) values were determined using nonlinear least-squared fitting of a 1:1 binding function. The data for the beta-mercaptoethanol-conjugated peptide in complex with HLA-A*01:01 and for the no target were all close to the baseline and overlap, and thus their apparent K_D values were not determined. Data shown here are from triplicate measurements. Error bars are within the size of the symbols.

FIG. 14 shows results from binding titrations of purified antibodies to the KRAS(G12C)-AMG510 conjugate presented by endogenous MHC molecules on the cell surface. Raji cells were first incubated with the KRAS(G12C)-AMG510 conjugate or the KRAS(wild type) peptide, and excess conjugate and peptide were washed away. Surface-bound antibody levels detected using a fluorescently labeled secondary antibody are shown as a function of IgG concentration used for staining. Apparent dissociation constant (K_D) values were determined using nonlinear least-squared fitting of a 1:1 binding function. Data shown here are from triplicate measurements.

FIG. 15 shows results from an antibody binding to a KRAS(G12C)-expressing cell line pretreated with AMG510. The non-small cell lung cancer cell line H358 was incubated with AMG510 for 2 days and then stained with antibodies targeting the KRAS G12C peptide-AMG510 conjugate or an isotype control, followed by detection with a secondary antibody. (A) Flow cytometry histograms. (B) Quantification of the median fluorescence intensity of H358 cells treated with or without AMG510. The antibodies used are indicated along the horizontal axis. (C) Quantification of the median fluorescence intensity of H358 cells and HEK293T cells (a negative control) treated with or without AMG510 and stained with the AMRA3-7 antibody.

FIG. 16 is related to Example 2 and shows binding of P2AMR-1 IgG to cells preincubated with the KRAS(G12C) peptide-AMG510 conjugate, KRAS(wild type) peptide, or no peptide. The antibody was precomplexed with a dye-labeled secondary antibody in order to enhance the effective binding (avidity). The antibody bound to the Raji cells that express HLA-A*03:01 when the cells were incubated with the conjugate.

FIG. 17 shows results from binding of purified antibodies in the IgG format to the indicated drug-peptide/MHC complexes as measured using the multiplex bead binding assay (MBBA).

FIGS. 18 and 19 are discussed in Example 4.

FIG. 20 shows results from binding titration curves of AMR-A3-7 and AMR-A3-7D displayed on the yeast surface. Binding to HLA-A*03:01 presenting AMG510 conjugated to the Cys residue in the 9mer and 10mer RAS (G12C) peptides, VVGACGVGK (SEQ ID NO: 1) and VVVGACGVGK (SEQ ID NO: 2), respectively, is shown.

FIGS. 23 and 24 show results from deep mutational scanning of CDR-L3 and CDR-H3 of AMR-A3-7D and OEA2-5, respectively.

Specific and non-limiting examples of antibody sequences that bind in an MHC-drug conjugate-specific manner are as follows:

Exemplary Antibody Clones Binding to KRAS(G12C)-AMG510 conjugate presented on HLA-A*03:01 and HLA-A*11:01. CDR residues (Kabat scheme) in bold.

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AMRA3-2
 V_L : (SEQ ID NO: 32)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQSGWSYPIT
FGQGTKVEIKRTV
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 202, respectively, in order of appearance)

V_H : (SEQ ID NO: 33)
EVQLVESGGGLVQPGGSLRLSCAASGFTFYSSYIHWRQAPGKGLEWVA
SISPYYGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
SSYYALDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 173, 203, and 204, respectively, in order of appearance)

AMRA3-7
 V_L : (SEQ ID NO: 34)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQISYVYSLI
TFQGTKVEIKRTV
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 205, respectively, in order of appearance)

V_H : (SEQ ID NO: 35)
EVQLVESGGGLVQPGGSLRLSCAASGFTFSSYIHWRQAPGKGLEWVA
SISSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
GGWPAMDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 206, 207, and 171, respectively, in order of appearance)

AMRA3-7KK
 V_L : (SEQ ID NO: 36)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQISYVKKLI
TFQGTKVEIKRTV
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 168, respectively, in order of appearance)

V_H : (SEQ ID NO: 37)
EVQLVESGGGLVQPGGSLRLSCAASGFTFSSYIHWRQAPGKGLEWVA
SISSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
GGWPAMDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 206, 207, and 171, respectively, in order of appearance)

AMRA3-7D
 V_L : (SEQ ID NO: 3)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQISYVKKLI
TFQGTKVEIKRTV
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 168, respectively, in order of appearance)

30

-continued
 V_H : (SEQ ID NO: 4)
EVQLVESGGGLVQPGGSLRLSCAASGFTFSYDYSIHWRQAPGKGLEWVA
SISSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
GGWPAMDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 169, 170, and 171, respectively, in order of appearance)

10 AMRA3-8
 V_L : (SEQ ID NO: 38)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDLATYYCQQYQYGYNLI
TFGQGTKVEIKRTV
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 208, respectively, in order of appearance)

20 V_H : (SEQ ID NO: 39)
EVQLVESGGGLVQPGGSLRLSCAASGFTISYSSSIHWRQAPGKGLEWVA
SIYSYSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
YSYGWVPGWRAIDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 210, and 211, respectively, in order of appearance)

30 AMRA3-11
 V_L : (SEQ ID NO: 40)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQSSSVYKLL
TFGQGTKVEIKRTV
(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 212, respectively, in order of appearance)

35 V_H : (SEQ ID NO: 41)
EVQLVESGGGLVQPGGSLRLSCAASGFTVYYSSSIHWRQAPGKGLEWVA
SISSSSYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTALYYCAR
GGPGWYRAMDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 213, and 214, respectively, in order of appearance)

45 AMRA3-15
 V_L : (SEQ ID NO: 42)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY
SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQSSSLITF
GQGTKVEIKRTV
55 (Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

50 V_H : (SEQ ID NO: 43)
EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSSSIHWRQAPGKGLEWVA
SISSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR
GYFYYGWAMAFDYWGQGTLVTVSS
(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 170, and 216, respectively, in order of appearance)

-continued

AMRA3-17

V_L:

(SEQ ID NO: 44)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSQWYEPLI

TFGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 217, respectively, in order of appearance)

V_H:

(SEQ ID NO: 45)

EVQLVESGGGLVQPGGSLRLSCAASGFTIYSSYIHWRQAPGKGLEWVA

SISSSSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

SYSYMSQWG WYQYS GMDYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 173, 170, and 218, respectively, in order of appearance)

AMRA3-18

V_L:

(SEQ ID NO: 46)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQGSYT YRLI

TFGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 219, respectively, in order of appearance)

V_H:

(SEQ ID NO: 47)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSYSSSIHWRQAPGKGLEWVA

SISSSSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YAWWAHGLDYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 220, and 221, respectively, in order of appearance)

AMRA3-21

V_L:

(SEQ ID NO: 48)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQASYWYNLF

TFGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 222, respectively, in order of appearance)

V_H:

(SEQ ID NO: 49)

EVQLVESGGGLVQPGGSLRLSCAASGFTIISYSSSIHWRQAPGKGLEWVA

SISSYSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

QYSMHPFWGYGM DYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 206, 207, and 223, respectively, in order of appearance)

AMRA3-22

V_L:

(SEQ ID NO: 50)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSDMPPITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 224, respectively, in order of appearance)

-continued

V_H:

(SEQ ID NO: 51)

EVQLVESGGGLVQPGGSLRLSCAASGFTFYSSSIHWRQAPGKGLEWVA

YIYSSSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

PVNYYYQGALDYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 225, and 226, respectively, in order of appearance)

10 AMRA3-23

V_L:

(SEQ ID NO: 52)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQYYFPITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 227, respectively, in order of appearance)

20

V_H:

(SEQ ID NO: 53)

EVQLVESGGGLVQPGGSLRLSCAASGFTVYSSSIHWRQAPGKGLEWVA

SISPSSGTYYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YHYMF EYDKGESKWGYGYCFDYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 228, and 229, respectively, in order of appearance)

30 AMRA11-1

V_L:

(SEQ ID NO: 54)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSQYFPITF

35 GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 230, respectively, in order of appearance)

V_H:

(SEQ ID NO: 55)

EVQLVESGGGLVQPGGSLRLSCAASGFTISYSSSIHWRQAPGKGLEWVA

SISSYSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

NSWSWYSGVGM DYWGQGTLTVSS

45 (Bolded CDR sequences are disclosed as SEQ ID NOS 209, 221, and 232, respectively, in order of appearance)

AMRA11-2

V_L:

(SEQ ID NO: 56)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSSSSLITF

GQGTKVEIKRTV

55 (Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 57)

EVQLVESGGGLVQPGGSLRLSCAASGFTI SSSSIHWRQAPGKGLEWVA

SISSYSSSTYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YPYGWG WGGSGLDYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 233, and 234, respectively, in order of appearance)

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-continued

AMRA11-15
VL:
(SEQ ID NO: 58)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQDFQYLIT

FGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 235, respectively, in order of appearance)

V_H:

(SEQ ID NO: 59)
EVQLVESGGGLVQPGGSLRLSCAASGFTVYSSIHWVRQAPGKGLEWVA

SISSYYGSTYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

GEK WALDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 236, and 237, respectively, in order of appearance)

AMRA11-16

VL:
(SEQ ID NO: 60)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQYMYQQPLI

TFCQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 238, respectively, in order of appearance)

V_H:

(SEQ ID NO: 61)
EVQLVESGGGLVQPGGSLRLSCAASGFTVYSSIHWVRQAPGKGLEWVA

SISSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

EPIYNWYGMWDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 170, and 239, respectively, in order of appearance)

AMRA311-2

VL:
(SEQ ID NO: 62)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQSLWWPITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 240, respectively, in order of appearance)

V_H:

(SEQ ID NO: 63)
EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSIHWVRQAPGKGLEWVA

SISSYSGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

HGSYGSWALDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 241, and 242, respectively, in order of appearance)

AMRA311-10

VL:
(SEQ ID NO: 64)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQYFYFPITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 243, respectively, in order of appearance)

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-continued

V_H:
(SEQ ID NO: 65)
EVQLVESGGGLVQPGGSLRLSCAASGFTFYSSSIHWVRQAPGKGLEWVA

SISSYYGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

ASYYSGYSSYPYMGLDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 244, and 245, respectively, in order of appearance)

10 **AMRA311-14**

V_L:
(SEQ ID NO: 66)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQGSYRNPLL

TFCQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 246, respectively, in order of appearance)

20 V_H:

(SEQ ID NO: 67)
EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSYIHWVRQAPGKGLEWVA

SISSSSGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

25 **MNWSHYAMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 206, 220, and 247, respectively, in order of appearance)

AMRA311-16

V_L:
(SEQ ID NO: 68)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQSSSSLITF

35 **GQGTKVEIKRTV**

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 69)
EVQLVESGGGLVQPGGSLRLSCAASGFTISSLSSSIHWVRQAPGKGLEWVA

YISSYSGYTYYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YWYGHYHSYFGLDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 248, and 249, respectively, in order of appearance)

AMRA311-17

V_L:
(SEQ ID NO: 70)
DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRSGSRSGTDFLTISLQPEDFATYYCQQSSSSLITF

45 **GQGTKVEIKRTV**

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 71)
EVQLVESGGGLVQPGGSLRLSCAASGFTISSLSSSIHWVRQAPGKGLEWVA

SISSYSGSTYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YPYGSHVYTGLDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 250, and 251, respectively, in order of appearance)

-continued

AMRA311-18

V_L:

(SEQ ID NO: 72)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISLQPEDFATYYCQQWNWADYLV

TFGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 252, respectively, in order of appearance)

V_H:

(SEQ ID NO: 73)

EVQLVESGGGLVQPGGSLRLSCAASGFTISSLQVQAPGKGLEWVA

SIYSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**VISSRYWGWGVAFDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 253, and 254, respectively, in order of appearance)

AMRA311-19

V_L:

(SEQ ID NO: 74)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISLQPEDFATYYCQQYWWYSLIT

FGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 255, respectively, in order of appearance)

V_H:

(SEQ ID NO: 75)

EVQLVESGGGLVQPGGSLRLSCAASGFTVYSSSIHWVRQAPGKGLEWVA

YIYSSSGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

RSFPQWYNGSYTPWPAMDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 225, and 256, respectively, in order of appearance)

AMRA311-20

V_L:

(SEQ ID NO: 76)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISLQPEDFATYYCQQYMWYPVTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 257, respectively, in order of appearance)

V_H:

(SEQ ID NO: 77)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSIHWVRQAPGKGLEWVA

SIYSYSSYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**PFYWGERYALDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 258, and 259, respectively, in order of appearance)

Antibody clones that bind preferentially to KRAS (G12C)-AMG510 conjugate presented on HLA-A*03:01 relative to the same conjugate presented on HLA-A*11:01.

CDR residues (Kabat scheme) in bold.

AMRA3-5

V_L:

(SEQ ID NO: 78)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISLQPEDFATYYCQQSYSTLVTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 260, respectively, in order of appearance)

V_H:

(SEQ ID NO: 79)

EVQLVESGGGLVQPGGSLRLSCAASGFTFYSSSIHWVRQAPGKGLEWVA

SIYSSYGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**IYGWSYQWGAGMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 261, and 262, respectively, in order of appearance)

AMRA3-6

V_L:

(SEQ ID NO: 80)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISLQPEDFATYYCQQSSSSLITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

30 *V_H:*

(SEQ ID NO: 81)

EVQLVESGGGLVQPGGSLRLSCAASGFTISYSSSIHWVRQAPGKGLEWVA

SIYPPYGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**GGDYYWGWYWVAMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 263, and 264, respectively, in order of appearance)

AMRA3-9

V_L:

(SEQ ID NO: 82)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTIXSLQPEDFATYYCQKSSSSLITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 265, respectively, in order of appearance)

35 *V_H:*

(SEQ ID NO: 83)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSYIHWRQAPGKGLEWVA

SISSSYGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**MYYYTYPGMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 173, 266, and 267, respectively, in order of appearance)

AMRA3-10

V_L:

(SEQ ID NO: 84)

60 DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFTLTISLQPEDFATYYCQKGSSYLLTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 268, respectively, in order of appearance)

-continued

 V_H :

(SEQ ID NO: 85)

EVQLVESGGGLVQPGGSLRLSCAASGFTIYSYSSIHWVRQAPGKGLEWVA

SISPSSGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**HYGGWSHYMSGMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 206, 269, and 270, respectively, in order of appearance)

AMRA3-13

 V_L :

(SEQ ID NO: 86)

DIQMTQSPSSLSASVGDRVITIC**RASQSVSSAVAWYQQKPGKAPKLLIY****SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQNYYYYHKLI****TFGQGTKVEIKRTV**

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 271, respectively, in order of appearance)

 V_H :

(SEQ ID NO: 87)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSYSSIHWVRQAPGKGLEWVA

SISSSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**GRYGGMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 272, and 273, respectively, in order of appearance)

AMRA3-19

 V_L :

(SEQ ID NO: 88)

DIQMTQSPSSLSASVGDRVITIC**RASQSVSSAVAWYQQKPGKAPKLLIY****SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQLSYYVKLI****TFGQGTKVEIKRTV**

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 274, respectively, in order of appearance)

 V_H :

(SEQ ID NO: 89)

EVQLVESGGGLVQPGGSLRLSCAASGFTFYSSSIHWRQAPGKGLEWVA

SISSSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**GWYKAMDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 272, and 275, respectively, in order of appearance)

AMRA3-24

 V_L :

(SEQ ID NO: 90)

DIQMTQSPSSLSASVGDRVITIC**RASQSVSSAVAWYQQKPGKAPKLLIY****SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSSSSLITE****GQGTKVEIKRTV**

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

 V_H :

(SEQ ID NO: 91)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSYSSSIHWRQAPGKGLEWVA

SISSSYGYSTYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR**MYYYYYPGIDYWGQGTLVTVSS**

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 276, and 277, respectively, in order of appearance)

Antibody clones that bind preferentially to KRAS(G12C)-AMG510 conjugate presented on HLA-A*11:01 relative to KRAS(G12C)-AMG510 conjugate presented on HLA-A*03:01.

5

AMRA11-3

 V_L :

(SEQ ID NO: 92)

DIQMTQSPSSLSASVGDRVITIC**RASQSVSSAVAWYQQKPGKAPKLLIY**

10

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDLATYYCQQYYYFPITF

15

GQGTTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 278, respectively, in order of appearance)

 V_H :

(SEQ ID NO: 93)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSIHWRQAPGKGLEWVA

20

SISPYYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

25

SPYYWYQYFWGGLDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 279, and 280, respectively, in order of appearance)

30

AMRA11-4

 V_L :

(SEQ ID NO: 94)

DIQMTQSPSSLSASVGDRVITIC**RASQSVSSAVAWYQQKPGKAPKLLIY**

35

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSSSSLITE

40

GQGTTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

45

 V_H :

(SEQ ID NO: 95)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSYSSSIHWRQAPGKGLEWVA

50

SISSSGTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

55

SPYWNNMSAMDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 170, and 281, respectively, in order of appearance)

60

AMRA11-7

 V_L :

(SEQ ID NO: 96)

DIQMTQSPSSLSASVGDRVITIC**RASQSVSSAVAWYQQKPGKAPKLLIY**

65

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQGWWNPFTF

70

GQGTTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 282, respectively, in order of appearance)

75

 V_H :

(SEQ ID NO: 97)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSYSIHWRQAPGKGLEWVA

80

SISPYYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

85

WSWQYYSGHSSWGLDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 206, 279, and 283, respectively, in order of appearance)

-continued

AMRA11-8

V_L:

(SEQ ID NO: 98)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQSWYFPPLTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 284, respectively, in order of appearance)

V_H:

(SEQ ID NO: 99)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSSIHWWVRQAPGKGLEWVA

SIYSYYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WYNEYHHDDYWDAMDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 231, and 285, respectively, in order of appearance)

AMRA11-9

V_L:

(SEQ ID NO: 100)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQQSSSLITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 101)

EVQLVESGGGLVQPGGSLRLSCAASGFTLYYSSIHWWVRQAPGKGLEWVA

SISSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WMYWWWSFALDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 170, and 286, respectively, in order of appearance)

AMRA11-10

V_L:

(SEQ ID NO: 102)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQSYLWPITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 287, respectively, in order of appearance)

V_H:

(SEQ ID NO: 103)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSSIHWWVRQAPGKGLEWVA

SIYSYYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WQYHYNWYGMWDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 231, and 288, respectively, in order of appearance)

AMRA11-11

V_L:

(SEQ ID NO: 104)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQYPMSLITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 289, respectively, in order of appearance)

-continued

V_H:

(SEQ ID NO: 105)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSYSSIHWWVRQAPGKGLEWVA

SISPYSGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

GYDDYAGLDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 290, and 291, respectively, in order of appearance)

AMRA11-12

V_L:

(SEQ ID NO: 106)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQYYYFPITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 278, respectively, in order of appearance)

20

V_H:

(SEQ ID NO: 107)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSYSSIHWWVRQAPGKGLEWVA

SISPYYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WESEYSGTYEDYWGMDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 292, 279, and 293, respectively, in order of appearance)

30 AMRA11-13

V_L:

(SEQ ID NO: 108)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQYMWWPITF

35 GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 294, respectively, in order of appearance)

V_H:

(SEQ ID NO: 109)

EVQLVESGGGLVQPGGSLRLSCAASGFTISYSSIHWWVRQAPGKGLEWVA

SISSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

TGYWQGYLALDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 213, and 295, respectively, in order of appearance)

45 AMRA11-14

V_L:

(SEQ ID NO: 110)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSGRSGTDFLTISLQPEDFATYYCQQSSSSLITF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 111)

EVQLVESGGGLVQPGGSLRLSCAASGFTISYSSIHWWVRQAPGKGLEWVA

SISSSSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YYYYWNSTPAMDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 209, 170, and 296, respectively, in order of appearance)

60

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41

-continued

AMRA11-18

V_L:

(SEQ ID NO: 112)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSYGYPVTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 297, respectively, in order of appearance)

V_H:

(SEQ ID NO: 113)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSIHWVRQAPGKGLEWVA

SISSSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WYNSSWYSSNWWYKGFGMDYWQGQTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 272, and 298, respectively, in order of appearance)

AMRA11-20

V_L:

(SEQ ID NO: 114)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQYSSLLTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 299, respectively, in order of appearance)

V_H:

(SEQ ID NO: 115)

EVQLVESGGGLVQPGGSLRLSCAASGFTFYSSSIHWVRQAPGKGLEWVA

SISSSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

TSYTYPVITYYGFDYWGQGTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 272, and 300, respectively, in order of appearance)

AMRA11-22

V_L:

(SEQ ID NO: 116)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSWYYPLTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 301, respectively, in order of appearance)

V_H:

(SEQ ID NO: 117)

EVQLVESGGGLVQPGGSLRLSCAASGFTLYSSSIHWVRQAPGKGLEWVA

SISSSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YRYSSWNRAIDYWQGQTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 272, and 302, respectively, in order of appearance)

AMRA11-24

V_L:

(SEQ ID NO: 118)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSYWWPLTF

GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 303, respectively, in order of appearance)

42

-continued

V_H:

(SEQ ID NO: 119)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSIHWVRQAPGKGLEWVA

SISSSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WSKSPWYYQGIDYWQGQTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 231, and 304, respectively, in order of appearance)

15 Exemplary Antibody Clones Binding to BTK-Ibrutinib conjugate presented on HLA-A*01:01. CDR residues (Kabat scheme) in bold.

20 IBA1-4

V_L:

(SEQ ID NO: 120)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQYHYWASLI

25 TFGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 305, respectively, in order of appearance)

30 V_H:

(SEQ ID NO: 121)

EVQLVESGGGLVQPGGSLRLSCAASGFTVSSSIHWVRQAPGKGLEWVA

SISSYSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

35 QYSSSYVWPGMDYWQGQTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 306, and 307, respectively, in order of appearance)

IBA1-7

V_L:

(SEQ ID NO: 122)

DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSYWWKSLV

45 TFGQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 308, respectively, in order of appearance)

V_H:

(SEQ ID NO: 123)

50 **EVQLVESGGGLVQPGGSLRLSCAASGFTLSSSIHWVRQAPGKGLEWVA**

SISSYSGSTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

55 MHYSWQEYSSYDYGMDYWQGQTLTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 244, and 309, respectively, in order of appearance)

IBA1-8

V_L:

(SEQ ID NO: 124)

60 **DIQMTQSPSSLSASVGDRVTITCRASQSVSSAVAWYQQKPGKAPKLLIY**

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQPYYPPLTF

65 GQGTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 310, respectively, in order of appearance)

-continued

V_H:

(SEQ ID NO: 125)

EVQLVESGGGLVQPGGSLRLSCAASGFTISYSSIHWWVRQAPGKGLEWVA

SIYPSYGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WQGYYQPALDYWGQGTLVTVSS

(Bolted CDR sequences are disclosed as SEQ ID NOS 209, 311, and 312, respectively, in order of appearance)

IBA1-12

V_L:

(SEQ ID NO: 126)

DIQMTQSPSSL SASVGDRV TITCRAS QSVSSA VAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSSKYYPI

TFCQGTTKVEIKRTV

(Bolted CDR sequences are disclosed as SEQ ID NOS 166, 167, and 313, respectively, in order of appearance)

V_H:

(SEQ ID NO: 127)

EVQLVESGGGLVQPGGSLRLSCAASGFTI SSSSIHWWVRQAPGKGLEWVA

SISPYYGYTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

WGYGWWYWG LDYWGQGTLVTVSS

(Bolted CDR sequences are disclosed as SEQ ID NOS 215, 279, and 314, respectively, in order of appearance)

IBA1-13

V_L:

(SEQ ID NO: 128)

DIQMTQSPSSL SASVGDRV TITCRAS QSVSSA VAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQGHDMNPVT

FGQGTTKVEIKRTV

(Bolted CDR sequences are disclosed as SEQ ID NOS 166, 167, and 315, respectively, in order of appearance)

V_H:

(SEQ ID NO: 129)

EVQLVESGGGLVQPGGSLRLSCAASGFTLSSSIHWWVRQAPGKGLEWVA

SIYSSYGYTYYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YYYYWYGGMDYWGQGTLVTVSS

(Bolted CDR sequences are disclosed as SEQ ID NOS 215, 316, and 317, respectively, in order of appearance)

IBA1-19

V_L:

(SEQ ID NO: 130)

DIQMTQSPSSL SASVGDRV TITCRAS QSVSSA VAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSWMSDSL

TFCQGTTKVEIKRTV

(Bolted CDR sequences are disclosed as SEQ ID NOS 166, 167, and 318, respectively, in order of appearance)

V_H:

(SEQ ID NO: 131)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSYSSIHWWVRQAPGKGLEWVA

SIYPSSGYTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

GWYWMAWDYAMDYWGQGTLVTVSS

-continued

(Bolted CDR sequences are disclosed as SEQ ID NOS 209, 319, and 320, respectively, in order of appearance)

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IBA1-21

V_L:

(SEQ ID NO: 132)

DIQMTQSPSSL SASVGDRV TITCRAS QSVSSA VAWYQQKPGKAPKLLIY

10

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQM QYSGWL I

TFCQGTTKVEIKRTV

(Bolted CDR sequences are disclosed as SEQ ID NOS 166, 167, and 321, respectively, in order of appearance)

15

appearance)

V_H:

(SEQ ID NO: 133)

EVQLVESGGGLVQPGGSLRLSCAASGFTI SSSSIHWWVRQAPGKGLEWVA

20

SISSYYGYTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YY SYSSGYG YY DYFDWGAM DYWGQGTLVTVSS

25

(Bolted CDR sequences are disclosed as SEQ ID NOS 215, 322, and 323, respectively, in order of appearance)

30

(Exemplary Antibody Clones binding to EGFR-Osimertinib conjugate presented on HLA-A*02:01. CDR residues (Kabat scheme) in bold.

OEA2-1

V_L:

(SEQ ID NO: 134)

DIQMTQSPSSL SASVGDRV TITCRAS QSVSSA VAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQSSSLITF

GQGTTKVEIKRTV

40

(Bolted CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 135)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSIHWWVRQAPGKGLEWVA

45

SISSSSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YYGYVWGGYWG WY SKALD YWGQGTLVTVSS

50

(Bolted CDR sequences are disclosed as SEQ ID NOS 215, 170, and 324, respectively, in order of appearance)

OEA2-5

V_L:

(SEQ ID NO: 5)

DIQMTQSPSSL SASVGDRV TITCRAS QSVSSA VAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISLQPEDFATYYCQQYSWPIF

GQGTTKVEIKRTV

60

(Bolted CDR sequences are disclosed as SEQ ID NOS 166, 167, and 172, respectively, in order of appearance)

V_H:

(SEQ ID NO: 6)

EVQLVESGGGLVQPGGSLRLSCAASGFTI SSSYIHWWVRQAPGKGLEWVA

65

YISPSYGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

EYVTMALD YWGQGTLVTVSS

45

-continued

(Bolded CDR sequences are disclosed as SEQ ID NOS 173, 174, and 175, respectively, in order of appearance)

OEA2-12

V_L:

(SEQ ID NO: 136)

DIQMTQSPSSLSASVGDRVITTCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISSSLQPEDFATYYCQQYDWNNTYLV

TFGQGKTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 325, respectively, in order of appearance)

V_H:

(SEQ ID NO: 137)

EVQLVESGGGLVQPGGSLRLSCAASGFTIYSSSIHWRQAPGKGLEWVA

SISSYYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

YQYYGSLLYSQQWAMDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 322, and 326, respectively, in order of appearance)

OEA2-16

V_L:

(SEQ ID NO: 138)

DIQMTQSPSSLSASVGDRVITTCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISSSLQPEDFATYYCQQSSSLITF

GQGKTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

V_H:

(SEQ ID NO: 139)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSSIHWRQAPGKGLEWVA

SISSSSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

SPSSPYFM SWGWYWQY GIDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 170, and 327, respectively, in order of appearance)

OEA2-21

V_L:

(SEQ ID NO: 140)

DIQMTQSPSSLSASVGDRVITTCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISSSLQPEDFATYYCQQSSWGGLVT

FGQGKTKVEIKRTV

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 328, respectively, in order of appearance)

V_H:

(SEQ ID NO: 141)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSSIHWRQAPGKGLEWVA

SISPSYGYTSYADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

DMYEWWHWAIDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 173, 329, and 330, respectively, in order of appearance)

OEA2-24

V_L:

(SEQ ID NO: 142)

DIQMTQSPSSLSASVGDRVITTCRASQSVSSAVAWYQQKPGKAPKLLIY

SASSLYSGVPSRFSRSRGTDFTLTISSSLQPEDFATYYCQQSSSLITF

GQGKTKVEIKRTV

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-continued

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 180, respectively, in order of appearance)

5 V_H:

(SEQ ID NO: 143)

EVQLVESGGGLVQPGGSLRLSCAASGFTFSSSSIHWRQAPGKGLEWVA

SISSSSGSTS YADSVKGRFTISADTSKNTAYLQMNSLRAEDTAVYYCAR

10 YGHYLYVYWGWG WYWSAALDYWGQGTLVTVSS

(Bolded CDR sequences are disclosed as SEQ ID NOS 215, 170, and 331, respectively, in order of appearance)

15 References Related to Example 3

- Miller K R, Koide A, Leung B, Fitzsimmons J, Yoder B, Yuan H, Jay M, Sidhu S S, Koide S, Collins E J. T cell receptor-like recognition of tumor in vivo by synthetic antibody fragment. PloS one. 2012; 7 (8): e43746. Epub 2012 Aug. 24. doi: 10.1371/journal.pone.0043746. PubMed PMID: 22916301; PMCID: 3423377.
- Hattori T, Koide A, Panchenko T, Romero L A, Teng K W, Corrado A D, Koide S. Multiplex bead binding assays using off-the-shelf components and common flow cytometers. J Immunol Methods. 2020;112952. Epub 2020 Dec. 29. doi: 10.1016/j.jim.2020.112952. PubMed PMID: 33358997.

30 Example 4

This Example demonstrates single-chain Diabody (scDb) formats of Hapimmune antibodies and their effectiveness in cell killing. Data from non-limiting embodiments are presented in FIGS. 18 and 19. The results are summarized as in the brief descriptions of FIGS. 18 and 19.

- To obtain the results for FIG. 18, Raji cells and T2 cells (ATCC) were cultured in RPMI supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin. The 40 cytotoxic effect of scDbs was measured by following the protocol published previously (ref: PMID 26813960). Briefly, Raji cells or T2 cells were stained with carboxyfluorescein succinimidyl ester (CSFE, ThermoFisher, 65-0850-84), then incubated with the final 10 μ M KRAS(G12C)-45 AMG510 conjugate or 1 μ M EGFR-Osimertinib in the presence of 10 μ g/mL human beta-2 microglobulin for 4 hr. The cells were harvested using centrifugation and washed in media to remove the unbound conjugate and peptide. Peptide-drug-pulsed cells were then co-cultured with human 50 T-cells (E:T=3:1) in the presence of single-chain Diabodies (scDbs) for 19-21 hr. After incubation, cells were harvested and washed with PBS, then stained with Fixable Viability Dye eFluor660 (ThermoFisher, 65-0864-14). After washing cells, the cells were analyzed on iQue screener (Sartorius).
- 55 To obtain the results for FIG. 19, lung cancer cell lines were cultured in RPMI supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin. For cytotoxicity assays, 1×10^4 cells/well were seeded in 96-well flat bottom plates and incubated at 37° C., 5% CO₂ for 24 hours. Media 60 were replaced with fresh media supplemented with 100 nM AMG510 or DMSO, then the cells were incubated for 24 hours at 37° C. After incubation, cells were co-cultured with human T cells (E:T=5:1) and AMRA3-7_UCHT1 scDb in the presence of 100 IU/mL IL-2 for 24 hr at 37° C. Cell 65 viability was assessed by using PrestoBlue™ Cell Viability Reagent (ThermoFisher, A13261). Cytotoxicity was calculated by taking the fluorescent signal of a given well,

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subtracting the fluorescent signal from the wells that contain only T-cells, and normalizing to the fluorescent signal from the wells without scDb.

Exemplary Single-Chain Diabody Clones Targeting Both HLA-A*03:01 RAS-AMG510 and HLA-A*11:01 RAS-AMG510

The italicized sequences represent AviTag and HisTag, respectively.

AMRA3-7_UCHT1_scDb
 (SEQ ID NO: 144, sequence without tag disclosed as SEQ ID NO: 161)
 DIVRSIDIQMTQSPSSLSASVGDRVITICRASQSVSSAVAWYQQKPGKAPK
 LLIYSASSLYSGVPSRSGSRSGTDFTLTISSLQPEDFATYYCQQISYVY
 SLITFGQGTKVEIKGGGSEVQLQQSGPELVKGASMKISCKASGYSFTG
 YTMMWVKQSHGKNLEWMGLINPYKGVSTYNQFKDKATLTVDKSSSTAYM
 ELLSLTSEDAVYYCARSGYYGDSDWYFDVWGQGTTLTVSSGGGGSGGG
 SGGGSDIQMTQTTSSLSASLGDRVТИCRASQDIRNYLNWYQQKPDGT
 KLLIYYTSRLHSGVPSKFGSGSGTDSLTISNLEQEDIATYFCQQGNTL
 PWTFAAGTKLEIKGGGSEVOLVESGGGLVQPGSRLSCAASGFTFSSY
 SIHWVRQAPGKGLEWVASIYSSYGYTSYADSVKGRFTISADTSKNTAYLQ
 MNSLRAEDTAVYYCARGGWYPAMDYWGQTLTVSSLEGGGLNDIFEAQ
 KIEWHESRHHHHHH

AMRA311-16_UCHT1_scDb
 (SEQ ID NO: 145, sequence without tag disclosed as SEQ ID NO: 162)
 DIVRSIDIQMTQSPSSLSASVGDRVITICRASQSVSSAVAWYQQKPGKAPK
 LLIYSASSLYSGVPSRSGSRSGTDFTLTISSLQPEDFATYYCQQSSSL
 ITFGQGTKVEIKGGGSEVQLQQSGPELVKGASMKISCKASGYSFTGYT
 MNWVKQSHGKNLEWMGLINPYKGVSTYNQFKDKATLTVDKSSSTAYMEL
 LSLSLTSEDAVYYCARSGYYGDSDWYFDVWGQGTTLTVSSGGGGSGGG
 GGGSDIQMTQTTSSLSASLGDRVТИCRASQDIRNYLNWYQQKPDGT
 LIYYTSRLHSGVPSKFGSGSGTDSLTISNLEQEDIATYFCQQGNTL
 TFAGGKLEIKGGGSEVQLVESGGGLVQPGSRLSCAASGFTISSSI
 HWVRQAPGKGLEWAVIYSSYGYTYYADSVKGRFTISADTSKNTAYLQMN
 SLRAEDTAVYYCARYWYGHYHSYFGLDYWGQGLTVSSLEGGGLNDIF
 EAQKIEWHESRHHHHHH

Exemplary single-chain Diabody Clones Targeting HLA-A*02:01 EGFR-Osimertinib

The italicized sequences represent AviTag and HisTag, respectively.

OE2-5_UCHT1_scDb
 (SEQ ID NO: 146, sequence without tag disclosed as SEQ ID NO: 163)
 DIVRSIDIQMTQSPSSLSASVGDRVITICRASQSVSSAVAWYQQKPGKAPK
 LLIYSASSLYSGVPSRSGSRSGTDFTLTISSLQPEDFATYYCQQSYWP
 ITFGQGTKVEIKGGGSEVQLQQSGPELVKGASMKISCKASGYSFTGYT
 MNWVKQSHGKNLEWMGLINPYKGVSTYNQFKDKATLTVDKSSSTAYMEL
 LSLSLTSEDAVYYCARSGYYGDSDWYFDVWGQGTTLTVSSGGGGSGGG
 GGGSDIQMTQTTSSLSASLGDRVТИCRASQDIRNYLNWYQQKPDGT
 VKL

48

-continued

LIYYTSRLHSGVPSKFGSGSGTDSLTISNLEQEDIATYFCQQGNTL
 TFAGGKLEIKGGGSEVOLVESGGGLVQPGSRLSCAASGFTISSYI
 5 HWVRQAPGKGLEWAVIYSSYGYTSYADSVKGRFTISADTSKNTAYLQMN
 SLRAEDTAVYYCAREVYTMALDYWGQGLTVSSLEGGGLNDIFEAQKI
 EWHEHESRHHHHHH
 10 OEA2-21_UCHT1_scDb
 (SEQ ID NO: 147, sequence without tag disclosed as SEQ ID NO: 164)
 DIVRSIDIQMTQSPSSLSASVGDRVITICRASQSVSSAVAWYQQKPGKAPK
 15 LLIYSASSLYSGVPSRSGSRSGTDFTLTISSLQPEDFATYYCQQSSWGG
 LVTFGQGTKVEIKGGGSEVQLQQSGPELVKGASMKISCKASGYSFTGY
 TMNWVKQSHGKNLEWMGLINPYKGVSTYNQFKDKATLTVDKSSSTAYM
 20 LLSLTSEDAVYYCARSGYYGDSDWYFDVWGQGTTLTVSSGGGGSGGG
 GGGSDIQMTQTTSSLSASLGDRVТИCRASQDIRNYLNWYQQKPDGT
 25 VKL
 LLIYYTSRLHSGVPSKFGSGSGTDSLTISNLEQEDIATYFCQQGNTL
 WTPAGGKLEIKGGGSEVOLVESGGGLVQPGSRLSCAASGFTFSSY
 30 IHWVRQAPGKGLEWAVIYSSYGYTSYADSVKGRFTISADTSKNTAYLQMN
 NSLRAEDTAVYYCARDMYEWWHWAIDYWGQGLTVSSLEGGGLNDIFE
 AQKIEWHESRHHHHHH

Example 5

This Example demonstrates scDb and 2+1 CrossMab antibodies constructed with the AMRA3-7 clone and their effectiveness in cell killing. Data from non-limiting embodiments are presented in FIGS. 21-22.

To obtain the results for FIG. 21 (A) and FIG. 22 (B), Raji cells (ATCC) were cultured in RPMI supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin. 40 The cytotoxic effects of scDb and CrossMab were measured by following the protocol published previously (ref: PMID 26813960). Briefly, Raji cells were stained with carboxyfluorescein succinimidyl ester (CSFE, ThermoFisher, 65-0850-84), then incubated with 10 μ M KRAS(G12C)-AMG510 conjugate or 10 μ M KRAS(WT) peptide (final concentrations) in the presence of 10 μ g/mL human beta-2 microglobulin for 4 hr. The cells were harvested by centrifugation and washed in media to remove the unbound 45 conjugate and peptide. Peptide-drug-pulsed cells were then co-cultured with human T-cells (E:T=3:1) in the presence of scDb or CrossMab for 19 hr. After incubation, cells were harvested and washed with PBS, then stained with Fixable Viability Dye eFluor660 (ThermoFisher, 65-0864-14). After 50 washing again, cells were analyzed on iQue screener (Satorius).

To obtain the results for FIGS. 21 (B) and 22 (C), H2122 cells (ATCC) were cultured in RPMI supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin. 60 For cytotoxicity assays, 5×10^3 cells/well were seeded in 96-well flat bottom plates in the presence of 1 mM AMG510 or DMSO and 5 μ g/mL human beta-2 microglobulin, and then were incubated at 37° C., 5% CO₂ for 48 hours. After incubation, cells were co-cultured with human T cells (E:T=10:1) and AMRA3-7D scDb or CrossMab in the presence of 10 ng/ml IL7 and IL 15 for 24 hr at 37° C. Dead 65 cells were measured by using CytoTox-Glo cytotoxic assay

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(Promega, G9290). The luminescent signal of a given well was calculated by subtracting the signal from wells that contain H2122 and T-cells without scDb or CrossMab constructs.

To obtain the results for FIG. 22 (A), Jurkat and Raji cells (ATCC) were cultured in RPMI supplemented with 10% fetal bovine serum (FBS) and penicillin/streptomycin. Cells were washed twice with PBS, then incubated with AMRA3-7D CrossMab at 4° C. for 30 min. After washing three times with PBS containing 1% BSA (PBS/BSA), cells were stained with Alexa647 Goat Anti-Human IgG Fc (Jackson ImmunoResearch, 109-605-098). After incubation, cells were washed three times with PBS/BSA and analyzed on iQue screener (Sartorius).

AMRA3-7D_UCHT1_scDb

(SEQ ID NO: 148, sequence without tag disclosed as SEQ ID NO: 165)

DIVRSIDIQMTQSPSSLSASVGDRVTITC**RASQSVSSAVAWYQQKPGKAPK**
LILIYASSSLYSGVPSRSGSRSGTDFTLTISLQPEDFATYYCQQQISYVK****

KLITFGQGTKVEIKGGGGSEVQLQQSGPELVKGASMKISCKASGY**SFTG
 YTMNWVKQSHGKLNLEWMLINPFGKGVSTYNQFKDKATLTVDKSSSTAYM
 ELLSLTSEDAVYYCARSGYYGSDDWYFDVWGQGTTLTVSSGGGGSGGG
 SGGGSDIQMTQTTSSLSASLGDRVТИSCRASQDIRNYLNWYQQKPDGT
 KLLIYYTSRLHSGVPSKFGSGSGTDYSLTISNLEQEDIATYFCQQQNTL
 PWTFAAGGTKEI KGGGGSEVQLVESGGGLVQPGGSLRLSCAASGFTFS**DY**
 SIHWVRQAPGKGLEWVA**SISSSGSTSYADSVKGRFTISADTSKNTAYLQ**
 MNSLRAEDTAVYYC**ARGGWYPAMDYWGQGTLTVVSSLEGGGLNDIFEAQ****

KIEWHESRHHHHHH

(Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, 168, 169, 170, and 171, respectively, in order of appearance)

AMRA3-7D_CrossMab

>Chain A. QMY30735.1

(SEQ ID NO: 149)

EVQLLESGGGLVQPGGSLRLSCAASGFTFS**TYAMN**WVRQAPGKGLEW**SR**
 IRSKYNNYATYYADSVKGRFTISRDDSKNTLYL**QMN**SLRAEDTAVYYCVR
 HGNFGNSYVSWFAYWGQGTLTVSSASVAAPS**VIFPPSDEQL**KSGTASV
 VCLNNFYPREAKV**QWKVDNALQGSNQESVTEQDSKDSTYS**SSLTL**TL**
 KADYEKHKVYACEVTH**QGLSSPVTKSFNR**GEC

>Chain B.

(SEQ ID NO: 150)

EVQLVESGGGLVQPGGSLRLSCAASGFTFS**DY**SIHWRQAPGKGLEW**VAS**
 I**SISSSGSTSYADSVKGRFTISADTSKNTAYLQ**MNSLRAEDTAVYYC**ARGG**
 WYPAMDYWGQGTLTVSSASTKGPSVFLAPSSK**STSGGTAALGCLVEDY**
 FPEPVTVWSNGALTSGVHTFP**AVLQSSGLYSLSSVVTV**PSSSLG**TQTYI**
 CNVNHKPSNTKVDEKVEPKSCDG**GGGGGGGGGSQAVV**TQEP**SLTVSPGGTV**
 TLTCGSSTGAVTTSNANWV**QEKPGQA**FRLGLIGGTNKRAPGTPARFSGSL
 LGGKAAL**TLSGAQ**PEDEAEYYCALWYSNLWV**FGGGT**KLT**VLSSASTKGP**
 VFPLAPSSK**STSGGTAALGCLV**DYFPEPVTVWSN**GALTSGVHTFP**AVL
 QSSGLYSLSSVVTV**PSSSLG**TQTYI**CNVNHKPSNTKVDKK**VEPK**SCDKTH**
 TCPPCPA**PEAAGG**PSVFLFPPK**KDTLM**ISRTPEV**TCVVVD**VSHEDPEV**K**

50

-continued

FNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVS
 NKALGAPIEKTISKAKGQPREPQVYTLPPCRDELTKNQVSLWCLVKGFYP
 SDIAVEWESNGQPENNYKTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFS
 CSVMEALHNHYTQKSLSLSP
 >Chain C.
 (SEQ ID NO: 151)
 EVQLVESGGGLVQPGGSLRLSCAASGFTFS**DY**SIHWRQAPGKGLEW**VAS**
 I**SISSGSTSYADSVKGRFTISADTSKNTAYLQ**MNSLRAEDTAVYYC**ARGG**
 WYPAMDYWGQGTLTVSSASTKGPSVFLAPSSK**STSGGTAALGCLVEDY**
 FPEPVTVWSNGALTSGVHTFP**AVLQSSGLYSLSSVVTV**PSSSLG**TQTYI**
 CNVNHKPSNTKVDEKVEPKSCDKTHTC**CPPCPA**PEAAGG**PSVFLFPPKPKD**
 TLMISRTPEV**TCVVVD**VSHEDPEV**KFNWY**VDGVEVHNAKTKPREEQYNST
 YRVVSVLTVLHQDWLNGKEYKCKVSNKALGAPIEKTISKAKGQPREPQVC
 TLPPSRDELT**KNQVSL**CAVKG**FYPSDI**AVEWESNGQPENNYKTPPVLD
 SDGSFFL**VSKLT**VDKSRWQQGNVFSCSVMEALHNHYTQKSLSLSP
 >Chain D.
 (SEQ ID NO: 152)
 DIQMTQSPSSLSASVGDRVTITC**RASQSVSSAVAWYQQKPGKAPK**LI**Y**
 ASSLYSGVPSRSGSRSGTDFTLT**ISLQPEDFATYYC**QQQISYVK**KL**ITF
 GQG**TKV**EIKRTVAAPSVF**I**FPPSDRK**L**KG**T**ASV**V**CV**LNNF**PREAKV**QW**
 KVDNAL**QSGNSQESV**TEQDSKD**STYS**SSL**TL**SKADYEKHKVYACEV**TH**
 QGLSSPVTKSFNR**GEC**

FIG. 21 shows the cytotoxic effect of AMRA3-7D in the single-chain Diabody (scDb) format. (A) Raji cells were first pulsed with AMG510 conjugated to a peptide corresponding to a fragment of KRAS(G12C) or a control peptide corresponding to KRAS(wild type). Pulsed cells were co-cultured with human T cells (Effector:Target=3:1) in the presence of scDb at the indicated concentrations. After incubation for 17 hours, dead cells were stained and detected by flow cytometry. (B) H2122 cells were first incubated with AMG510 or DMSO. The cells were then co-cultured with human T cells (Effector:Target=10:1) in the presence of scDb at the indicated concentrations. After incubation, dead cells were measured by detecting a distinct intracellular protease activity released from membrane-compromised cells. Data shown here are from triplicate measurements. Error bars indicate the s.d. Where error bars are not visible, the errors are smaller than the symbols. Anti-HLA-A3 is a positive control.

FIG. 22 shows the CD3 binding properties and cytotoxic effects of AMRA3-7D in the CrossMab format. (A) Binding titration curve of AMRA3-7 CrossMab to Jurkat (CD3 positive) and Raji (CD3 negative) cells. The apparent K_D value is shown. Note that cells were not pulsed with any peptides. (B, C) Cytotoxic effects of AMRA3-7D CrossMab on Raji cells pulsed with an exogenous peptide-drug conjugate (B) and on H2122 cells treated with the drug (C). Methods are the same as in FIG. 22. Data shown here are from triplicate measurements. Error bars indicate the s.d. Where error bars are not visible, the errors are smaller than the symbols. Anti-HLA-A3 serves as a positive control.

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Example 6

This example demonstrates deep mutational analysis of the AMR-A3-7D and OEA2-5 antibodies. Data from non-limiting embodiments is presented in FIGS. 23 and 24.

To identify mutations in CDRs of AMR-A3-7D and OEA2-5 that retain antigen binding, we performed deep mutational scanning on residues CDR-L3 and CDR-H3. In the yeast display format, each of the CDR-L3 and CDR-H3 residues were mutated to all genetically encoded amino acids using the NNK codon (N=A, T, G and C; K=G and T), one residue at a time. The resulting pool of mutants was combined, and the library was subjected to FACS using the relevant antigen, i.e., AMG510-KRAS(G12C) peptide in complex with HLA-A*03:01 for AMR-A3-7D and Osimertinib-EGFR in complex with HLA-A*02:01. We used different antigen concentrations in order to adjust the stringency of library sorting. Vectors recovered from binding-capable and binding-incapable pools were analyzed by deep sequencing on an Illumina MiSeq instrument. Mutations found in different pools were deduced from the DNA sequencing analysis.

From this analysis, the disclosure provides the following permissible mutations at each CDR position as shown in the tables below. As references, the VL and VH sequences of the parent clones are shown, with the analyzed CDR residues in bold and italics. In embodiments, the disclosure includes each mutation alone, and all combination of mutations. Thus, as evident from the Tables, the disclosure included the described CDRs with 1, 2, 3, 4, 5, 6, 7, or 8 mutations as indicated in the Tables. The disclosure includes additional amino acid chances, such as in CDR1, CDR2, and in the framework sequences.

AMRA3-7D
V_L:
 (SEQ ID NO: 3)
DIQMTQSPSS LSASVGDRV^T ITCRASQSV^S SAVAWYQQKP
GKAPKLLIYS ASSLYSGVPS RFSGSRSGTD FTLLTISSLQP
EDFATYYCQQ ISYVKKLITF GQGTKVEIKR TV
 (Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 168, respectively, in order of appearance)
V_H:
 (SEQ ID NO: 4)
EVQLVESGGG LVQPGGSLRL SCAASGFTFS DYSIHWWVRQA
PGKGLEWVAS ISSSSGSTSY ADSVKGRFTI SADTSKNTAY
LQMNSLRAED TAVYYCARGG WYPAMDYWQQ GTLTVVSS
 (Bolded CDR sequences are disclosed as SEQ ID NOS 169, 170, and 171, respectively, in order of appearance)

TABLE A

Parental and consensus sequences disclosed as SEQ ID NOS 153-154, respectively			
Position (VL)	Parental amino acid residue	Permissible mutation	Consensus amino acid residue
91	I	A, L, P, S, T, V	A, I, L, P, S, T, V
92	S	K, R, T	K, R, S, T
93	Y	F	F, Y
94	V	I, R, T	I, R, T, V
95	K	A, E, H, L, M, N, Q, R, S, T, Y	A, E, H, K, L, M, N, Q, R, S, T, Y

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TABLE A-continued

Parental and consensus sequences disclosed as SEQ ID NOS 153-154, respectively			
Position (VL)	Parental amino acid residue	Permissible mutation	Consensus amino acid residue
96	K	R	K, R
97	L	A, C, D, E, G, H, K, M, N, P, Q, R, S, T, V, W	A, C, D, E, G, H, K, L, M, N, P, Q, R, S, T, V, W
98	I	L, V	I, L, V

TABLE B

Parental and consensus sequences disclosed as SEQ ID NOS 155-156, respectively			
Position (VH)	Parental amino acid residue	Permissible mutation	Consensus amino acid residue
99	G	G	G
100	G	A, C, E, H, K, L, M, N, P, Q, R, S, T, W, Y	A, C, E, G, H, K, L, M, N, P, Q, R, S, T, W, Y
101	W	G, P, R, T	G, P, R, T, W
102	Y	A, D, E, F, G, H, I, K, M, N, Q, R, S, T, W	A, D, E, F, G, H, I, K, M, N, Q, R, S, T, W, Y
103	P	A, V	A, P, V
104	A	C, G, K, L, M, Q, R, S, T, Y	A, C, G, K, L, M, Q, R, S, T, Y

Deep Mutational Scanning of OEA2-5

V_L:
 (SEQ ID NO: 5)
DIQMTQSPSS LSASVGDRV^T ITCRASQSV^S SAVAWYQQKP
GKAPKLLIYS ASSLYSGVPS RFSGSRSGTD FTLLTISSLQP
EDFATYYCQQ ISYVKKLITF GQGTKVEIKR TV
 (Bolded CDR sequences are disclosed as SEQ ID NOS 166, 167, and 172, respectively, in order of appearance)
V_H:
 (SEQ ID NO: 6)
EVQLVESGGG LVQPGGSLRL SCAASGFTFS SSYIHWWVRQA
PGKGLEWVAY ISPSSYGSTSY ADSVKGRFTI SADTSKNTAY
LQMNSLRAED TAVYYCAREY VTMALDYWGQ GTLTVVSS
 (Bolded CDR sequences are disclosed as SEQ ID NOS 173, 174, and 175, respectively, in order of appearance)

TABLE C

Parental and consensus sequences disclosed as SEQ ID NOS 157-158, respectively			
Position (VH)	Parental amino acid residue	Permissible mutation	Consensus amino acid residue
91	Y		Y
92	S	A	A, S
93	Y	A, C, D, E, F, G, H, I, K, L, M, N, Q, R, S, T,	A, C, D, E, F, G, H, I, K, L, M, N, Q, R, S, T, V, W, Y
94	W	V, W	W

TABLE C-continued

Parental and consensus sequences disclosed as SEQ ID NOS 157-158, respectively			
Position (VH)	Parental amino acid residue	Permissible mutation	Consensus amino acid residue
95	P		P
96	I	E, P	E, I, P

TABLE D

Parental and consensus sequences disclosed as SEQ ID NOS 159-160, respectively			
Position (VH)	Parental amino acid residue	Permissible mutation	Consensus amino acid residue
99	E	D	D, E
100	Y	E, F, L, Q, S, W	E, F, L, Q, S, W, Y
101	V	T, I	I, T, V
102	T	A, E, N, S	A, E, N, S, T
103	M	F, Y	F, M, Y
104	A	C, G, S, T	A, C, G, S, T
105	L	A, C, E, F, H, I, K, M, N, P, Q, S, T, V	A, C, E, F, H, I, K, L, M, N, P, Q, S, T, V

As will be evident from the foregoing tables, in one embodiment, the binding partner comprises a light chain that comprises a complementary determining region 3 (CDR3) that comprises the sequence SEQ ID NO: 154 and a heavy chain that comprises a CDR3 that comprises the sequence of SEQ ID NO:156. In another embodiment, the binding partner the binding partner comprises a light chain that comprises a CDR3 that comprises the sequence of SEQ ID NO:158 and a heavy chain that comprises a CDR3 that comprises the sequence of SEQ ID NO:160.

FIG. 23 describes deep mutational scanning analysis of the CDR-L3 and H3 residues of AMR-A3-7D. (a) Representative flow cytometry profiles of yeast cells displaying AMRA3-7D or its deep mutational scanning library popu-

lations. Binding to 5 nM HLA-A*03:01 presenting AMG510 conjugated to the Cys residue in the peptides, VVGACGVGK (SEQ ID NO: 1), was measured. The profile of the parental antibody, AMRA3-7D, is shown on the left, and those for sorted subsets of the deep mutational scanning library are shown on the right. The library was sorted with 1, 3, and 10 nM target, referred to as conditions 1, 2 and 3, respectively, and the nonbinder pool was sorted with 50 nM target. (b) The prevalence of mutations in the sorted subsets of the deep mutational scanning library is presented in a heat map format. The number of deep sequencing reads were normalized to the total reads for each sorted pool and multiplied by 1000. Crosses indicate the wild-type amino acid.

FIG. 24 describes results from deep mutational scanning analysis of CDR-L3 and H3 residues of OEA2-5. (a) Representative flow cytometry profiles of yeast cells displaying OEA2-5 in single-chain Fv format and its deep mutational scanning library populations. Binding to 1.5 nM streptavidin tetramer saturated with biotinylated HLA-A*02:01 presenting Osimertinib conjugated to the Cys residue in the peptides, QLMPFGCLL (SEQ ID NO: 30), was measured. The profile of the parental antibody, OEA2-5, is shown on the left, and those for sorted subsets of the deep mutational scanning library are shown on the right. The library was sorted with 12.5, 2.5, and 0.5 nM target, referred to as conditions 1, 2 and 3, respectively, and the nonbinder pool was sorted with 12.5 nM target. (b) The prevalence of mutations in the sorted subsets of the deep mutational scanning library is presented in a heat map format. The number of deep sequencing reads were normalized to the total reads for each sorted pool and multiplied by 1000. Crosses indicate the wild-type amino acid.

Other embodiments of the disclosure will be apparent to those skilled in the art from consideration of the specification and practice of the disclosure disclosed herein. It is intended that the specification and examples be considered as exemplary only, with a true scope and spirit of the disclosure being indicated by the following claims.

SEQUENCE LISTING

```

Sequence total quantity: 420
SEQ ID NO: 1      moltype = AA length = 9
FEATURE           Location/Qualifiers
REGION            1..9
note = Description of Artificial Sequence: Synthetic peptide
source             1..9
mol_type = protein
organism = synthetic construct

SEQUENCE: 1
VVGACGVGK

SEQ ID NO: 2      moltype = AA length = 10
FEATURE           Location/Qualifiers
REGION            1..10
note = Description of Artificial Sequence: Synthetic peptide
source             1..10
mol_type = protein
organism = synthetic construct

SEQUENCE: 2
VVVGACGVGK

SEQ ID NO: 3      moltype = AA length = 112
FEATURE           Location/Qualifiers
REGION            1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source             1..112

```

-continued

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mol_type = protein
organism = synthetic construct

SEQUENCE: 3
DIQMTQSPSS LSASVGRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ ISYVKKLITF QQGTVKEIKR TV      112

SEQ ID NO: 4          moltype = AA length = 118
FEATURE             Location/Qualifiers
REGION              1..118
note = Description of Artificial Sequence: Synthetic
polypeptide
source               1..118
mol_type = protein
organism = synthetic construct

SEQUENCE: 4
EVQLVESGGG LVQPGGSLRL SCAASGFTFS DYSIHWRQPA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGG WYPAMDYWGQ GTLTVSS     118

SEQ ID NO: 5          moltype = AA length = 110
FEATURE             Location/Qualifiers
REGION              1..110
note = Description of Artificial Sequence: Synthetic
polypeptide
source               1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 5
DIQMTQSPSS LSASVGRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ YSYWPITPGQ GTKVEIKRTV      110

SEQ ID NO: 6          moltype = AA length = 118
FEATURE             Location/Qualifiers
REGION              1..118
note = Description of Artificial Sequence: Synthetic
polypeptide
source               1..118
mol_type = protein
organism = synthetic construct

SEQUENCE: 6
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSYIHWRQPA PGKGLEWVAY ISPSSYGS 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCAREY VTMALDYWGQ GTLTVSS     118

SEQ ID NO: 7          moltype = AA length = 18
FEATURE             Location/Qualifiers
source               1..18
mol_type = protein
organism = Thosea asigna virus

SEQUENCE: 7
EGRGSSLTCG DVEENPGP                                         18

SEQ ID NO: 8          moltype = AA length = 17
FEATURE             Location/Qualifiers
source               1..17
mol_type = protein
organism = Teschovirus A

SEQUENCE: 8
ATNFSLKQAG DVENPGP                                         17

SEQ ID NO: 9          moltype = AA length = 19
FEATURE             Location/Qualifiers
source               1..19
mol_type = protein
organism = Equine rhinitis A virus

SEQUENCE: 9
QCTNYALKLA GDVESNPGP                                         19

SEQ ID NO: 10         moltype = AA length = 21
FEATURE             Location/Qualifiers
source               1..21
mol_type = protein
organism = Foot-and-mouth disease virus

SEQUENCE: 10
VKQTLNFDLK LAGDVESNPG P                                         21

SEQ ID NO: 11         moltype = AA length = 8
FEATURE             Location/Qualifiers
REGION              1..8
note = Description of Artificial Sequence: Synthetic peptide
source               1..8

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-continued

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mol_type = protein
organism = synthetic construct

SEQUENCE: 11
GGGSAAA                                         8

SEQ ID NO: 12          moltype = AA  length = 20
FEATURE           Location/Qualifiers
REGION            1..20
note = Description of Artificial Sequence: Synthetic peptide
source             1..20
mol_type = protein
organism = synthetic construct

SEQUENCE: 12
GGGGSGGGGS GGGGSGGGGS                           20

SEQ ID NO: 13          moltype = AA  length = 6
FEATURE           Location/Qualifiers
REGION            1..6
note = Description of Artificial Sequence: Synthetic 6xHis
tag
source             1..6
mol_type = protein
organism = synthetic construct

SEQUENCE: 13
HHHHHHH                                         6

SEQ ID NO: 14          moltype = AA  length = 107
FEATURE           Location/Qualifiers
REGION            1..107
note = Description of Artificial Sequence: Synthetic
polypeptide
source             1..107
mol_type = protein
organism = synthetic construct

SEQUENCE: 14
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS  60
RFGSGRSRSGTD FTLTISSLQP EDFATYYCQQ DWYFPITFGQ GTKVEIK                107

SEQ ID NO: 15          moltype = AA  length = 128
FEATURE           Location/Qualifiers
REGION            1..128
note = Description of Artificial Sequence: Synthetic
polypeptide
source             1..128
mol_type = protein
organism = synthetic construct

SEQUENCE: 15
EVQLVESGGG LVQPGGSLRL SCAASGFTFS SYYIHWRQQA PGKGLEWVAS ISPSSGSTYY  60
ADSVVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYG GRSYWQKQDS YFYQHGLDYW    120
QQTLVSS                                         128

SEQ ID NO: 16          moltype = AA  length = 107
FEATURE           Location/Qualifiers
REGION            1..107
note = Description of Artificial Sequence: Synthetic
polypeptide
source             1..107
mol_type = protein
organism = synthetic construct

SEQUENCE: 16
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS  60
RFGSGRSRSGTD FTLTISSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIK                107

SEQ ID NO: 17          moltype = AA  length = 125
FEATURE           Location/Qualifiers
REGION            1..125
note = Description of Artificial Sequence: Synthetic
polypeptide
source             1..125
mol_type = protein
organism = synthetic construct

SEQUENCE: 17
EVQLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWWRQQA PGKGLEWVAS ISSYSGYTSY  60
ADSVVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSY SYSEFRYYYS GQGMDYWGQG    120
TLVSS                                         125

SEQ ID NO: 18          moltype = AA  length = 107
FEATURE           Location/Qualifiers
REGION            1..107

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source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..107
mol_type = protein
organism = synthetic construct

SEQUENCE: 18
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTISSSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIK 107

SEQ ID NO: 19      moltype = AA length = 123
FEATURE          Location/Qualifiers
REGION           1..123
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..123
mol_type = protein
organism = synthetic construct

SEQUENCE: 19
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWVRQA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSN YGWRWHLVGM DYWGQQTLVT 120
VSS                      123

SEQ ID NO: 20      moltype = AA length = 107
FEATURE          Location/Qualifiers
REGION           1..107
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..107
mol_type = protein
organism = synthetic construct

SEQUENCE: 20
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTISSSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIK 107

SEQ ID NO: 21      moltype = AA length = 123
FEATURE          Location/Qualifiers
REGION           1..123
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..123
mol_type = protein
organism = synthetic construct

SEQUENCE: 21
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWVRQA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSP YVYYWYMGF DYWGQQTLVT 120
VSS                      123

SEQ ID NO: 22      moltype = AA length = 15
FEATURE          Location/Qualifiers
REGION           1..15
source          note = Description of Artificial Sequence: Synthetic peptide
1..15
mol_type = protein
organism = synthetic construct

SEQUENCE: 22
YKLVVVGACG VGKSA                                         15

SEQ ID NO: 23      moltype = AA length = 10
FEATURE          Location/Qualifiers
REGION           1..10
source          note = Description of Artificial Sequence: Synthetic peptide
1..10
mol_type = protein
organism = synthetic construct

SEQUENCE: 23
SSSSCSSSSW                                         10

SEQ ID NO: 24      moltype = AA length = 110
FEATURE          Location/Qualifiers
REGION          1..110
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 24
QSVLIQPRSV SGSPGQSVTI SCTGTSSDVG GYNVYWSYQQ HPGKAPKLMI YDVSKRPSGV 60
PDRFSGSKSG NTASLTVSGL QAEDEADYYC GSYADTDTIV FGTGKLTQLV 110

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-continued

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SEQ ID NO: 25      moltype = AA length = 119
FEATURE
REGION          Location/Qualifiers
1..119
note = Description of Artificial Sequence: Synthetic
polypeptide
1..119
source           mol_type = protein
organism = synthetic construct

SEQUENCE: 25
QVQLVQSEPE VKKPQSSVKL SCKASGGTFS TDAITWVRQA PGQGLEMMGG IIPLLDSVDY 60
AQKFQGRVTV SADKSTGTAY MEVRSLGSED TAKYYCAKWS SVDTGLDYWG QGTLTVSS 119

SEQ ID NO: 26      moltype = AA length = 126
FEATURE
REGION          Location/Qualifiers
1..126
note = Description of Artificial Sequence: Synthetic
polypeptide
1..126
source           mol_type = protein
organism = synthetic construct

SEQUENCE: 26
QVQLQESGPG LVKPSETLSL TCTVSGDSII NDPHYWGWIR QSPGKGLEWI GSTSHSGHTY 60
FNPSLKSRRVS MSIDVAKNQF SLNVRSTAA DTAVYYCARM RYYYSGTYPV YYFDYWGQGT 120
LVTVSS          126

SEQ ID NO: 27      moltype = AA length = 110
FEATURE
REGION          Location/Qualifiers
1..110
note = Description of Artificial Sequence: Synthetic
polypeptide
1..110
source           mol_type = protein
organism = synthetic construct

SEQUENCE: 27
SVLTQPPSA SGTPGQRVTI SCSGSSSNIG SNFVSWYQQL PGTAPKLLIS SNNQRPSGVP 60
DRFGSKSDT SASLAISGLQ SEDEADYYCA AWDDSLNGPV FGGGTQLTVAL 110

SEQ ID NO: 28      moltype = AA length = 121
FEATURE
REGION          Location/Qualifiers
1..121
note = Description of Artificial Sequence: Synthetic
polypeptide
1..121
source           mol_type = protein
organism = synthetic construct

SEQUENCE: 28
QVQLVQSEAE VKKPQSSVKV SCKASGGTFS RYGVSWVRQA PGQGLEMMGG IIPMFGTANY 60
AQKFQGRVTI TADESTSTAY MELRSLRSED TAVYYCARGD NSAYSDAFNI WGQGTMVTVS 120
S               121

SEQ ID NO: 29      moltype = AA length = 9
FEATURE
source           Location/Qualifiers
1..9
mol_type = protein
organism = Homo sapiens

SEQUENCE: 29
VVGAGGVGK          9

SEQ ID NO: 30      moltype = AA length = 9
FEATURE
REGION          Location/Qualifiers
1..9
note = Description of Artificial Sequence: Synthetic peptide
1..9
source           mol_type = protein
organism = synthetic construct

SEQUENCE: 30
QLMPFGCLL          9

SEQ ID NO: 31      moltype = AA length = 10
FEATURE
REGION          Location/Qualifiers
1..10
note = Description of Artificial Sequence: Synthetic peptide
1..10
source           mol_type = protein
organism = synthetic construct

SEQUENCE: 31
YMANGCLLN          10

SEQ ID NO: 32      moltype = AA length = 111

```

-continued

FEATURE Location/Qualifiers
REGION 1..111
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..111
mol_type = protein
organism = synthetic construct

SEQUENCE: 32
DIQMTQSPSS LSASVGDRV TITCRASQSV SAVA WYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ SGWSYPITFG QGTKVEIKRT V 111

SEQ ID NO: 33 moltype = AA length = 117
FEATURE Location/Qualifiers
REGION 1..117
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..117
mol_type = protein
organism = synthetic construct

SEQUENCE: 33
EVOLVESGGG LVQPGGSLRL SCAASGFTFY SSYIHWRQ A PGKGLEWVAS ISPYYGSTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSS YYALDYWGQ TLTVSS 117

SEQ ID NO: 34 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 34
DIQMTQSPSS LSASVGDRV TITCRASQSV SAVA WYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ ISYVYSLITF GQGTKVEIKR TV 112

SEQ ID NO: 35 moltype = AA length = 118
FEATURE Location/Qualifiers
REGION 1..118
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..118
mol_type = protein
organism = synthetic construct

SEQUENCE: 35
EVQLVESGGG LVQPGGSLRL SCAASGFTFS SYSIHWRQ A PGKGLEWVAS IYSSYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGG WYPAMDYWGQ GTLTVSS 118

SEQ ID NO: 36 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 36
DIQMTQSPSS LSASVGDRV TITCRASQSV SAVA WYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ ISYVKKLITF GQGTKVEIKR TV 112

SEQ ID NO: 37 moltype = AA length = 118
FEATURE Location/Qualifiers
REGION 1..118
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..118
mol_type = protein
organism = synthetic construct

SEQUENCE: 37
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SYSIHWRQ A PGKGLEWVAS IYSSYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGG WYPAMDYWGQ GTLTVSS 118

SEQ ID NO: 38 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

-continued

SEQUENCE: 38
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSLQP EDLATYYCQQ YOYGYNLITF GQGTKVEIKR TV 112

SEQ ID NO: 39 moltype = AA length = 124
 FEATURE Location/Qualifiers
 REGION 1..124
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..124
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 39
 EVQLVESGGG LVQPGGSLRL SCAASGFTIS YSSIHWVRQA PGKGLEWVAS IYSYSGSTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYS YGWVGPGWRA IDYWGQGTLV 120
 TVSS 124

SEQ ID NO: 40 moltype = AA length = 112
 FEATURE Location/Qualifiers
 REGION 1..112
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..112
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 40
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SSSVYKLLTF GQGTKVEIKR TV 112

SEQ ID NO: 41 moltype = AA length = 120
 FEATURE Location/Qualifiers
 REGION 1..120
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..120
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 41
 EVQLVESGGG LVQPGGSLRL SCAASGFTVY YSSIHWVRQA PGKGLEWVAS ISSSYSYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TALYYCARGG PGWYRAMDYW GQGTLTVSS 120

SEQ ID NO: 42 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 42
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 43 moltype = AA length = 123
 FEATURE Location/Qualifiers
 REGION 1..123
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..123
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 43
 EVQLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWVRQA PGKGLEWVAS ISSSSGSTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGY FYYGWWAMAF DYWGQGTLVT 120
 VSS 123

SEQ ID NO: 44 moltype = AA length = 112
 FEATURE Location/Qualifiers
 REGION 1..112
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..112
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 44
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SQWYEPLITF GQGTKVEIKR TV 112

SEQ ID NO: 45 moltype = AA length = 127

-continued

FEATURE
REGION Location/Qualifiers
1..127
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..127
mol_type = protein
organism = synthetic construct

SEQUENCE: 45
EVQLVESGGG LVQPGGSLRL SCAASGFTIY SSYIHWRQQA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSY SYMSQWGWYQ YSGMDYWQGQ 120
TLTVSS 127

SEQ ID NO: 46 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 46
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ GSYTYRLITF GQGTKVEIKR TV 112

SEQ ID NO: 47 moltype = AA length = 119
FEATURE Location/Qualifiers
REGION 1..119
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..119
mol_type = protein
organism = synthetic construct

SEQUENCE: 47
EVQLVESGGG LVQPGGSLRL SCAASGFTVS YSSIHWVRQQA PGKGLEWVAS ISSSSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYA WWAHGLDYWG QGTLTVSS 119

SEQ ID NO: 48 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 48
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ ASYWYNLPTF GQGTKVEIKR TV 112

SEQ ID NO: 49 moltype = AA length = 123
FEATURE Location/Qualifiers
REGION 1..123
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..123
mol_type = protein
organism = synthetic construct

SEQUENCE: 49
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SYSIHWRQQA PGKGLEWVAS IYSSYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARQY SMHFPWGYGM DYWGQGTLVT 120
VSS 123

SEQ ID NO: 50 moltype = AA length = 110
FEATURE Location/Qualifiers
REGION 1..110
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 50
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SDMPPITFGQ GTKVEIKRTV 110

SEQ ID NO: 51 moltype = AA length = 121
FEATURE Location/Qualifiers
REGION 1..121
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..121

-continued

```

mol_type = protein
organism = synthetic construct

SEQUENCE: 51
EVQLVESGGG LVQPGGSLRL SCAASGFTFY SSSIHWVRQA PGKGLEWVAY IYSSSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARPV NYYYQGALDY WGQGTLVTVS 120
S 121

SEQ ID NO: 52      moltype = AA length = 110
FEATURE           Location/Qualifiers
REGION            1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source             1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 52
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGSRSGTD FTLTSSLQP EDFATYYCQQ YYVFPITFGQ GTKVEIKRTV 110

SEQ ID NO: 53      moltype = AA length = 130
FEATURE           Location/Qualifiers
REGION            1..130
note = Description of Artificial Sequence: Synthetic
               polypeptide
source             1..130
mol_type = protein
organism = synthetic construct

SEQUENCE: 53
EVOLVESGGG LVQPGGSLRL SCAASGFTVY SSSIHWVRQA PGKGLEWVAS ISPSSGYTYY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARIH YMFEYDKGES KWGYYGFDYW 120
GQGTLTVSS 130

SEQ ID NO: 54      moltype = AA length = 110
FEATURE           Location/Qualifiers
REGION            1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source             1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 54
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGSRSGTD FTLTSSLQP EDFATYYCQQ SQYFPITFGQ GTKVEIKRTV 110

SEQ ID NO: 55      moltype = AA length = 122
FEATURE           Location/Qualifiers
REGION            1..122
note = Description of Artificial Sequence: Synthetic
               polypeptide
source             1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 55
EVOLVESGGG LVQPGGSLRL SCAASGFTIS YSSIHWVRQA PGKGLEWVAS IYSYYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARNS WSWYSGVGMD YWGQGTLTV 120
SS 122

SEQ ID NO: 56      moltype = AA length = 110
FEATURE           Location/Qualifiers
REGION            1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source             1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 56
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGSRSGTD FTLTSSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 57      moltype = AA length = 123
FEATURE           Location/Qualifiers
REGION            1..123
note = Description of Artificial Sequence: Synthetic
               polypeptide
source             1..123
mol_type = protein
organism = synthetic construct

SEQUENCE: 57
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSSIHWVRQA PGKGLEWVAS ISSYSSSTYY 60

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ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYP YGWWGGSGL DYWGQGTLVT 120
VSS 123

SEQ ID NO: 58 moltype = AA length = 111
FEATURE Location/Qualifiers
REGION 1..111
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..111
mol_type = protein
organism = synthetic construct

SEQUENCE: 58
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ FDFQYLITFG QGTKVEIKRT V 111

SEQ ID NO: 59 moltype = AA length = 117
FEATURE Location/Qualifiers
REGION 1..117
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..117
mol_type = protein
organism = synthetic construct

SEQUENCE: 59
EVQLVESGGG LVQPGGSLRL SCAASGFTVY YSSIHWRQQA PGKGLEWVAS IYSYYGSTYY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGE KWALDYWGQG TLTVSS 117

SEQ ID NO: 60 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 60
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ YMYYQPLITF GQGTKVEIKR TV 112

SEQ ID NO: 61 moltype = AA length = 121
FEATURE Location/Qualifiers
REGION 1..121
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..121
mol_type = protein
organism = synthetic construct

SEQUENCE: 61
EVQLVESGGG LVQPGGSLRL SCAASGFTVY YSSIHWRQQA PGKGLEWVAS ISSSSGSTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCAREP YNYNWYGMDS WGQGTLVTVS 120
S 121

SEQ ID NO: 62 moltype = AA length = 110
FEATURE Location/Qualifiers
REGION 1..110
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 62
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SLWWPITFGQ GTKVEIKRTV 110

SEQ ID NO: 63 moltype = AA length = 121
FEATURE Location/Qualifiers
REGION 1..121
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..121
mol_type = protein
organism = synthetic construct

SEQUENCE: 63
EVQLVESGGG LVQPGGSLRL SCAASGFTVS SSSIHWRQQA PGKGLEWVAS IYSYSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARHG SYGSWWALDY WGQGTLVTVS 120
S 121

SEQ ID NO: 64 moltype = AA length = 110
FEATURE Location/Qualifiers

-continued

REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 64
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ YFYFPITFGQ GTKVEIKRTV 110

SEQ ID NO: 65 moltype = AA length = 128
 FEATURE Location/Qualifiers
 REGION 1..128
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..128
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 65
 EVQLVESGGG LVQPGGSLRL SCAASGFTFY SSSIHWVRQA PGKGLEWVAS ISSYYGSTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARAS YYSGYGSSYP YYMGLDYWGQ 120
 GTLTVTVSS 128

SEQ ID NO: 66 moltype = AA length = 112
 FEATURE Location/Qualifiers
 REGION 1..112
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..112
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 66
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ GSYRNPLTF GQGTKVEIKR TV 112

SEQ ID NO: 67 moltype = AA length = 119
 FEATURE Location/Qualifiers
 REGION 1..119
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..119
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 67
 EVQLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWVRQA PGKGLEWVAS ISSSSGYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARMM WSHYAMDYWG QGTLTVSS 119

SEQ ID NO: 68 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 68
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 69 moltype = AA length = 123
 FEATURE Location/Qualifiers
 REGION 1..123
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..123
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 69
 EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSSIHWVRQA PGKGLEWVAY ISSYSGYTYY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYW YGHYHSYFGL DYWGQGTLVT 120
 VSS 123

SEQ ID NO: 70 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide

source 1..110
 mol_type = protein

-continued

organism = synthetic construct

SEQUENCE: 70
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 71 moltype = AA length = 122
FEATURE Location/Qualifiers
REGION 1..122
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..122
mol_type = protein
organism = synthetic construct

SEQUENCE: 71
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSSIHWRQPA PGKGLEWVAS ISSYSGSTYY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYP YGSHVYTGLD YWGQGTLTV 120
SS 122

SEQ ID NO: 72 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 72
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ WNWADYLVFQ GQGTKVEIKR TV 112

SEQ ID NO: 73 moltype = AA length = 124
FEATURE Location/Qualifiers
REGION 1..124
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..124
mol_type = protein
organism = synthetic construct

SEQUENCE: 73
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSSIHWRQPA PGKGLEWVAS IYSSSGSTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARVY SSRYWGWGVA FDYWGQGTLV 120
TVSS 124

SEQ ID NO: 74 moltype = AA length = 111
FEATURE Location/Qualifiers
REGION 1..111
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..111
mol_type = protein
organism = synthetic construct

SEQUENCE: 74
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ YYWYSLITFG QG GTKVEIKRTV 111

SEQ ID NO: 75 moltype = AA length = 128
FEATURE Location/Qualifiers
REGION 1..128
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..128
mol_type = protein
organism = synthetic construct

SEQUENCE: 75
EVQLVESGGG LVQPGGSLRL SCAASGFTVY SSSIHWRQPA PGKGLEWVAY IYSSSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARRS FPQWYNGSYT PWPAMDYWGQ 120
GTLVTVSS 128

SEQ ID NO: 76 moltype = AA length = 110
FEATURE Location/Qualifiers
REGION 1..110
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 76
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ YMWWPVTFGQ GTKVEIKRTV 110

-continued

SEQ ID NO: 77 moltype = AA length = 121
 FEATURE Location/Qualifiers
 REGION 1..121
 note = Description of Artificial Sequence: Synthetic polypeptide
 source 1..121
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 77
 EVQLVESGGG LVQPGGSLRL SCAASGFTVS SSSIHWRQQA PGKGLEWVAS IYSYSSYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARPF YWGERYALDY WGQGTLVTVS 120
 S 121

SEQ ID NO: 78 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 78
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SYSTLVTFGQ GTKVEIKRTV 110

SEQ ID NO: 79 moltype = AA length = 123
 FEATURE Location/Qualifiers
 REGION 1..123
 note = Description of Artificial Sequence: Synthetic polypeptide
 source 1..123
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 79
 EVQLVESGGG LVQPGGSLRL SCAASGFTFY SSSIHWRQQA PGKGLEWVAS IYSSYGSTS 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARIY GWSYQGWAGM DYWGQGTLVT 120
 VSS 123

SEQ ID NO: 80 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 80
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 81 moltype = AA length = 124
 FEATURE Location/Qualifiers
 REGION 1..124
 note = Description of Artificial Sequence: Synthetic polypeptide
 source 1..124
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 81
 EVQLVESGGG LVQPGGSLRL SCAASGFTIS YSSIHWRQQA PGKGLEWVAS IYPYYGSTS 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGG DYYWGWWVA MDYWGQGTLV 120
 TVSS 124

SEQ ID NO: 82 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic polypeptide
 VARIANT 76
 note = MOD_RES - Any amino acid
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 82
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLTIXSLQP EDFATYYCQK SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 83 moltype = AA length = 120

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FEATURE Location/Qualifiers
REGION 1..120
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..120
mol_type = protein
organism = synthetic construct

SEQUENCE: 83
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSYIHWRQA PGKGLEWVAS ISSSYGSTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARMY YYTYPGMDYW GQGTLVTSS 120

SEQ ID NO: 84 moltype = AA length = 110
FEATURE Location/Qualifiers
REGION 1..110
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 84
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTSSLQF EDFATYYCQQ GSSYLLTFGQ GTKVEIKRTV 110

SEQ ID NO: 85 moltype = AA length = 124
FEATURE Location/Qualifiers
REGION 1..124
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..124
mol_type = protein
organism = synthetic construct

SEQUENCE: 85
EVOLVESGGG LVQPGGSLRL SCAASGFTIY SYSIHWRQA PGKGLEWVAS ISPSSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYH YGGWSHYMSG MDYWGQGTLV 120
TVSS 124

SEQ ID NO: 86 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 86
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTSSLQF EDFATYYCQQ NYYYHKLITF GQGTKVEIKR TV 112

SEQ ID NO: 87 moltype = AA length = 117
FEATURE Location/Qualifiers
REGION 1..117
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..117
mol_type = protein
organism = synthetic construct

SEQUENCE: 87
EVOLVESGGG LVQPGGSLRL SCAASGFTFS YSSIHWRQA PGKGLEWVAS ISSSYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCAGR YGGMDYWGQG TLTVSS 117

SEQ ID NO: 88 moltype = AA length = 112
FEATURE Location/Qualifiers
REGION 1..112
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 88
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTSSLQF EDFATYYCQQ LSYVYKLITF GQGTKVEIKR TV 112

SEQ ID NO: 89 moltype = AA length = 117
FEATURE Location/Qualifiers
REGION 1..117
note = Description of Artificial Sequence: Synthetic
polypeptide
source 1..117
mol_type = protein

-continued

SEQUENCE: 89
 EVOLVESGGG LVQPGGSLRL SCAASGFTFY SSSIHWVRQA PGKGLEWVAS ISSSYGYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGW YKAMDYWGQG TLTVSS 117

SEQ ID NO: 90 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 90
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAQYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFGSGRSRGTD FTLTISLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 91 moltype = AA length = 120
 FEATURE Location/Qualifiers
 REGION 1..120
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..120
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 91
 EVOLVESGGG LVQPGGSLRL SCAASGFTFS YSSIHWVRQA PGKGLEWVAS ISSSYGSTYY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARMY YYYYPGIDYW GQGTLTVSS 120

SEQ ID NO: 92 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 92
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAQYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFGSGRSRGTD FTLTISLQP EDLATYYCQQ YYFPITFGQ GTKVEIKRTV 110

SEQ ID NO: 93 moltype = AA length = 125
 FEATURE Location/Qualifiers
 REGION 1..125
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..125
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 93
 EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWVRQA PGKGLEWVAS ISPYYGYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSP YYWYQYFYGW GLDYWGQGTL 120
 VTVSS 125

SEQ ID NO: 94 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 94
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAQYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFGSGRSRGTD FTLTISLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 95 moltype = AA length = 122
 FEATURE Location/Qualifiers
 REGION 1..122
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..122
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 95
 EVOLVESGGG LVQPGGSLRL SCAASGFTFS YSSIHWVRQA PGKGLEWVAS ISSSSGTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSP YWWNYMSAMD YWGQGTLTV 120
 SS 122

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SEQ ID NO: 96 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 96
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTTLTISSLQP EDFATYYCQQ GWWWPFTFGQ GTKVEIKRTV 110

SEQ ID NO: 97 moltype = AA length = 125
 FEATURE Location/Qualifiers
 REGION 1..125
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..125
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 97
 EVQLVESGGG LVQPGGSLRL SCAASGFTVS SYSIHWVRQA PGKGLEWVAS ISPYYGYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWS WQYYSGHSSW GLDYWGQGTL 120
 VTVSS 125

SEQ ID NO: 98 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 98
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTTLTISSLQP EDFATYYCQQ SWYFPLTFQG GTKVEIKRTV 110

SEQ ID NO: 99 moltype = AA length = 125
 FEATURE Location/Qualifiers
 REGION 1..125
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..125
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 99
 EVQLVESGGG LVQPGGSLRL SCAASGFTVS SSSIHWVRQA PGKGLEWVAS IYSYYGYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWY NEYYHDYYWD AMDYWGQGTL 120
 VTVSS 125

SEQ ID NO: 100 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 100
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTTLTISSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 101 moltype = AA length = 120
 FEATURE Location/Qualifiers
 REGION 1..120
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..120
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 101
 EVQLVESGGG LVQPGGSLRL SCAASGFTLY YSSIHWRQA PGKGLEWVAS ISSSSGSTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWM YWWSFALDYW GQGTLVTVSS 120

SEQ ID NO: 102 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide

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source          1..110
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 102
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTSSLQP EDFATYYCQQ SYLWPITFGQ GTKVEIKRTV           110

SEQ ID NO: 103      moltype = AA  length = 122
FEATURE          Location/Qualifiers
REGION           1..122
note = Description of Artificial Sequence: Synthetic
               polypeptide
source          1..122
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 103
EVOLVESGGG LVQPGGSLRL SCAASGFTVS SSSIHWRQA PGKGLEWVAS IYSYYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWQ YHYNWYGMQ YWGQGLTVV 120
SS                         122

SEQ ID NO: 104      moltype = AA  length = 110
FEATURE          Location/Qualifiers
REGION           1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source          1..110
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 104
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTSSLQP EDFATYYCQQ YPMSLITFGQ GTKVEIKRTV           110

SEQ ID NO: 105      moltype = AA  length = 119
FEATURE          Location/Qualifiers
REGION           1..119
note = Description of Artificial Sequence: Synthetic
               polypeptide
source          1..119
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 105
EVOLVESGGG LVQPGGSLRL SCAASGFTVS YSSIHWRQA PGKGLEWVAS ISPYSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGY DYYAGLDYWQ QGTLTVSS 119

SEQ ID NO: 106      moltype = AA  length = 110
FEATURE          Location/Qualifiers
REGION           1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source          1..110
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 106
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFSGSRSGTD FTLTSSLQP EDFATYYCQQ YYFPITFGQ GTKVEIKRTV           110

SEQ ID NO: 107      moltype = AA  length = 127
FEATURE          Location/Qualifiers
REGION           1..127
note = Description of Artificial Sequence: Synthetic
               polypeptide
source          1..127
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 107
EVOLVESGGG LVQPGGSLRL SCAASGFTVS YYSIHWRQA PGKGLEWVAS ISPYSGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWE SEYSGTYEDY WAGMDYWGQG 120
TLTVSS                         127

SEQ ID NO: 108      moltype = AA  length = 110
FEATURE          Location/Qualifiers
REGION           1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source          1..110
               mol_type = protein
               organism = synthetic construct
SEQUENCE: 108
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60

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RFSGSRSGTD FTLLTISLQP EDFATYYCQQ YMWWPITFGQ GTKVEIKRTV 110

SEQ ID NO: 109 moltype = AA length = 121
 FEATURE Location/Qualifiers
 REGION 1..121
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..121
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 109
 EVQLVESGGG LVQPGGSLRL SCAASGFTIS YSSIHWVRQA PGKGLEWVAS ISSSYSYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARTG YWQGYLALDY WGQGTLVTVS 120
 S 121

SEQ ID NO: 110 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 110
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLLTISLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 111 moltype = AA length = 123
 FEATURE Location/Qualifiers
 REGION 1..123
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..123
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 111
 EVQLVESGGG LVQPGGSLRL SCAASGFTIS YSSIHWVRQA PGKGLEWVAS ISSSSGTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARTY YYWNSTPAM DYWGQGTLVT 120
 VSS 123

SEQ ID NO: 112 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 112
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLLTISLQP EDFATYYCQQ SYGYPVTFQ GTKVEIKRTV 110

SEQ ID NO: 113 moltype = AA length = 129
 FEATURE Location/Qualifiers
 REGION 1..129
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..129
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 113
 EVQLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHHWVRQA PGKGLEWVAS ISSSYGYTSY 60
 ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWY NSSWYYSNWW YKGFGMDYWG 120
 QGTLVTVSS 129

SEQ ID NO: 114 moltype = AA length = 110
 FEATURE Location/Qualifiers
 REGION 1..110
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..110
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 114
 DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
 RFSGSRSGTD FTLLTISLQP EDFATYYCQQ YYSSLFTFGQ GTKVEIKRTV 110

SEQ ID NO: 115 moltype = AA length = 124
 FEATURE Location/Qualifiers

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mol_type = protein
organism = synthetic construct

SEQUENCE: 121
EVQLVESGGG LVQPGGSLRL SCAASGFTVS SSSIHWRQQA PGKGLEWVAS IYSYSGSTYY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARQY SSSYYWPQGM DYWGQGTLVT 120
VSS 123

SEQ ID NO: 122      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = Description of Artificial Sequence: Synthetic
               polypeptide
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 122
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGSRSGTD FTLTSSLQP EDFATYYCQQ SYWWKSLVTF GQGTKVEIKR TV 112

SEQ ID NO: 123      moltype = AA length = 126
FEATURE          Location/Qualifiers
REGION           1..126
note = Description of Artificial Sequence: Synthetic
               polypeptide
source            1..126
mol_type = protein
organism = synthetic construct

SEQUENCE: 123
EVOLVESGGG LVQPGGSLRL SCAASGFTLS SSSIHWRQQA PGKGLEWVAS ISSYYGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARMH YSWQEYYSYD WGMDYWGQGT 120
LTVSS 126

SEQ ID NO: 124      moltype = AA length = 110
FEATURE          Location/Qualifiers
REGION           1..110
note = Description of Artificial Sequence: Synthetic
               polypeptide
source            1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 124
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGSRSGTD FTLTSSLQP EDFATYYCQQ PYYPLITFGQ GTKVEIKRTV 110

SEQ ID NO: 125      moltype = AA length = 120
FEATURE          Location/Qualifiers
REGION           1..120
note = Description of Artificial Sequence: Synthetic
               polypeptide
source            1..120
mol_type = protein
organism = synthetic construct

SEQUENCE: 125
EVOLVESGGG LVQPGGSLRL SCAASGFTIS YSSIHWRQQA PGKGLEWVAS IYPSYGSTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWQ GYYQPALDYW GQGTLTVSS 120

SEQ ID NO: 126      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
note = Description of Artificial Sequence: Synthetic
               polypeptide
source            1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 126
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGSRSGTD FTLTSSLQP EDFATYYCQQ SSKYYYPTF GQGTKVEIKR TV 112

SEQ ID NO: 127      moltype = AA length = 121
FEATURE          Location/Qualifiers
REGION           1..121
note = Description of Artificial Sequence: Synthetic
               polypeptide
source            1..121
mol_type = protein
organism = synthetic construct

SEQUENCE: 127
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSSIHWRQQA PGKGLEWVAS ISPYYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARWG YGWWYGLDY WGQGTLTVS 120

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S	121
SEQ ID NO: 128	moltype = AA length = 111
FEATURE	Location/Qualifiers
REGION	1..111
	note = Description of Artificial Sequence: Synthetic
	polypeptide
source	1..111
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 128	
DIQMTQSPSS LSASVGRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60	
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ GHDMNPVTFG QG GTKVEIKRT V 111	
SEQ ID NO: 129	moltype = AA length = 120
FEATURE	Location/Qualifiers
REGION	1..120
	note = Description of Artificial Sequence: Synthetic
	polypeptide
source	1..120
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 129	
EVQLVESGGG LVQPGGSLRL SCAASGFTLS SSSIHWVRQA PGKGLEWVAS IYSSYGYTYY 60	
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYY YYWYGGMDYW GQGTLTVSS 120	
SEQ ID NO: 130	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = Description of Artificial Sequence: Synthetic
	polypeptide
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 130	
DIQMTQSPSS LSASVGRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60	
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ SWMSDSLITF GQGTKVEIKR TV 112	
SEQ ID NO: 131	moltype = AA length = 123
FEATURE	Location/Qualifiers
REGION	1..123
	note = Description of Artificial Sequence: Synthetic
	polypeptide
source	1..123
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 131	
EVQLVESGGG LVQPGGSLRL SCAASGFTFS YSSIHWVRQA PGKGLEWVAS IYPSSGYTSY 60	
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARGW WYWMAWDYAM DYWGQGTLVT 120	
VSS	123
SEQ ID NO: 132	moltype = AA length = 112
FEATURE	Location/Qualifiers
REGION	1..112
	note = Description of Artificial Sequence: Synthetic
	polypeptide
source	1..112
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 132	
DIQMTQSPSS LSASVGRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60	
RFSGSRSGTD FTLTISSLQP EDFATYYCQQ MQYSGWLITF GQGTKVEIKR TV 112	
SEQ ID NO: 133	moltype = AA length = 130
FEATURE	Location/Qualifiers
REGION	1..130
	note = Description of Artificial Sequence: Synthetic
	polypeptide
source	1..130
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 133	
EVQLVESGGG LVQPGGSLRL SCAASGFTIS SSSIHWVRQA PGKGLEWVAS ISSYYGYTSY 60	
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYY SYSSGYGYD YFDWGAMDYW 120	
GQGTLTVSS	130
SEQ ID NO: 134	moltype = AA length = 110
FEATURE	Location/Qualifiers
REGION	1..110

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source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 134
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTISSSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 135      moltype = AA length = 129
FEATURE          Location/Qualifiers
REGION           1..129
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..129
mol_type = protein
organism = synthetic construct

SEQUENCE: 135
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWRQQA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYY GYWGGYWGW WYSKALDYWG 120
QGTLTVSS                                129

SEQ ID NO: 136      moltype = AA length = 112
FEATURE          Location/Qualifiers
REGION           1..112
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..112
mol_type = protein
organism = synthetic construct

SEQUENCE: 136
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTISSSLQP EDFATYYCQQ YDWNYYLVTQ GQGTKEIKR TV 112

SEQ ID NO: 137      moltype = AA length = 126
FEATURE          Location/Qualifiers
REGION           1..126
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..126
mol_type = protein
organism = synthetic construct

SEQUENCE: 137
EVOLVESGGG LVQPGGSLRL SCAASGFTIY SSSIHWRQQA PGKGLEWVAS ISSYYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARYQ YYGSLYYSQQ WAMDYWGQGT 120
LTVSS                                126

SEQ ID NO: 138      moltype = AA length = 110
FEATURE          Location/Qualifiers
REGION           1..110
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 138
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAWYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRSGTD FTLTISSSLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 139      moltype = AA length = 129
FEATURE          Location/Qualifiers
REGION           1..129
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..129
mol_type = protein
organism = synthetic construct

SEQUENCE: 139
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWRQQA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARSP SSPYFMSWGW YWQYGINYWG 120
QGTLTVSS                                129

SEQ ID NO: 140      moltype = AA length = 111
FEATURE          Location/Qualifiers
REGION           1..111
source          note = Description of Artificial Sequence: Synthetic
                polypeptide
1..111
mol_type = protein

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organism = synthetic construct

SEQUENCE: 140
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAQYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRGTD FTLTISLQP EDFATYYCQQ SSWGGLVTFG QGTKVEIKRT V 111

SEQ ID NO: 141
FEATURE moltype = AA length = 121
REGION Location/Qualifiers
1..121
note = Description of Artificial Sequence: Synthetic polypeptide
source 1..121
mol_type = protein
organism = synthetic construct

SEQUENCE: 141
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSYIHWRQA PGKGLEWVAS ISPSYGYTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARDM YEWHHWAIDY WGQGTLVTVS 120
S 121

SEQ ID NO: 142
FEATURE moltype = AA length = 110
REGION Location/Qualifiers
1..110
note = Description of Artificial Sequence: Synthetic polypeptide
source 1..110
mol_type = protein
organism = synthetic construct

SEQUENCE: 142
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAQYQQKP GKAPKLLIYS ASSLYSGVPS 60
RFGSGRSRGTD FTLTISLQP EDFATYYCQQ SSSSLITFGQ GTKVEIKRTV 110

SEQ ID NO: 143
FEATURE moltype = AA length = 129
REGION Location/Qualifiers
1..129
note = Description of Artificial Sequence: Synthetic polypeptide
source 1..129
mol_type = protein
organism = synthetic construct

SEQUENCE: 143
EVOLVESGGG LVQPGGSLRL SCAASGFTFS SSSIHWRQA PGKGLEWVAS ISSSSGTSY 60
ADSVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARCG HYLYYWGWG YWSAALDYWG 120
QGTTLVTVSS 129

SEQ ID NO: 144
FEATURE moltype = AA length = 514
REGION Location/Qualifiers
1..514
note = Description of Artificial Sequence: Synthetic polypeptide
source 1..514
mol_type = protein
organism = synthetic construct

SEQUENCE: 144
DIVRSIDIQMT QSPSSLSASV GDRVITICRA SQSVSSAVW YQQKPGKAPL LLIYSASSLY 60
SGVPSRFSGS RSGTDFTLTI SSLQPEDFAT YYCQQISVY SLITFGQGTK VEIKGGGSEVQ 120
LQLQSGPSEL VKPGASMKIS CKASGYSTFG YTMMWVKQSH GKNLWMLINP NPYKVSTYNQ 180
QFKDKATLTV DVKSSSTAYM ELLSLTSEDS AVYYCARSGY YGDSDWYFDV WGGTTLTVS 240
SGGGGSGGGG SGGGGSDIQM QTQTSLSAS LGDRVTISCR ASQDIRNYLNW YWQKPDGTGV 300
KLLIYTTSRL HSGVPKFSD SGSGTDYSLT NLNEQEBIA TYFCQGQNTL PWTFAGGTKL 360
EIKGGGSEVQ OLVESGGGLV QPGGSRLSC ISANGFTFSYY SIHWRQAPG KGLEWVASIY 420
SSYGYTSYAD SVKGRTFISA DTSKNTAYLQ MNSLRAEDTA VYYCARGGWY PAMDYWGQGT 480
LTVVSSLEGG GGLNDIFEAQ KIEWHESRHH HHHH 514

SEQ ID NO: 145
FEATURE moltype = AA length = 517
REGION Location/Qualifiers
1..517
note = Description of Artificial Sequence: Synthetic polypeptide
source 1..517
mol_type = protein
organism = synthetic construct

SEQUENCE: 145
DIVRSIDIQMT QSPSSLSASV GDRVITICRA SQSVSSAVW YQQKPGKAPL LLIYSASSLY 60
SGVPSRFSGS RSGTDFTLTI SSLQPEDFAT YYCQQSSSSL ITFGQGTKVE IKGGGSEVQ 120
LQLQSGPSELVK PGASMKISCK ASGYSTFGYT MNWVKQSHGK NLEWMLINP YKGVSTYNQ 180
FKDKATLTV DKSSTAYMEL LSLTSEDSAV YYCARSGYYG DSDWYFDVWG QGTTLTVSSG 240
GGGGGGGGGG GGGSDIQMTQ TTSSLASLGL DRVTICRAS QDIRNYLNW YQKPDGTVKL 300
LIYYTSRLNH GPVSKFSGSG SGDYSLTIS NLEQEDIATY FCQGQNTLPW TFAGGTKLEI 360
GGGGGSEVQL VESGGGLVQP GGRLRSLSCA SGFTTISSSI HWVRQAPGK LEVWAVY 420
SGYTYYADSV KGRFTISADT SKNTAYLQMN SLRAEDTAVY YCARYWYGHY HSYPGLDYG 480

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mol_type = protein
organism = synthetic construct

SEQUENCE: 150
EVQLVESGGG LVQPGGSLRL SCAASGFTFS DYSIHWVRQA PGKGLEWVAS ISSSSGSTS 60
ADSVKGRTI SADTSKNTAY LQMNSLRAED TAVYYCARGG WYPAMDYWGQ GTLVTVSSAS 120
TKGPSVFPPLA PSSKSTSGGT AALGCLVEDY FPEPVTVSWN SGALTSGVHT FPAVLQSSGL 180
YSLSSVVTVP SSSLGTQTYI CNVNHKPSNT KVDKEKVEPKS CDGGGGGGGG GSQAVTQEP 240
SLTVSPGGT TLTCGSSTGA VTTSNYANWV QEKPQAFRQ LIGGTNKRAP GTPARFSGSL 300
LGKAAALTLS GAQPEDAEAY YCALWYSNLW VFGGGTLITV LSSASTKGPS VFPLAPSSKS 360
TSGGTAAALGC LVKDYFPPEPV TVSWNSGALT SGVHTFPAVL QSSGLYSLSS VVTVPSSSLG 420
TQTYICVNHN KPSNTKVDKK VEPKSCDKTH TCPPCPAPEA AGGPSVFLFP PKPKDTLMIS 480
RTPEVTCVVV DVSHEDPEVK FNWYVDGVEV HNAKTKPREE QYNSTYRVVS VLTVLHQDWL 540
NGKEYKCVS NKALGAPIEK TISAKGQPR EPOVYTLPPC RDELTKNQVS LWCLVKGFYP 600
SDIAVEWSN QOPENNYKTT PPVLDSDGSF FLYSKLTVDK SRWQQGNVFS CSVMHEALHN 660
HYTQKSLSLS P 671

SEQ ID NO: 151      moltype = AA length = 446
FEATURE          Location/Qualifiers
REGION           1..446
                  note = Description of Artificial Sequence: Synthetic
                  polypeptide
source            1..446
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 151
EVQLVESGGG LVQPGGSLRL SCAASGFTFS DYSIHWVRQA PGKGLEWVAS ISSSSGSTS 60
ADSVKGRTI SADTSKNTAY LQMNSLRAED TAVYYCARGG WYPAMDYWGQ GTLVTVSSAS 120
TKGPSVFPPLA PSSKSTSGGT AALGCLVEDY FPEPVTVSWN SGALTSGVHT FPAVLQSSGL 180
YSLSSVVTVP SSSLGTQTYI CNVNHKPSNT KVDKEKVEPKS CDKTHTCPG PAPEAAGGPS 240
VFVLPFPKPKD TLMSIRTPEV TCVVVDVSH DPEVKFNWYV DGVEVHNAKT KPREEQYNST 300
YRVRVSVLTVL HQDWLNGKEY KCKVSNKALG APIEKTISKA KGQPREPQVC TLPPSRDELT 360
KNQVSLSCAV KGFYPSDIAV EWESNGQOPEN NYKTPPPVLD SDGSFFLVSK LTVDKSRWQQ 420
GNVFSCSVMH EALHNHYTQK SLSLSP 446

SEQ ID NO: 152      moltype = AA length = 216
FEATURE          Location/Qualifiers
REGION           1..216
                  note = Description of Artificial Sequence: Synthetic
                  polypeptide
source            1..216
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 152
DIQMTQSPSS LSASVGDRVT ITCRASQSVS SAVAQYQQKP GKAPKLLIYS ASSLYSGVPS 60
RIFSGSRSGTD FTLLTISSLQP EDFATYYCQQ ISYVKKLITF GQGTKVEIKR TVAAPSVFIF 120
PPSDRKLSKG TASVVCLLNN FYPREAKVQW KVNDALQSGN SQESVTEQDS KDSTYSLSS 180
LTLSKADYEK HKVYACEVTN QGLSSPVTKS FNRGEC 216

SEQ ID NO: 153      moltype = AA length = 8
FEATURE          Location/Qualifiers
REGION           1..8
                  note = Description of Artificial Sequence: Synthetic peptide
source            1..8
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 153
ISYVKKLI 8

SEQ ID NO: 154      moltype = length =
SEQUENCE: 154
000

SEQ ID NO: 155      moltype = AA length = 6
FEATURE          Location/Qualifiers
REGION           1..6
                  note = Description of Artificial Sequence: Synthetic peptide
source            1..6
                  mol_type = protein
                  organism = synthetic construct
SEQUENCE: 155
GGWYPA 6

SEQ ID NO: 156      moltype = length =
SEQUENCE: 156
000

SEQ ID NO: 157      moltype = AA length = 6
FEATURE          Location/Qualifiers
REGION           1..6

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source note = Description of Artificial Sequence: Synthetic peptide
 1..6
 mol_type = protein
 organism = synthetic construct
 SEQUENCE: 157
 YSYWPI 6

SEQ ID NO: 158 moltype = length =
 SEQUENCE: 158
 000

SEQ ID NO: 159 moltype = AA length = 7
 FEATURE Location/Qualifiers
 REGION 1..7
 note = Description of Artificial Sequence: Synthetic peptide
 1..7
 source mol_type = protein
 organism = synthetic construct
 SEQUENCE: 159 EYVTMAL 7

SEQ ID NO: 160 moltype = length =
 SEQUENCE: 160
 000

SEQ ID NO: 161 moltype = AA length = 488
 FEATURE Location/Qualifiers
 REGION 1..488
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 1..488
 source mol_type = protein
 organism = synthetic construct
 SEQUENCE: 161 DIVRSIDIQMT QSPSSLSASV GDRVITCRA SQSVSSAVAW YQQKPGKAPK LLIYSASSLY 60
 SGVPSRSGS RSGTDFLTG SSLQPEDFAT YYCQQISVYY SLITFGQGTK VEIKGGGSE 120
 VQLQQSGPEL VKPGASMKIS CKASGYSFVG YTMMNVVKOSH GKNLEWMGLI NPYKGVSTYN 180
 QKFKDATALT VDKSSSTAYM ELLSLTSEDS AVYYCARSGY YGDSDWYFDV WGGQGTTLTVS 240
 SGGGGSGGGG SGGGGSDIQM TQTTSSLAS LGDRVTISCR ASQDIRNYLN WYQQKPDGTV 300
 KLLIYYTSRL HSGVPSKFSG SGSGTDYSLT ISNLEQEDIA TYFCQQGNTL PWTFAAGGTLK 360
 EIKGGGGSEV QLVESGGGLV QPGGSLRLSC AASGFTFSYY SIHWVRQAPG KGLEWVASIY 420
 SSYGYTSYAD SVKGFRFTA DTSKNTAYLQ MNSLRAEDTA VYYCARGGWY PAMDYWGQGT 480
 LTVSSLE 488

SEQ ID NO: 162 moltype = AA length = 491
 FEATURE Location/Qualifiers
 REGION 1..491
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 1..491
 source mol_type = protein
 organism = synthetic construct
 SEQUENCE: 162 DIVRSIDIQMT QSPSSLSASV GDRVITCRA SQSVSSAVAW YQQKPGKAPK LLIYSASSLY 60
 SGVPSRSGS RSGTDFLTG SSLQPEDFAT YYCQQSSSSL ITFGQGTKVE IKGGGSEVQ 120
 LQQSGPELVK PGASMKISCK ASGYSGFTGYT MNWVKQSHGK NLEWMGLINP YKGVSTYNQK 180
 FKDKATALTVD KSSSTAYMEL LSLTSEDSAV YYCARSGYYG DSDWYFDVWG QGTTLTVSSG 240
 GGGGGGGGG GGGSDIQMTQ TTSSLASLG DRVТИSCRAS QDIRNYLNWY QQKPDGTVKL 300
 LIYYTSRLHS GPVSKFSGSG SGTDYSLTIS NLEQEDIATY FCQQGNTLPW TFAGGKLEI 360
 KGGGGSEVQL VESGGGLVQP GGSLRLSCAA SGFTISSSSII HWVRQAPGKG LEWVAYISSY 420
 SGYTYYADSV KGRFTISADT SKNTAYLQM SLRAEDTAVY YCARYWYGHY HSYFGLDYWG 480
 QTGLTVSSLE 491

SEQ ID NO: 163 moltype = AA length = 486
 FEATURE Location/Qualifiers
 REGION 1..486
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 1..486
 source mol_type = protein
 organism = synthetic construct
 SEQUENCE: 163 DIVRSIDIQMT QSPSSLSASV GDRVITCRA SQSVSSAVAW YQQKPGKAPK LLIYSASSLY 60
 SGVPSRSGS RSGTDFLTG SSLQPEDFAT YYCQQYSWY ITFGQGTKVE IKGGGSEVQ 120
 LQQSGPELVK PGASMKISCK ASGYSGFTGYT MNWVKQSHGK NLEWMGLINP YKGVSTYNQK 180
 FKDKATALTVD KSSSTAYMEL LSLTSEDSAV YYCARSGYYG DSDWYFDVWG QGTTLTVSSG 240
 GGGGGGGGG GGGSDIQMTQ TTSSLASLG DRVТИSCRAS QDIRNYLNWY QQKPDGTVKL 300
 LIYYTSRLHS GPVSKFSGSG SGTDYSLTIS NLEQEDIATY FCQQGNTLPW TFAGGKLEI 360
 KGGGGSEVQL VESGGGLVQP GGSLRLSCAA SGFTISSSSYI HWVRQAPGKG LEWVAYISPS 420

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YGSTSYADSV KGRFTISADT SKNTAYLQMN SLRAEDTAVY YCAREYVTMA LDYWQGQTLV	480
TVSSLE	486

SEQ ID NO: 164 moltype = AA length = 490
 FEATURE Location/Qualifiers
 REGION 1..490
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..490
 mol_type = protein
 organism = synthetic construct
 SEQUENCE: 164
 DIVRSDIQMT QSPSSLSASV GDRVTITCRA SQSVSSAVAW YQQKPGKAPK LLIYSASSLY 60
 SGVPSRFSGS RSGTDFTLTI SSLQPEDFAT YYCQQSSWGG LVTFGQGCKV EIKGGGSEV 120
 QLQQSGPELV KPGASMKIS KASGYSFTGY TMMWVKQSHG KNLEWMGLIN PYKGVSTYNQ 180
 KFKDKATLTV DKSSSTAYME LLSLTSEDSA VYYCARSGY GDSDWYFDVV GQGTTLTVSS 240
 GGGGSGGGGS GGGGSDIQM QTTSLSLSDA GDRVTISCR A SODIRNYLNW YQQKPDGTWK 300
 LLIYYTTSRLH SGVPSKFSGS GSGTDYSLTI SNLEQEDIAT YFCQQGNTLP WTFAGGTKLE 360
 IKGGGGSEVO LIVESGGGLVQ PGGSRLSCA ASGFTFSSSY IHWVRQAPGK GLEWVAVISP 420
 SYGYTSYADS VKGRFTISAD TSKNTAYLQM NSLAEDTAV YYCARDMYEW WHWAIDYWQ 480
 GTLVTVSLE 490

SEQ ID NO: 165 moltype = AA length = 488
 FEATURE Location/Qualifiers
 REGION 1..488
 note = Description of Artificial Sequence: Synthetic
 polypeptide
 source 1..488
 mol_type = protein
 organism = synthetic construct
 SEQUENCE: 165
 DIVRSDIQMT QSPSSLSASV GDRVTITCRA SQSVSSAVAW YQQKPGKAPK LLIYSASSLY 60
 SGVPSRFSGS RSGTDFTLTI SSLQPEDFAT YYCQQISYVK KLITFGQGCKV VEIKGGGSE 120
 VQLQQSGPELV KPGASMKIS CKASGYSFTGY TMMWVKQSHG KNLEWMGLIN NPYKGVSTYN 180
 QKFKDKATLTV DVKSSSTAYM ELLSLTSEDS AVYYCARSGY YGDSDWYFDVV WGQGTTLTVS 240
 SGGGGSGGGG SGGGGSDIQM TQTTSLSAS LGDRVTISCR ASQDIRNYLN WYQQKPDGTWV 300
 KLLIYYTTSRL HSGVPSKFSGS SGSGTDYSLT ISNLEQEDIA TYFCQQGNTL PWTFAGGTKL 360
 EIKGGGGSEV QLVESGGGLVQ PGGSRLSCA AASGFTFSYD SIHWVRQAPG KGLEWVAVIS 420
 SSSGSTSYAD SVKGRFTISA DTSKNTAYLQ MNSLRAEDTA VYYCARGGWY PAMDYWGQGT 480
 LTVSSLE 488

SEQ ID NO: 166 moltype = AA length = 11
 FEATURE Location/Qualifiers
 source 1..11
 mol_type = protein
 organism = synthetic construct
 SEQUENCE: 166
 RASQSVSSAV A 11

SEQ ID NO: 167 moltype = AA length = 7
 FEATURE Location/Qualifiers
 source 1..7
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 167
 SASSLYS 7

SEQ ID NO: 168 moltype = AA length = 11
 FEATURE Location/Qualifiers
 source 1..11
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 168
 QQISYVKKLI T 11

SEQ ID NO: 169 moltype = AA length = 5
 FEATURE Location/Qualifiers
 source 1..5
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 169
 DYSIH 5

SEQ ID NO: 170 moltype = AA length = 17
 FEATURE Location/Qualifiers
 source 1..17
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 170

-continued

SISSSSGSTS YADSVKG	17
SEQ ID NO: 171	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 171	
GGWYPAMDY	9
SEQ ID NO: 172	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 172	
QQSYWPIT	9
SEQ ID NO: 173	moltype = AA length = 5
FEATURE	Location/Qualifiers
source	1..5
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 173	
SSYIH	5
SEQ ID NO: 174	moltype = AA length = 17
FEATURE	Location/Qualifiers
source	1..17
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 174	
YISPSYGSTS YADSVKG	17
SEQ ID NO: 175	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 175	
EYVTMALDY	9
SEQ ID NO: 176	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 176	
QQDWYWFPIT	9
SEQ ID NO: 177	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 177	
FTFSSYYIH	9
SEQ ID NO: 178	moltype = AA length = 17
FEATURE	Location/Qualifiers
source	1..17
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 178	
SISPSSGSTY YADSVKG	17
SEQ ID NO: 179	moltype = AA length = 21
FEATURE	Location/Qualifiers
source	1..21
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 179	
YGGRSYWQKQ DSYFYQHGLD Y	21
SEQ ID NO: 180	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct

-continued

SEQUENCE: 180		
QQSSSSLIT	9	
SEQ ID NO: 181	moltype = AA length = 9	
FEATURE	Location/Qualifiers	
source	1..9	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 181		
FTFSSSIH	9	
SEQ ID NO: 182	moltype = AA length = 17	
FEATURE	Location/Qualifiers	
source	1..17	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 182		
SISSYSGYTS YADSVKG	17	
SEQ ID NO: 183	moltype = AA length = 18	
FEATURE	Location/Qualifiers	
source	1..18	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 183		
SYSYSEFRYY YSGQGMDY	18	
SEQ ID NO: 184	moltype = AA length = 14	
FEATURE	Location/Qualifiers	
source	1..14	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 184		
SNYGWRWHLV GMDY	14	
SEQ ID NO: 185	moltype = AA length = 8	
FEATURE	Location/Qualifiers	
source	1..8	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 185		
TFSSSIH	8	
SEQ ID NO: 186	moltype = AA length = 14	
FEATURE	Location/Qualifiers	
source	1..14	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 186		
SPYVYYWYMV GFDY	14	
SEQ ID NO: 187	moltype = AA length = 14	
FEATURE	Location/Qualifiers	
source	1..14	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 187		
TGTSSDVGGY NYVS	14	
SEQ ID NO: 188	moltype = AA length = 7	
FEATURE	Location/Qualifiers	
source	1..7	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 188		
DVSKRPS	7	
SEQ ID NO: 189	moltype = AA length = 10	
FEATURE	Location/Qualifiers	
source	1..10	
	mol_type = protein	
	organism = synthetic construct	
SEQUENCE: 189		
GSYADTDTIV	10	
SEQ ID NO: 190	moltype = AA length = 5	
FEATURE	Location/Qualifiers	
source	1..5	
	mol_type = protein	

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	organism = synthetic construct	
SEQUENCE: 190		
TDAIT		5
SEQ ID NO: 191	moltype = AA length = 17	
FEATURE	Location/Qualifiers	
source	1..17	
	mol_type = protein	
SEQUENCE: 191	organism = synthetic construct	
GIPIPLLDSVD YAQRFQG		17
SEQ ID NO: 192	moltype = AA length = 10	
FEATURE	Location/Qualifiers	
source	1..10	
	mol_type = protein	
SEQUENCE: 192	organism = synthetic construct	
WSSVDTGLDY		10
SEQ ID NO: 193	moltype = AA length = 7	
FEATURE	Location/Qualifiers	
source	1..7	
	mol_type = protein	
SEQUENCE: 193	organism = synthetic construct	
NDPHYWG		7
SEQ ID NO: 194	moltype = AA length = 16	
FEATURE	Location/Qualifiers	
source	1..16	
	mol_type = protein	
SEQUENCE: 194	organism = synthetic construct	
STSHSGHTYF NPSLKS		16
SEQ ID NO: 195	moltype = AA length = 16	
FEATURE	Location/Qualifiers	
source	1..16	
	mol_type = protein	
SEQUENCE: 195	organism = synthetic construct	
MRYYYSGTYP VYYFDY		16
SEQ ID NO: 196	moltype = AA length = 13	
FEATURE	Location/Qualifiers	
source	1..13	
	mol_type = protein	
SEQUENCE: 196	organism = synthetic construct	
SGSSSNIGSN PVS		13
SEQ ID NO: 197	moltype = AA length = 7	
FEATURE	Location/Qualifiers	
source	1..7	
	mol_type = protein	
SEQUENCE: 197	organism = synthetic construct	
SNNQRPS		7
SEQ ID NO: 198	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 198	organism = synthetic construct	
AAWDDSLNGP V		11
SEQ ID NO: 199	moltype = AA length = 5	
FEATURE	Location/Qualifiers	
source	1..5	
	mol_type = protein	
SEQUENCE: 199	organism = synthetic construct	
RYGVSV		5
SEQ ID NO: 200	moltype = AA length = 17	
FEATURE	Location/Qualifiers	
source	1..17	

-continued

SEQUENCE: 200 GIIPMFGTAN YAQKFQG	mol_type = protein organism = synthetic construct	
		17
SEQ ID NO: 201 FEATURE source	moltype = AA length = 12 Location/Qualifiers 1..12 mol_type = protein organism = synthetic construct	
SEQUENCE: 201 GDNSAYSDAF NI		12
SEQ ID NO: 202 FEATURE source	moltype = AA length = 10 Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 202 QQSGWWSYPIT		10
SEQ ID NO: 203 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 203 SISPYYGSTS YADSVKG		17
SEQ ID NO: 204 FEATURE source	moltype = AA length = 8 Location/Qualifiers 1..8 mol_type = protein organism = synthetic construct	
SEQUENCE: 204 SSYYALDY		8
SEQ ID NO: 205 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 205 QQISYYVSLI T		11
SEQ ID NO: 206 FEATURE source	moltype = AA length = 5 Location/Qualifiers 1..5 mol_type = protein organism = synthetic construct	
SEQUENCE: 206 SYSIH		5
SEQ ID NO: 207 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 207 SIYSSYGYTS YADSVKG		17
SEQ ID NO: 208 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 208 QQYQYGYNLI T		11
SEQ ID NO: 209 FEATURE source	moltype = AA length = 5 Location/Qualifiers 1..5 mol_type = protein organism = synthetic construct	
SEQUENCE: 209 YSSIH		5
SEQ ID NO: 210 FEATURE	moltype = AA length = 17 Location/Qualifiers	

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source	1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 210 SIYSYSGSGTS YADSVKG		17
SEQ ID NO: 211 FEATURE source	moltype = AA length = 15 Location/Qualifiers 1..15 mol_type = protein organism = synthetic construct	
SEQUENCE: 211 YSYGVVGPGW RAIDY		15
SEQ ID NO: 212 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 212 QQSSSVYKLL T		11
SEQ ID NO: 213 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 213 SISSSSYTS YADSVKG		17
SEQ ID NO: 214 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 214 GGPGWYRAMD Y		11
SEQ ID NO: 215 FEATURE source	moltype = AA length = 5 Location/Qualifiers 1..5 mol_type = protein organism = synthetic construct	
SEQUENCE: 215 SSSIH		5
SEQ ID NO: 216 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 216 GYFYYGWWAM AFDY		14
SEQ ID NO: 217 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 217 QQSQWYEPLI T		11
SEQ ID NO: 218 FEATURE source	moltype = AA length = 18 Location/Qualifiers 1..18 mol_type = protein organism = synthetic construct	
SEQUENCE: 218 SYSYMSQWGW YQYSGMDY		18
SEQ ID NO: 219 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 219 QQGSYTYRLI T		11
SEQ ID NO: 220	moltype = AA length = 17	

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FEATURE source	Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 220 SISSSSGYTS YADSVKG		17
SEQ ID NO: 221 FEATURE source	moltype = AA length = 10 Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 221 YAWWAHGLDY		10
SEQ ID NO: 222 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 222 QQASYWYNLF T		11
SEQ ID NO: 223 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 223 QYSMHFPWGY GMDY		14
SEQ ID NO: 224 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 224 QQSDMPPIT		9
SEQ ID NO: 225 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 225 YIYSSSGYTS YADSVKG		17
SEQ ID NO: 226 FEATURE source	moltype = AA length = 12 Location/Qualifiers 1..12 mol_type = protein organism = synthetic construct	
SEQUENCE: 226 PVNYYYQQGAL DY		12
SEQ ID NO: 227 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 227 QQYYVFPIT		9
SEQ ID NO: 228 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 228 SISPSSGGYTS YADSVKG		17
SEQ ID NO: 229 FEATURE source	moltype = AA length = 21 Location/Qualifiers 1..21 mol_type = protein organism = synthetic construct	
SEQUENCE: 229 YHYMFNEYDKG ESKWGYYGFD Y		21

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SEQ ID NO: 230	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 230	QQSQYFPIT	9
SEQ ID NO: 231	moltype = AA length = 17 Location/Qualifiers source 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 231	SIYSYYGTS YADSVKG	17
SEQ ID NO: 232	moltype = AA length = 13 Location/Qualifiers source 1..13 mol_type = protein organism = synthetic construct	
SEQUENCE: 232	NSWSWYSGVG MDY	13
SEQ ID NO: 233	moltype = AA length = 17 Location/Qualifiers source 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 233	SISSYSSSTY YADSVKG	17
SEQ ID NO: 234	moltype = AA length = 14 Location/Qualifiers source 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 234	YPYGWGWGGS GLDY	14
SEQ ID NO: 235	moltype = AA length = 10 Location/Qualifiers source 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 235	QQFDFQYLIT	10
SEQ ID NO: 236	moltype = AA length = 17 Location/Qualifiers source 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 236	SIYSYYGSTY YADSVKG	17
SEQ ID NO: 237	moltype = AA length = 8 Location/Qualifiers source 1..8 mol_type = protein organism = synthetic construct	
SEQUENCE: 237	GEKWALDY	8
SEQ ID NO: 238	moltype = AA length = 11 Location/Qualifiers source 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 238	QQYMYYQPLI T	11
SEQ ID NO: 239	moltype = AA length = 12 Location/Qualifiers source 1..12 mol_type = protein organism = synthetic construct	
SEQUENCE: 239	EPYNYNWYGM DY	12

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SEQ ID NO: 240 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 240 QQSLWWPIT		9
SEQ ID NO: 241 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 241 SIYSYSGYTS YADSVKG		17
SEQ ID NO: 242 FEATURE source	moltype = AA length = 12 Location/Qualifiers 1..12 mol_type = protein organism = synthetic construct	
SEQUENCE: 242 HGSYGSWWAL DY		12
SEQ ID NO: 243 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 243 QQYFYFPIT		9
SEQ ID NO: 244 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 244 SISSYYGSTS YADSVKG		17
SEQ ID NO: 245 FEATURE source	moltype = AA length = 19 Location/Qualifiers 1..19 mol_type = protein organism = synthetic construct	
SEQUENCE: 245 ASYYSGYGSS YPYYMGLDY		19
SEQ ID NO: 246 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 246 QQGSYRNPLL T		11
SEQ ID NO: 247 FEATURE source	moltype = AA length = 10 Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 247 MNWSHYAMDY		10
SEQ ID NO: 248 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 248 YISSYSGYTY YADSVKG		17
SEQ ID NO: 249 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 249		

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YWYGHYHSYF GLDY	14
SEQ ID NO: 250 FEATURE source SEQUENCE: 250 SISSYSGSTY YADSVKG	17
SEQ ID NO: 251 FEATURE source SEQUENCE: 251 YPYGSHVYTG LDY	13
SEQ ID NO: 252 FEATURE source SEQUENCE: 252 QQQNWADYLV T	11
SEQ ID NO: 253 FEATURE source SEQUENCE: 253 SIYSSSGSTS YADSVKG	17
SEQ ID NO: 254 FEATURE source SEQUENCE: 254 VYSSRYWGWG VAFDY	15
SEQ ID NO: 255 FEATURE source SEQUENCE: 255 QQYYWYSLIT	10
SEQ ID NO: 256 FEATURE source SEQUENCE: 256 RSFPQWYNGS YTPWPAMDY	19
SEQ ID NO: 257 FEATURE source SEQUENCE: 257 QQYMWWPVT	9
SEQ ID NO: 258 FEATURE source SEQUENCE: 258 SIYSYSSYTS YADSVKG	17
SEQ ID NO: 259 FEATURE source SEQUENCE: 259 SIYSSYTS YADSVKG	12

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SEQUENCE: 259 PFYWGERYAL DY		12
SEQ ID NO: 260 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 260 QQSYSTLV		9
SEQ ID NO: 261 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 261 SIYSSYGSTS YADSVKG		17
SEQ ID NO: 262 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 262 IYGWSYQGWA GMDY		14
SEQ ID NO: 263 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 263 SIYPYYGSTS YADSVKG		17
SEQ ID NO: 264 FEATURE source	moltype = AA length = 15 Location/Qualifiers 1..15 mol_type = protein organism = synthetic construct	
SEQUENCE: 264 GGDYYWGWW VAMDY		15
SEQ ID NO: 265 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 265 QKSSSSLIT		9
SEQ ID NO: 266 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 266 SISSSYGSTS YADSVKG		17
SEQ ID NO: 267 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 267 MYYYTYPGMD Y		11
SEQ ID NO: 268 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 268 QKGSSYLLT		9
SEQ ID NO: 269 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein	

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SEQUENCE: 269 SISPSSGYTS YADSVKG	organism = synthetic construct	
SEQ ID NO: 270 FEATURE source	moltype = AA length = 15 Location/Qualifiers 1..15 mol_type = protein organism = synthetic construct	17
SEQUENCE: 270 YHYGGWSHYM SGMDY		15
SEQ ID NO: 271 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 271 QQNYYYYHKLI T		11
SEQ ID NO: 272 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 272 SISSSYGTY S YADSVKG		17
SEQ ID NO: 273 FEATURE source	moltype = AA length = 8 Location/Qualifiers 1..8 mol_type = protein organism = synthetic construct	
SEQUENCE: 273 GRYGGMDY		8
SEQ ID NO: 274 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 274 QQLSYVYKLI T		11
SEQ ID NO: 275 FEATURE source	moltype = AA length = 8 Location/Qualifiers 1..8 mol_type = protein organism = synthetic construct	
SEQUENCE: 275 GWYKAMDY		8
SEQ ID NO: 276 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 276 SISSSYGSTY YADSVKG		17
SEQ ID NO: 277 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 277 MYYYYYPGID Y		11
SEQ ID NO: 278 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 278 QQYYYFPIT		9
SEQ ID NO: 279 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17	

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SEQUENCE: 279 SISPYYGYTS YADSVKG	mol_type = protein organism = synthetic construct	
		17
SEQ ID NO: 280 FEATURE source	moltype = AA length = 16 Location/Qualifiers 1..16	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 280 SPYYWYQYFY GWGLDY		16
SEQ ID NO: 281 FEATURE source	moltype = AA length = 13 Location/Qualifiers 1..13	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 281 SPYWWNYMSA MDY		13
SEQ ID NO: 282 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 282 QQGWWWPPFT		9
SEQ ID NO: 283 FEATURE source	moltype = AA length = 16 Location/Qualifiers 1..16	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 283 WSWQYYSGHS SWGLDY		16
SEQ ID NO: 284 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 284 QQSWYFPLT		9
SEQ ID NO: 285 FEATURE source	moltype = AA length = 16 Location/Qualifiers 1..16	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 285 WYNÉYYHDYY WDAMDY		16
SEQ ID NO: 286 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 286 WMYWWSFALD Y		11
SEQ ID NO: 287 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 287 QQSYLWPIT		9
SEQ ID NO: 288 FEATURE source	moltype = AA length = 13 Location/Qualifiers 1..13	
	mol_type = protein organism = synthetic construct	
SEQUENCE: 288 WQYHYNWYG MDY		13
SEQ ID NO: 289 FEATURE	moltype = AA length = 9 Location/Qualifiers	

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source	1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 289 QQYPMSLIT		9
SEQ ID NO: 290 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 290 SISPYSGYTS YADSVKG		17
SEQ ID NO: 291 FEATURE source	moltype = AA length = 10 Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 291 GYDYYAGLDY		10
SEQ ID NO: 292 FEATURE source	moltype = AA length = 5 Location/Qualifiers 1..5 mol_type = protein organism = synthetic construct	
SEQUENCE: 292 YYSIH		5
SEQ ID NO: 293 FEATURE source	moltype = AA length = 12 Location/Qualifiers 1..12 mol_type = protein organism = synthetic construct	
SEQUENCE: 293 WESEYSGTYE DY		12
SEQ ID NO: 294 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 294 QQYMWWPIT		9
SEQ ID NO: 295 FEATURE source	moltype = AA length = 12 Location/Qualifiers 1..12 mol_type = protein organism = synthetic construct	
SEQUENCE: 295 TGYWQGYLAL DY		12
SEQ ID NO: 296 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 296 YYYYWNSTP AMDY		14
SEQ ID NO: 297 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 297 QQSYGYPVT		9
SEQ ID NO: 298 FEATURE source	moltype = AA length = 20 Location/Qualifiers 1..20 mol_type = protein organism = synthetic construct	
SEQUENCE: 298 WYNSSSWYYSN WWYKGFGMDY		20
SEQ ID NO: 299	moltype = AA length = 9	

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FEATURE source	Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 299 QQYYSSLFT		9
SEQ ID NO: 300 FEATURE source	moltype = AA length = 15 Location/Qualifiers 1..15 mol_type = protein organism = synthetic construct	
SEQUENCE: 300 TSYTYPVVITY YGFDY		15
SEQ ID NO: 301 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 301 QQSWYYPLT		9
SEQ ID NO: 302 FEATURE source	moltype = AA length = 13 Location/Qualifiers 1..13 mol_type = protein organism = synthetic construct	
SEQUENCE: 302 YRYSSWNRGA IDY		13
SEQ ID NO: 303 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 303 QQSYWWPLT		9
SEQ ID NO: 304 FEATURE source	moltype = AA length = 13 Location/Qualifiers 1..13 mol_type = protein organism = synthetic construct	
SEQUENCE: 304 WSKSPWYYQG IDY		13
SEQ ID NO: 305 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 305 QQHYHWASLI T		11
SEQ ID NO: 306 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 306 SIYSYSGSTY YADSVKG		17
SEQ ID NO: 307 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 307 QYSSYYVWP GMDY		14
SEQ ID NO: 308 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 308 QQSYWWKSLV T		11

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SEQ ID NO: 309	moltype = AA length = 17
FEATURE	Location/Qualifiers
source	1..17
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 309	
MHYSWQEYYS YDWGMDY	17
SEQ ID NO: 310	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 310	
QQPYYPLIT	9
SEQ ID NO: 311	moltype = AA length = 17
FEATURE	Location/Qualifiers
source	1..17
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 311	
SIYPSYGSTS YADSVKG	17
SEQ ID NO: 312	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 312	
WQGYYQPALD Y	11
SEQ ID NO: 313	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 313	
QQSSKKYYPI T	11
SEQ ID NO: 314	moltype = AA length = 12
FEATURE	Location/Qualifiers
source	1..12
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 314	
WGYGWYWGL DY	12
SEQ ID NO: 315	moltype = AA length = 10
FEATURE	Location/Qualifiers
source	1..10
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 315	
QQGHDMNPVT	10
SEQ ID NO: 316	moltype = AA length = 17
FEATURE	Location/Qualifiers
source	1..17
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 316	
SIYSSYGYTY YADSVKG	17
SEQ ID NO: 317	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 317	
YYYYWYGGMD Y	11
SEQ ID NO: 318	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 318	
QQSWMSDSL T	11

-continued

SEQ ID NO: 319 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct
SEQUENCE: 319 SIYPSSGYTS YADSVKG	17
SEQ ID NO: 320 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct
SEQUENCE: 320 GWWYWMAMDY AMDY	14
SEQ ID NO: 321 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 321 QQMQYSGWLI T	11
SEQ ID NO: 322 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct
SEQUENCE: 322 SISSYYGYTS YADSVKG	17
SEQ ID NO: 323 FEATURE source	moltype = AA length = 21 Location/Qualifiers 1..21 mol_type = protein organism = synthetic construct
SEQUENCE: 323 YYSYSSGYGY YDYFDWGAMD Y	21
SEQ ID NO: 324 FEATURE source	moltype = AA length = 20 Location/Qualifiers 1..20 mol_type = protein organism = synthetic construct
SEQUENCE: 324 YYGYVWGGYW GWWYSKALDY	20
SEQ ID NO: 325 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 325 QQYDWNYLYV T	11
SEQ ID NO: 326 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct
SEQUENCE: 326 YQYYGSLYY S QQWAMDY	17
SEQ ID NO: 327 FEATURE source	moltype = AA length = 20 Location/Qualifiers 1..20 mol_type = protein organism = synthetic construct
SEQUENCE: 327 SPSSPYFMSW GWWQYGIDY	20
SEQ ID NO: 328 FEATURE source	moltype = AA length = 10 Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct
SEQUENCE: 328	

-continued

QQSSWGLVLT	10
SEQ ID NO: 329 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct
SEQUENCE: 329 SISPSYGYTS YADSVKG	17
SEQ ID NO: 330 FEATURE source	moltype = AA length = 12 Location/Qualifiers 1..12 mol_type = protein organism = synthetic construct
SEQUENCE: 330 DMYEWWHWAI DY	12
SEQ ID NO: 331 FEATURE source	moltype = AA length = 20 Location/Qualifiers 1..20 mol_type = protein organism = synthetic construct
SEQUENCE: 331 YGHYLYYWG WGWWSAALDY	20
SEQ ID NO: 332 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 332 QQASYVKL I T	11
SEQ ID NO: 333 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 333 QQLSYVKL I T	11
SEQ ID NO: 334 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 334 QQPSYVKL I T	11
SEQ ID NO: 335 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 335 QQSSYVKL I T	11
SEQ ID NO: 336 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 336 QQTSYVKL I T	11
SEQ ID NO: 337 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct
SEQUENCE: 337 QQVSYVKL I T	11
SEQ ID NO: 338 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct

-continued

SEQUENCE: 338	
QQIKYVKKLI T	11
SEQ ID NO: 339	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 339	
QQIRYVKKLI T	11
SEQ ID NO: 340	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 340	
QQITYVKKLI T	11
SEQ ID NO: 341	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 341	
QQISFVKKLI T	11
SEQ ID NO: 342	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 342	
QQISYIKKLI T	11
SEQ ID NO: 343	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 343	
QQISYRKKLI T	11
SEQ ID NO: 344	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 344	
QQISYTKKLI T	11
SEQ ID NO: 345	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 345	
QQISYVAKLI T	11
SEQ ID NO: 346	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 346	
QQISYVEKLI T	11
SEQ ID NO: 347	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 347	
QQISYVHKLI T	11
SEQ ID NO: 348	moltype = AA length = 11
FEATURE	Location/Qualifiers
source	1..11
	mol_type = protein

-continued

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organism = synthetic construct
SEQUENCE: 348
QQISYVLKLI T                               11

SEQ ID NO: 349      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 349      organism = synthetic construct
QQISYVMKLI T                               11

SEQ ID NO: 350      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 350      organism = synthetic construct
QQISYVNKLI T                               11

SEQ ID NO: 351      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 351      organism = synthetic construct
QQISYVQKLI T                               11

SEQ ID NO: 352      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 352      organism = synthetic construct
QQISYVRKLI T                               11

SEQ ID NO: 353      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 353      organism = synthetic construct
QQISYVSKLI T                               11

SEQ ID NO: 354      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 354      organism = synthetic construct
QQISYVTKLI T                               11

SEQ ID NO: 355      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 355      organism = synthetic construct
QQISYVYKLI T                               11

SEQ ID NO: 356      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 356      organism = synthetic construct
QQISYVKRLI T                               11

SEQ ID NO: 357      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11
                  mol_type = protein
SEQUENCE: 357      organism = synthetic construct
QQISYVKKAI T                               11

SEQ ID NO: 358      moltype = AA length = 11
FEATURE          Location/Qualifiers
source           1..11

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-continued

SEQUENCE: 358	mol_type = protein organism = synthetic construct	
QQISYVKC1 T		11
SEQ ID NO: 359	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 359	organism = synthetic construct	
QQISYVKKDI T		11
SEQ ID NO: 360	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 360	organism = synthetic construct	
QQISYVKKEI T		11
SEQ ID NO: 361	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 361	organism = synthetic construct	
QQISYVKKGI T		11
SEQ ID NO: 362	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 362	organism = synthetic construct	
QQISYVKKHI T		11
SEQ ID NO: 363	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 363	organism = synthetic construct	
QQISYVKKKI T		11
SEQ ID NO: 364	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 364	organism = synthetic construct	
QQISYVKKMI T		11
SEQ ID NO: 365	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 365	organism = synthetic construct	
QQISYVKKNI T		11
SEQ ID NO: 366	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 366	organism = synthetic construct	
QQISYVKKPI T		11
SEQ ID NO: 367	moltype = AA length = 11	
FEATURE	Location/Qualifiers	
source	1..11	
	mol_type = protein	
SEQUENCE: 367	organism = synthetic construct	
QQISYVKKQI T		11
SEQ ID NO: 368	moltype = AA length = 11	
FEATURE	Location/Qualifiers	

-continued

source	1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 368 QQISYVKKRI T		11
SEQ ID NO: 369 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 369 QQISYVKKSI T		11
SEQ ID NO: 370 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 370 QQISYVKKTI T		11
SEQ ID NO: 371 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 371 QQISYVKKVI T		11
SEQ ID NO: 372 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 372 QQISYVKKWI T		11
SEQ ID NO: 373 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 373 QQISYVKLL T		11
SEQ ID NO: 374 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 374 QQISYVKKLV T		11
SEQ ID NO: 375 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 375 GAWYPAMDY		9
SEQ ID NO: 376 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 376 GCWYPAMDY		9
SEQ ID NO: 377 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 377 GEWYPAMDY		9
SEQ ID NO: 378	moltype = AA length = 9	

-continued

FEATURE source	Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 378 GHWYPAMDY		9
SEQ ID NO: 379 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 379 GWYYPAMDY		9
SEQ ID NO: 380 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 380 GLWYPAMDY		9
SEQ ID NO: 381 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 381 GMWYPAMDY		9
SEQ ID NO: 382 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 382 GNWYPAMDY		9
SEQ ID NO: 383 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 383 GPWYPAMDY		9
SEQ ID NO: 384 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 384 GQWYPAMDY		9
SEQ ID NO: 385 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 385 GRWYPAMDY		9
SEQ ID NO: 386 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 386 GSWYPAMDY		9
SEQ ID NO: 387 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 387 GTWYPAMDY		9

-continued

SEQ ID NO: 388	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 388	GWYWYPAMDY
9	
SEQ ID NO: 389	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 389	GYWYPAMDY
9	
SEQ ID NO: 390	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 390	GGGYPAMDY
9	
SEQ ID NO: 391	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 391	GGPYYPAMDY
9	
SEQ ID NO: 392	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 392	GGRYPAMDY
9	
SEQ ID NO: 393	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 393	GGTYYPAMDY
9	
SEQ ID NO: 394	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 394	GGWAPAMDY
9	
SEQ ID NO: 395	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 395	GGWPAMDY
9	
SEQ ID NO: 396	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 396	GGWEPAAMDY
9	
SEQ ID NO: 397	moltype = AA length = 9 Location/Qualifiers source 1..9 mol_type = protein organism = synthetic construct
SEQUENCE: 397	GGWFPAAMDY
9	

-continued

SEQ ID NO: 398	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 398	
GGWGPAMDY	9
SEQ ID NO: 399	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 399	
GGWHPAMDY	9
SEQ ID NO: 400	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 400	
GGWIPAMDY	9
SEQ ID NO: 401	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 401	
GGWKPAMDY	9
SEQ ID NO: 402	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 402	
GGWMPAMDY	9
SEQ ID NO: 403	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 403	
GGWNPAMDY	9
SEQ ID NO: 404	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 404	
GGWQPAMDY	9
SEQ ID NO: 405	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 405	
GGWRPAMDY	9
SEQ ID NO: 406	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 406	
GGWSPAMDY	9
SEQ ID NO: 407	moltype = AA length = 9
FEATURE	Location/Qualifiers
source	1..9
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 407	

-continued

GGWTPAMDY

9

SEQ ID NO: 408 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 408
 GGWPAMDY

9

SEQ ID NO: 409 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 409
 GGWYAAMDY

9

SEQ ID NO: 410 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 410
 GGWYVAMDY

9

SEQ ID NO: 411 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 411
 GGWYPCMDY

9

SEQ ID NO: 412 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 412
 GGWYPGMDY

9

SEQ ID NO: 413 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 413
 GGWYPKMDY

9

SEQ ID NO: 414 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 414
 GGWYPLMDY

9

SEQ ID NO: 415 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 415
 GGWYPMMDY

9

SEQ ID NO: 416 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 416
 GGWYPQMDY

9

SEQ ID NO: 417 moltype = AA length = 9
 FEATURE Location/Qualifiers
 source 1..9
 mol_type = protein
 organism = synthetic construct

SEQUENCE: 417
GGWYPRMDY

9

SEQ ID NO: 418
FEATURE
source
moltype = AA length = 9
Location/Qualifiers
1..9
mol_type = protein
organism = synthetic construct

SEQUENCE: 418
GGWYPSMDY

9

SEQ ID NO: 419
FEATURE
source
moltype = AA length = 9
Location/Qualifiers
1..9
mol_type = protein
organism = synthetic construct

SEQUENCE: 419
GGWYPTMDY

9

SEQ ID NO: 420
FEATURE
source
moltype = AA length = 9
Location/Qualifiers
1..9
mol_type = protein
organism = synthetic construct

SEQUENCE: 420
GGWYPYMDY

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What is claimed is:

1. A method comprising: administering to a human subject a pharmaceutical composition comprising an antibody or antigen binding fragment thereof, wherein the human subject has been pretreated with AMG510 and has a cancer that expresses a G12C mutant KRAS protein that comprises the amino acid sequence of SEQ ID NO: 1, wherein the antibody or antigen binding fragment thereof binds to a peptide conjugate in complex with a major histocompatibility complex (MHC) that is HLA-A*03:01, wherein the peptide conjugate comprises a peptide from the G12C mutant KRAS protein that comprises the amino acid sequence of SEQ ID NO: 1 covalently linked to the AMG510, and wherein the antibody or antigen binding fragment thereof comprises

- (i) a heavy chain variable region (VH) comprising
 - a heavy chain complementarity determining region 1 (HC CDR1) having the amino acid sequence set forth in SEQ ID NO: 169,
 - a HC CDR2 having the amino acid sequence set forth in SEQ ID NO: 170, and
 - a HC CDR3 having an amino acid sequence selected from the group consisting of the amino acid sequences set forth in SEQ ID NOS: 375-420; and
- (ii) a light chain variable region (VL) comprising
 - a light chain (LC) CDR1 having the amino acid sequence set forth in SEQ ID NO: 166,
 - a LC CDR2 having the amino acid sequence set forth in SEQ ID NO: 167, and
 - a LC CDR3 having an amino acid sequence selected from the group consisting of the amino acid sequences set forth in SEQ ID NOS: 332-374.

2. The method of claim 1, wherein the cancer is renal cell carcinoma, breast cancer, prostate cancer, pancreatic cancer, lung cancer, liver cancer, ovarian cancer, cervical cancer, colon cancer, esophageal cancer, glioma, glioblastoma or another brain cancer, stomach cancer, bladder cancer, testicular cancer, head and neck cancer, melanoma or another skin cancer, a sarcoma, or a blood cancer.

3. The method of claim 1, wherein the AMG510 is covalently linked to a Cys residue of SEQ ID NO: 1.

4. The method of claim 1, wherein a dissociation constant (Kd) of the antibody or antigen binding fragment thereof to the peptide conjugate in complex with the MHC is lower than a Kd of the antibody or antigen binding fragment thereof to the peptide conjugate not in complex with the MHC.

5. The method of claim 1, wherein the cancer is a lung cancer, a colon cancer, or a pancreatic cancer.

6. The method of claim 1, wherein the antibody or antigen binding fragment thereof is an intact antibody, an antigen-binding (Fab) fragment, an Fab' fragment, an (Fab')2 fragment, an Fd, an Fv, a dAb, a single-chain Diabody (scDb), a single-chain variable fragment (scFv), or a CrossMab.

7. A method for treating a cancer that expresses a G12C mutant KRAS protein that comprises the amino acid sequence of SEQ ID NO: 1 in a human subject in need thereof, the method comprising administering to the human subject a therapeutically effective amount of a pharmaceutical composition comprising an antibody or antigen binding fragment thereof, wherein the human subject has been pretreated with AMG510, wherein the antibody or antigen binding fragment thereof binds to a peptide conjugate in complex with a major histocompatibility complex (MHC) that is HLA-A*03:01, wherein the peptide conjugate comprises a peptide from the G12C mutant KRAS protein that comprises the amino acid sequence of SEQ ID NO: 1 covalently linked to the AMG510, and wherein the antibody or antigen binding fragment thereof comprises

- (i) a VH comprises
 - a HC CDR1 having the amino acid sequence set forth in SEQ ID NO: 169,
 - a HC CDR2 having the amino acid sequence set forth in SEQ ID NO: 170, and
 - a HC CDR3 having the amino acid sequence set forth in SEQ ID NO: 171; and
- (ii) a VL comprises
 - a LC CDR1 having the amino acid sequence set forth in SEQ ID NO: 166,
 - a LC CDR2 having the amino acid sequence set forth in SEQ ID NO: 167, and

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65

159

a LC CDR3 having the amino acid sequence set forth
in SEQ ID NO: 168.

8. The method of claim 7, wherein

(i) the VH comprises

a HC CDR1 having the amino acid sequence set forth 5
in SEQ ID NO: 169,

a HC CDR2 having the amino acid sequence set forth
in SEQ ID NO: 170, and

a HC CDR3 having the amino acid sequence set forth
in SEQ ID NO: 171; and 10

(ii) the VL comprises

a LC CDR1 having the amino acid sequence set forth
in SEQ ID NO: 166,

a LC CDR2 having the amino acid sequence set forth
in SEQ ID NO: 167, and 15

a LC CDR3 having the amino acid sequence set forth
in SEQ ID NO: 168; and

wherein the VH has the amino acid sequence as set forth
in SEQ ID NO: 4, and the VL has the amino acid
sequence as set forth in SEQ ID NO: 3. 20

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* * * *