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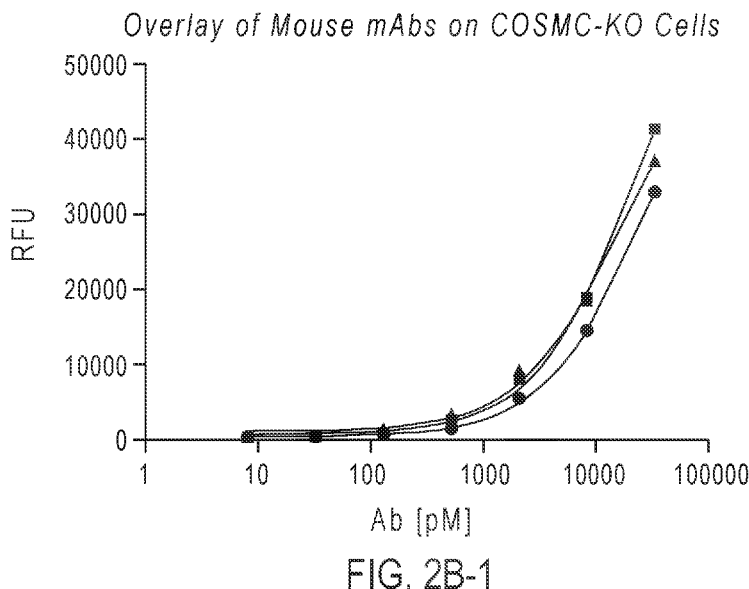
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(57) Abstract: The present disclosure relates to anti-glyco-cMET antibodies and antigen binding fragments thereof that specifically bind to a cancer-specific glycosylation variant of cMET and related fusion proteins and antibody-drug conjugates, as well as nucleic acids encoding such biomolecules. The present disclosure further relates to use of the antibodies, antigen-binding fragments, fusion proteins, antibody-drug conjugates and nucleic acids for cancer therapy.



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ANTI-GLYCO-CMET ANTIBODIES AND THEIR USES

1. CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims the priority benefit of U.S. provisional application no. 63/240,761, filed September 3, 2021, the contents of which are incorporated herein in their entireties by reference thereto.

2. BACKGROUND

[0002] Therapies redirecting T cell responses using chimeric antigen receptors (CARs) have emerged as a potent tool in cancer immunotherapies and have proved highly effective in haematological cancers, targeting antigens shared with nonessential tissues such as CD19 in B cell malignancies (Brentjens *et al.*, 2013, *Sci Transl Med.* 5(177):177ra38-177ra38; Grupp *et al.*, 2013, *N Engl J Med.* 368(16):1509–1518; Kalos *et al.*, 2011, *Sci Transl Med.* 3(95):95ra73-95ra73; Kochenderfer *et al.*, 2010, *Blood.* 116(20):4099–4102; Porter *et al.*, 2011, *N Engl J Med.* 365(8):725–733). However, adopting CAR therapies to solid tumours has been challenging because the majority of CAR targets are normal self-antigens overexpressed in solid cancers. As such, adverse effects due to cross-reactions with essential healthy tissues are often reported in studies targeting solid tumours with CAR T cells (Bin Hou *et al.*, 2019, *Dis Markers*, Article ID 3425291). To overcome the challenges of adopting CAR therapies to solid tumours, new cancer-specific antigens allowing selective targeting are required.

[0003] Many cancers express aberrantly glycosylated proteins that are distinct from healthy tissues. Such aberrantly glycosylated proteins contain glycopeptide epitopes that may be suitable for immunotherapy of solid tumors, but only few such glycopeptide epitopes have been identified.

[0004] The MET proto-oncogene encodes for the receptor tyrosine kinase that are widely expressed by cells of epithelial–endothelial origin (Brand-Saberi *et al.*, 1996, *Dev Biol.* 179(1):303-308; Heymann *et al.*, 1996, *Devel Biol.* 180(2):566-578; Bladt *et al.*, 1995, *Nature.* 376(6543):768771). Under normal conditions, c-MET signaling elicits a large variety of biological effects leading to increased cell growth, scattering and motility, invasion, protection from apoptosis, and angiogenesis (Sierra *et al.*, 2008, *J Exp Biol.* 205(7):1673-1655; Conrotto *et al.*, 2005, *Blood.* 105(11):4321-4329; Yi and Tsao, 2000, *Neoplasia.* 2(3):226-236; Silvagno *et al.*, 1995, *Arterioscler Thromb Vasc Biol.* 15(11):1857-1865), but in transformed epithelia improper activation of c-MET supports proliferation and invasive abilities of cancer cells (Benvenuti and Comoglio, 2007, *J Cell Physiol.* 213(2):316-325; Danilkovitch-Miagkova and Zbar, 2002, *J Clin Invest.* 109(7):863-867). Many studies have reported that c-MET is overexpressed in a variety of carcinomas. This includes lung, breast, ovary, kidney, colon, thyroid, liver, and gastric carcinomas (Knowles *et al.*, 2009, *Clin Cancer Res.* 15(11):3740-

3750; Lengyel *et al.*, 2005, *Int J Cancer*. 113(4):678-682; Tokunou *et al.*, 2001, *Am J Pathol*. 158(4):1451-1463; Ramirez *et al.*, 2000, *Clin Endocrinol (Oxf)*. 53(5):635-644; Tsao *et al.*, 1998, *Lung Cancer*. 20(1):1-16; Koochekpour *et al.*, 1997, *Cancer Res*. 57(23):5391-5398; Olivero *et al.*, 1996, *Br J Cancer*. 74(12):1862-1868; Tuck *et al.*, 1996, *Am J Pathol*. 148(1):225-232; Di Renzo *et al.*, 1995, *Cancer Res*. 55(5):1129-1138; Furukawa *et al.*, 1995, *Am J Pathol*. 147(4):889-895; Liu *et al.*, 1992, *Oncogene*. 7(1):181-185; Soman *et al.*, 1991, *Proc Natl Acad Sci USA*. 88(11):4892-4896; Houldsworth *et al.*, 1990, *Cancer Res*. 50(19):6417-6422).

Because of c-MET's importance in oncogenesis and cancer progression, c-MET is considered to be an important target in anticancer therapy (Trusolino *et al.*, 2010, *Nat Rev Mol Cell Bio*. 11(12):834-848; Migliore and Giordano, 2008, *Eur J Cancer*. 44(5):641-651; Peschard and Park, 2007, *Oncogene*. 26(9):1276-1285; Corso *et al.*, 2005, *Trends Mol Med*. 11(6):284-292). Several monoclonal antibodies that have displayed promising results in tumors with high HGF/c-MET levels, but most of these interfere primarily with c-MET's activation by HGF and are not suitably for immunotherapeutic targeting with cytotoxic strategies due to the prominent expression of c-MET in healthy tissue. Thus, there is a need for identification of glyco-cMET epitopes that are overexpressed in cancer cells and new therapeutic modalities, such as antibodies and CARs, which target such glyco-cMET epitopes.

3. SUMMARY

[0005] The disclosure captures the tumor specificity of glycopeptide variants by providing therapeutic and diagnostic agents based on antibodies and antigen binding fragments that are selective for cancer-specific epitopes of glyco-cMET. The antibodies and antigen-binding fragments advantageously bind to both the cMET backbone and its cancer specific O-linked glycans but not cMET on healthy tissues.

[0006] Accordingly, the present disclosure provides anti-glyco-cMET antibodies and antigen binding fragments thereof that bind to a cancer-specific glycosylation variant of cMET. The present disclosure further provides fusion proteins and antibody-drug conjugates comprising anti-glyco-cMET antibodies and antigen binding fragments, and nucleic acids encoding the anti-glyco-cMET antibodies, antigen binding fragments and fusion proteins.

[0007] The present disclosure further provides methods of using the anti-glyco-cMET antibodies, antigen-binding fragments, fusion proteins, antibody-drug conjugates and nucleic acids for cancer therapy.

[0008] In certain aspects, the disclosure provides bispecific and other multispecific anti-glyco-cMET antibodies and antigen binding fragments that bind to a cancer-specific glycosylation variant of cMET and to a second epitope. The second epitope can either be on cMET itself, on another protein co-expressed on cancer cells with cMET, or on another protein presented on a different cell, such as an activated T cell. Further, also disclosed are nucleic acids encoding such antibodies, including nucleic acids comprising codon-optimized coding regions and nucleic

acids comprising coding regions that are not codon-optimized for expression in a particular host cell.

[0009] The anti-glyco-cMET antibodies and binding fragments can be in the form of fusion proteins containing a fusion partner. The fusion partner can be useful to provide a second function, such as a signaling function of the signaling domain of a T cell signaling protein, a peptide modulator of T cell activation or an enzymatic component of a labeling system. Exemplary T cell signaling proteins include 4-1BB, CD28, CD2, and fusion peptides, e.g., CD28-CD3-zeta, 4-1BB-CD3-zeta, CD2-CD3-zeta, CD28-CD2-CD3-zeta, and 4-1BB-CD2-CD3-zeta. 4-1BB, also known as CD137, is a co-stimulatory receptor of T cells; CD2 is a co-stimulatory receptor of T and NK cells; CD3-zeta is a signal-transduction component of the T-cell antigen receptor. The moiety providing a second function can be a modulator of T cell activation, such as IL-15, IL-15R α , or an IL-15/IL-15R α fusion, can be an MHC-class I-chain-related (MIC) protein domain useful for making a MicAbody, or it can encode a label or an enzymatic component of a labeling system useful in monitoring the extent and/or location of binding *in vivo* or *in vitro*. Constructs encoding these prophylactically and therapeutically active biomolecules placed in the context of T cells, such as autologous T cells, provide a powerful platform for recruiting adoptively transferred T cells to prevent or treat a variety of cancers in some embodiments of the disclosure.

[0010] In certain aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain CDR sequences (as defined by Kabat, Chothia, IMGT or their combined region of overlap) of the anti-glyco-cMET antibodies 15C4.1D8.1G2 (sometimes referred to herein as "15C4"), 8H3.2B9.2C7 (sometimes referred to herein as "8H3"), 16E12.1D9.1B11 (sometimes referred to herein as "16E12"), 14E9, 19H2, or 39A3, or humanized counterparts of any one thereof. In some embodiments, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain variable sequences (or encoded by the nucleotide sequences) of the anti-glyco-cMET antibodies 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 of humanized counterparts thereof. The CDR and variable sequences (as well as their coding sequences) of the anti-glyco-cMET antibodies 15C4, 8H3, 16E12, 14E9, 19H2, AND 39A3 are set forth in Tables 1A through 1F, respectively. In certain aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain variable sequences (or encoded by the nucleotide sequences) set forth in Tables 1A through 1F. For clarity, when the term "anti-glyco-cMET antibody" is used in this document, it is intended to include monospecific and multi-specific (including bispecific) anti-glyco-cMET antibodies, antigen-binding fragments of the monospecific and multi-specific antibodies, and fusion proteins and conjugates containing the antibodies and their antigen-binding fragments, unless the context dictates otherwise. Likewise, when the term "anti-glyco-cMET antibody or antigen-binding fragment" is used, it is also

intended to include monospecific and multi-specific (including bispecific) anti-glyco-cMET antibodies and their antigen-binding fragments, together with fusion proteins and conjugates containing such antibodies and antigen-binding fragments, unless the context dictates otherwise.

[0011] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain CDR sequences (or encoded by the nucleotide sequences) set forth in Tables 1A-3H. The CDR sequences set forth in Tables 1A-1F include CDR sequences defined according to the IMGT (Lefranc *et al.*, 2003, *Dev Comparat Immunol* 27:55-77), Kabat (Kabat *et al.*, 1991, *Sequences of Proteins of Immunological Interest*, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, Md.), and Chothia (Al-Lazikani *et al.*, 1997, *J. Mol. Biol* 273:927-948) schemes for defining CDR boundaries. The CDR sequences set forth in Tables 1G-1I are consensus sequences derived from the CDR sequences set forth in Tables 1A through 1C ("Group 1" antibodies: 15C4, 8H3, 16E12) according to the IMGT, Kabat, and Chothia definitions, respectively. The CDR sequences set forth in Tables 1J-1L are consensus sequences derived from the CDR sequences set forth in Tables 1D through 1F ("Group 2" antibodies: 14E9, 19H2, and 39A3) according to the IMGT, Kabat, and Chothia definitions, respectively. The CDR sequences set forth in Tables 2A through 2F are the combined regions of overlap for the CDR sequences set forth in Tables 1A through 1F, respectively, with the IMGT, Kabat and Chothia sequences shown in underlined bold text. The CDR sequences set forth in Table 2G are the combined regions of overlap for the consensus CDR sequences set forth in Tables 2A-2C ("Group 1" antibodies: 15C4, 8H3, 16E12). The CDR sequences set forth in Table 2H are the combined regions of overlap for the consensus CDR sequences set forth in Tables 2D-2F ("Group 2" antibodies: 14E9, 19H2, and 39A3). The CDR sequences set forth in Tables 3A-3F are the common regions of overlap for the CDR sequences shown in Tables 1A-1F, respectively. The CDR sequences set forth in Table 3G are the common regions of overlap for the CDR sequences set forth in Tables 3A-3D ("Group 1" antibodies: 15C4, 8H3, 16E12). The CDR sequences set forth in Table 3H are the common regions of overlap for the CDR sequences set forth in Tables 3D-3F ("Group 2" antibodies: 14E9, 19H2, and 39A3). The framework sequences for such anti-glyco-cMET antibody and antigen-binding fragment can be the native murine framework sequences of the VH and VL sequences set forth in Tables 1A-1F or can be non-native (*e.g.*, humanized or human) framework sequences.

[0012] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain variable sequences of humanized anti-glyco-cMET antibody 8H3 set forth in Tables 4A through 4G.

Table 1A		
15C4.1D8.1G2 Sequences		
Description	Sequence	SEQ ID NO:
VH amino acid sequence (predicted mature)	QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWK QKPEQGLEWIGYFSPGNGDIKYNEKFKGKATLTADKSSS TAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGGQTSVT VSS	1
VL amino acid sequence (predicted mature)	NIVMTQSPKSMSMSVGERVTLSCKASENVGIYVSWYQQ KPEQSPKLLIYGPSNRYTGVPDRFTGSGSATDFTLTSSV QAEDLADYHCGQSYSYPFTFGSGTKLEIK	2
CDR-H1 amino acid sequence (IMGT definition)	GYTFTDHA	3
CDR-H2 amino acid sequence (IMGT definition)	FSPGNGDI	4
CDR-H3 amino acid sequence (IMGT definition)	KRSLPGPMDC	5
CDR-L1 amino acid sequence (IMGT definition)	ENVGIY	6
CDR-L2 amino acid sequence (IMGT definition)	GPS	7
CDR-L3 amino acid sequence (IMGT definition)	GQSYSYPFT	8
CDR-H1 amino acid sequence (Kabat definition)	DHAIH	9
CDR-H2 amino acid sequence (Kabat definition)	YFSPGNGDIKYNEKFKG	10
CDR-H3 amino acid sequence (Kabat definition)	SLPGPMDC	11
CDR-L1 amino acid sequence (Kabat definition)	KASENVGIYVS	12
CDR-L2 amino acid sequence (Kabat definition)	GPSNRYT	13

Table 1A 15C4.1D8.1G2 Sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Kabat definition)	GQSYSYPFT	14
CDR-H1 amino acid sequence (Chothia definition)	GYTFTDH	15
CDR-H2 amino acid sequence (Chothia definition)	SPGN GD	16
CDR-H3 amino acid sequence (Chothia definition)	SLPGPMDC	17
CDR-L1 amino acid sequence (Chothia definition)	KASENVGIYVS	18
CDR-L2 amino acid sequence (Chothia definition)	GPSNRYT	19
CDR-L3 amino acid sequence (Chothia definition)	GQSYSYPFT	20
VH nucleotide sequence (excl. signal sequence)	CAGGTT CAGCTGCAGCAGTCTGACGCTGAGTTGGTGA AACCTGGGGCTTCAGTGAAGATATCCTGCAAGGCTTC TGGCTACACCTTCACTGACCATGCTATTCCTGGGTG AAGCAGAAGCCTGAACAGGGCCTGGAATGGATTGGA TATTTTTCTCCCGGAAATGGTGATATTAAGTACAATGA GAAGTTCAAGGGCAAGGCCCACTGACTGCAGACAAA TCCTCCAGCACTGCCTACATGCAGCTCAACAGCCTGA CATCTGAGGATTCTGCAGTGTATTTCTGTAAAAGATCG CTACCGGGGCCTATGGACTGCTGGGGTCAAGGAACC TCAGTCACCGTCTCCTCA	21
VL nucleotide sequence (excl. signal sequence)	AACATTGTAATGACCCAATCTCCCAAATCCATGTCCAT GTCAGTGGGAGAGAGGGTCACCTTGAGCTGCAAGGC CAGTGAGAATGTGGGTATTTATGTATCCTGGTATCAAC AGAAACCAGAGCAGTCTCCTAAACTGCTGATATACGG GCCATCCAACCGGTACACTGGGGTCCCGATCGCTT CACAGGCAGTGGATCTGCAACAGATTTCACTCTGACC ATCAGCAGTGTGCAGGCTGAAGACCTTGACAGATTATC ACTGTGGACAGAGTTACAGCTATCCATTCACGTTCCGG CTCGGGGACAAAGTTGAAATAAAA	22

Table 1B		
8H3.2B9.2C7 Sequences		
Description	Sequence	SEQ ID NO:
VH amino acid sequence (predicted mature)	QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWK QRPEQGLEWIGYFSPGNGDIKYNEKFKDKATLTADKSSS TAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLV SS	23
VL amino acid sequence (predicted mature)	DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEK GKTNKLLIYSGSTLHSGIPSRFSGSGSGTDFTLTITSLAPE DFAMYFCQQHNEYPFTEGAGTKLELK	24
CDR-H1 amino acid sequence (IMGT definition)	GYTFTDHA	25
CDR-H2 amino acid sequence (IMGT definition)	FSPGNGDI	26
CDR-H3 amino acid sequence (IMGT definition)	KRSLPGDFDY	27
CDR-L1 amino acid sequence (IMGT definition)	KSVSEY	28
CDR-L2 amino acid sequence (IMGT definition)	SGS	29
CDR-L3 amino acid sequence (IMGT definition)	QQHNEYPFT	30
CDR-H1 amino acid sequence (Kabat definition)	DHAIH	31
CDR-H2 amino acid sequence (Kabat definition)	YFSPGNGDIKYNEKFKD	32
CDR-H3 amino acid sequence (Kabat definition)	SLPGDFDY	33
CDR-L1 amino acid sequence (Kabat definition)	RASKSVSEYLA	34
CDR-L2 amino acid sequence (Kabat definition)	SGSTLHS	35

Table 1B 8H3.2B9.2C7 Sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Kabat definition)	QQHNEYPFPT	36
CDR-H1 amino acid sequence (Chothia definition)	GYTFTDH	37
CDR-H2 amino acid sequence (Chothia definition)	SPGN GD	38
CDR-H3 amino acid sequence (Chothia definition)	SLPGDFDY	39
CDR-L1 amino acid sequence (Chothia definition)	RASKSVSEYLA	40
CDR-L2 amino acid sequence (Chothia definition)	SGSTLHS	41
CDR-L3 amino acid sequence (Chothia definition)	QQHNEYPFPT	42
VH nucleotide sequence (excl. signal sequence)	CAGGTT CAGCTGCAGCAGTCTGACGCTGAGTTGGTGA AACCTGGGGCTTCAGTGAAGATATCCTGCAAGGCTTC TGGCTACACCTTCACTGACCATGCTATTCCTGGGTG AAGCAGAGGCCTGAACAGGGCCTGGAATGGATTGGA TATTTTTCTCCCGGAAATGGTGATATTAAGTACAATGA GAAGTTCAAGGACAAGGCCCACTGACTGCAGACAAG TCCTCCAGCACTGCCTACATGCAGCTCAACAGCCTGA CATCTGAGGATTCTGCAGTGTATTTCTGTAAACGTTCC CTACCGGGGGACTTTGACTACTGGGGCCAAGGCACC ACTCTCACAGTCTCCTCA	43
VL nucleotide sequence (excl. signal sequence)	GATGTCCAGATAACCCAGTCTCCATCTTATCTTGCTGC ATCTCCTGGAGAAACCATTACTATTAATTGCCGGGCAA GTAAGAGCGTTAGCGAATATTTAGCCTGGTATCAAGA GAAACCTGGGAAAATAATAAGCTTCTTATCTACTCTG GATCCACTTTGCACTCTGGAATTCCATCAAGTTTCAGT GGCAGTGGATCTGGTACAGATTTCACTCTCACCATCA CTAGCCTGGCGCCTGAAGATTTTGCAATGTATTTCTGT CAACAGCATAATGAATACCCGTTACGTTTCGGTGCTG GGACCAAGCTGGAGCTGAAA	44

Table 1C		
16E12.1D9.1B11 Sequences		
Description	Sequence	SEQ ID NO:
VH amino acid sequence (predicted mature)	QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNDDVRYSEKFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLT VSS	45
VL amino acid sequence (predicted mature)	DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKPGKTIKPLIYSGSTLQTGTPSRFSGSGSGTDFSLTISLEPEDFAMYYCQQHNEYPPFTFGAGTKLELK	46
CDR-H1 amino acid sequence (IMGT definition)	GYTFTDHA	47
CDR-H2 amino acid sequence (IMGT definition)	FSPGNDDV	48
CDR-H3 amino acid sequence (IMGT definition)	KRSLPGDFDY	49
CDR-L1 amino acid sequence (IMGT definition)	KSINNY	50
CDR-L2 amino acid sequence (IMGT definition)	SGS	51
CDR-L3 amino acid sequence (IMGT definition)	QQHNEYPPFT	52
CDR-H1 amino acid sequence (Kabat definition)	DHAIH	53
CDR-H2 amino acid sequence (Kabat definition)	YFSPGNDDVRYSEKFKG	54
CDR-H3 amino acid sequence (Kabat definition)	SLPGDFDY	55
CDR-L1 amino acid sequence (Kabat definition)	RASKSINNYLV	56
CDR-L2 amino acid sequence (Kabat definition)	SGSTLQT	57

Table 1C 16E12.1D9.1B11 Sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Kabat definition)	QQHNEYPPFT	58
CDR-H1 amino acid sequence (Chothia definition)	GYTFTDH	59
CDR-H2 amino acid sequence (Chothia definition)	SPGNDD	60
CDR-H3 amino acid sequence (Chothia definition)	SLPGDFDY	61
CDR-L1 amino acid sequence (Chothia definition)	RASKSINNYLV	62
CDR-L2 amino acid sequence (Chothia definition)	SGSTLQT	63
CDR-L3 amino acid sequence (Chothia definition)	QQHNEYPPFT	64
VH nucleotide sequence (excl. signal sequence)	CAGGTT CAGCTGCAGCAGTCTGACGCTGAATTGGTGA AACCTGGGGCTTCAGTGAAGATATCCTGCAAGGCTTC TGGCTACACCTTCACTGACCATGCTATTCCTGGGTG AAGCAGAAGCCTGAACAGGGCCTGGAATGGATTGGA TATTTTTCTCCCGGAAATGATGATGTTAGGTACAGTGA GAAGTTCAAGGGCAAGGCCCACTGACTGCAGACAAA TCCTCCAGCACTGCCTACATGCAGCTCAACAGCCTGA CATCTGAGGATTCTGCAGTGTATTTCTGTAAACGTTCC CTACCGGGGGACTTTGACTACTGGGGCCAAGGCACC ACCCTCACAGTCTCCTCA	65
VL nucleotide sequence (excl. signal sequence)	GATGTCCAGATATCCCAGTCTCCATCTTATCTTGCTGC ATCTCCTGGAGAAACCATTACAATTAATTGCAGGGCAA GTAAGAGCATTAACTATTTAGTCTGGTATCAAGAG AAACCTGGGAAAATTAAGCCTCTTATCTACTCTGG ATCCACTTTGCAAACCTGGAACCTCCATCAAGGTTGAGT GGCAGTGGATCTGGTACAGATTTGAGTCTCACCATCA GTAGCCTGGAGCCTGAAGATTTTGCAATGTATTACTGT CAACAGCATAATGAATATCCGTTACGTTCCGGTGCTG GGACCAAGTTGGAGCTGAAA	66

Table 1D 14E9 Sequences		
Description	Sequence	SEQ ID NO:
VH amino acid sequence (predicted mature)	QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVR QAPGKGLEWVGCITYTGSGGNTYYATWAKGRFTVSETS STTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVG AFNLWGQGTLVTVSSGQPK	67
VL amino acid sequence (predicted mature)	DVVMTPASVGAAVGGTVTIKCQASQSSISNWLAWYQQ KPGQPPKLLIYSASYLESVPSRFSGSGSGTEFTLTISDL ECADAATYYCQCTYGSSGDSGSWDFGGGTEVVVKGDP V	68
CDR-H1 amino acid sequence (IMGT definition)	GIDFSSYW	69
CDR-H2 amino acid sequence (IMGT definition)	IYTGSGGNT	70
CDR-H3 amino acid sequence (IMGT definition)	ARMGYSAGYIGATYITVGAFNL	71
CDR-L1 amino acid sequence (IMGT definition)	QSSISNW	72
CDR-L2 amino acid sequence (IMGT definition)	SAS	73
CDR-L3 amino acid sequence (IMGT definition)	QCTYGSSGDSGSWD	74
CDR-H1 amino acid sequence (Kabat definition)	SYWIC	75
CDR-H2 amino acid sequence (Kabat definition)	CIYTGSGGNTYYATWAKG	76
CDR-H3 amino acid sequence (Kabat definition)	MGYSAGYIGATYITVGAFNL	77
CDR-L1 amino acid sequence (Kabat definition)	QASQSSISNWLA	78
CDR-L2 amino acid sequence (Kabat definition)	SASYLES	79

Table 1D 14E9 Sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Kabat definition)	QCTYGSSGDSGSWD	80
CDR-H1 amino acid sequence (Chothia definition)	GIDFSSY	81
CDR-H2 amino acid sequence (Chothia definition)	YTGSGGN	82
CDR-H3 amino acid sequence (Chothia definition)	MGYSAGYIGATYITVGAFNL	83
CDR-L1 amino acid sequence (Chothia definition)	QASQISISNWLA	84
CDR-L2 amino acid sequence (Chothia definition)	SASYLES	85
CDR-L3 amino acid sequence (Chothia definition)	QCTYGSSGDSGSWD	86
VH nucleotide sequence (excl. signal sequence)	CAGGAGCAGCTGGTGGAGTCCGGGGGAGGCCTGGT CGAGCCTGGGGCATCCCTGACACTCACCTGCAAAGC CTCTGGAATCGACTTCAGTAGCTACTGGATATGCTGG GTCCGCCAGGCTCCAGGGAAGGGGCTGGAGTGGGT CGGATGCATTTATACTGGTAGTGGTGGTAACACTTACT ACGCGACCTGGGCGAAGGGCCGATTCACCGTCTCCG AAACCTCGTCGACCACGGTGACTCTGCGAATGACCAG TCTGACGGCCGCGGACACGGCCACCTATTTCTGTGCA AGAATGGGGTATAGTGCTGGTTATATTGGTGCTACTTA TATTACCGTGGGTGCCTTTAATTTGTGGGGCCAGGGC ACCCTGGTCACCGTCTCGAGCGGACAGCCGAAA	87
VL nucleotide sequence (excl. signal sequence)	GATGTTGTGATGACCCAGACTCCAGCCTCCGTGGGG GCTGCTGTGGGAGGCACAGTCACCATCAAGTGCCAG GCCAGTCAGAGCATTAGCAACTGGTTAGCCTGGTATC AGCAGAAACCAGGGCAGCCTCCAAGCTCCTGATCTA TTCTGCATCCTATCTGGAATCTGGGGTCCCATCGCGG TTCAGCGGCAGTGGATCTGGGACAGAGTTCACTCTCA CCATCAGCGACCTGGAGTGTGCCGATGCTGCCACTTA CTACTGTCAATGTACTTATGGTAGTAGTGGTGATAGTG GTAGTTGGGATTTCCGGCGGAGGGACCGAGGTGGTGG TCAAAGGTGATCCCGTG	88

Table 1E		
19H2 Sequences		
Description	Sequence	SEQ ID NO:
VH amino acid sequence (predicted mature)	QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICW VRQAPGKGLEWIGCIYTGSSATDYANWARGRFTISKG SSPTVDLKMTSLTGADTGTYFCARMGYEDGYVGGVYTI VGAFNLWGQGLTLTVSSGQPK	89
VL amino acid sequence (predicted mature)	DVVMQTASPVSAAVGGTVTIKQASQTISSYLAWYQQ KPGQPPKLLIYATSYLESQVPSRFKGGSGGTQFTLTISGV QCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV	90
CDR-H1 amino acid sequence (IMGT definition)	GVAFSGSQW	91
CDR-H2 amino acid sequence (IMGT definition)	IYTGSSATD	92
CDR-H3 amino acid sequence (IMGT definition)	ARMGYEDGYVGGVYTIIVGAFNL	93
CDR-L1 amino acid sequence (IMGT definition)	QTISSY	94
CDR-L2 amino acid sequence (IMGT definition)	ATS	95
CDR-L3 amino acid sequence (IMGT definition)	QCSYSGSYSGSWT	96
CDR-H1 amino acid sequence (Kabat definition)	GSQWIC	97
CDR-H2 amino acid sequence (Kabat definition)	CIYTGSSATDYANWARG	98
CDR-H3 amino acid sequence (Kabat definition)	MGYEDGYVGGVYTIIVGAFNL	99
CDR-L1 amino acid sequence (Kabat definition)	QASQTISSYLA	100
CDR-L2 amino acid sequence (Kabat definition)	ATSYLES	101

Table 1E 19H2 Sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Kabat definition)	QCSYGSGYSGSWT	102
CDR-H1 amino acid sequence (Chothia definition)	GVAFSGSQ	103
CDR-H2 amino acid sequence (Chothia definition)	YTGSSAT	104
CDR-H3 amino acid sequence (Chothia definition)	MGYEDGYVGGVYTIVGAFNL	105
CDR-L1 amino acid sequence (Chothia definition)	QASQTISSYLA	106
CDR-L2 amino acid sequence (Chothia definition)	ATSYLES	107
CDR-L3 amino acid sequence (Chothia definition)	QCSYGSGYSGSWT	108
VH nucleotide sequence (excl. signal sequence)	CAGCAGCAGCTGGAGGAGTCCGGGGGAGGCCTGGT CAAGCCTGGGGCATCCCTGACACTCACCTGCCGAGC CTCTGGGGTTCGCCTTCAGTGGGAGCCAGTGGATATG TTGGGTCCGTCAGGCTCCAGGGAAGGGGCTGGAGTG GATCGGTTGCATTTATACTGGCAGTAGTGCTACTGATT ATTACGCGAACTGGGCGAGAGGCCGATTCACCATCTC CAAAGGCTCGTCGCCACGGTGGATCTGAAAATGACC AGTCTGACAGGCGCGGACTCGGGCACCTATTTCTGTG CGAGAATGGGGTATGAAGATGGTTATGTTGGTGGAGT TTATACTATCGTGGGTGCCTTTAACTTGTGGGGCCAG GGCACCTGGTCACCGTCTCGAGCGGACAGCCGAAA	109
VL nucleotide sequence (excl. signal sequence)	GATGTTGTGATGACCCAGACTGCATCCCCCGTGTCTG CAGCTGTGGGAGGCACAGTCACCATCAAGTGCCAGG CCAGTCAGACCATTAGTAGCTACTTAGCCTGGTATCA GCAGAAACCAGGGCAGCCTCCAAGCTCCTGATCTAT GCTACATCCTATCTGGAATCTGGGGTCCCGTCGCGAT TCAAAGGCAGTGGATCTGGGACACAGTTCACTCTCAC CATCAGCGGCGTGCAGTGTGACGATGCTGCCACTTAT TACTGTCAATGCAGTTATGGTAGTGGTTACAGTGGTA GTTGGACTTTCGGCGGAGGGACCGAGGTGGTGGTCA AAGGTGATCCCGTG	110

Table 1F		
39A3 Sequences		
Description	Sequence	SEQ ID NO:
VH amino acid sequence (predicted mature)	QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVR QAPGKGLEWIACMDNRVTYATWAKGRFTSSKTSSTTVT LQMTSLTAADTATYFCARGGYGGRGLVFNLWGQGTLVT VSSGQPK	111
VL amino acid sequence (predicted mature)	DPVLTQTTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWY QQKPGQPPKLLIYDASTLASGVPSRFKGSQSGTQFTLTI SGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVVKGDP V	112
CDR-H1 amino acid sequence (IMGT definition)	GLDFSGIYW	113
CDR-H2 amino acid sequence (IMGT definition)	MDNRV	114
CDR-H3 amino acid sequence (IMGT definition)	ARGGYGGRGLVFNL	115
CDR-L1 amino acid sequence (IMGT definition)	QSVYNNNE	116
CDR-L2 amino acid sequence (IMGT definition)	DAS	117
CDR-L3 amino acid sequence (IMGT definition)	QGIYYIGDWYSA	118
CDR-H1 amino acid sequence (Kabat definition)	GIYWAC	119
CDR-H2 amino acid sequence (Kabat definition)	CMDNRVTYATWAKG	120
CDR-H3 amino acid sequence (Kabat definition)	GGYGGRGLVFNL	121
CDR-L1 amino acid sequence (Kabat definition)	QSSQSVYNNNELS	122
CDR-L2 amino acid sequence (Kabat definition)	DASTLAS	123

Table 1F 39A3 Sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Kabat definition)	QGIYYIGDWYSA	124
CDR-H1 amino acid sequence (Chothia definition)	GLDFSGIY	125
CDR-H2 amino acid sequence (Chothia definition)	DNR	126
CDR-H3 amino acid sequence (Chothia definition)	GGYGGRGLVFNL	127
CDR-L1 amino acid sequence (Chothia definition)	QSSQSVYNNNELS	128
CDR-L2 amino acid sequence (Chothia definition)	DASTLAS	129
CDR-L3 amino acid sequence (Chothia definition)	QGIYYIGDWYSA	130
VH nucleotide sequence (excl. signal sequence)	CAGTCATTGGAGGAGTACGGGGGAGACCTGGTCAAG CCTGGGGCATCCCTGACACTCACCTGCACAGCCTCTG GGTTAGACTTCAGTGGCATCTACTGGGCATGCTGGGT CCGCCAGGCTCCAGGGAAGGGGCTGGAGTGGATCG CTTGCATGGATAATCGTGTTACATACGCGACCTGGGC GAAAGGCCGATTCACCAGCTCCAAAACCTCGTCGACC ACGGTGACTCTTCAAATGACCAGTCTGACAGCCGCGG ACACGGCCACATATTTCTGTGCGAGAGGGGTTATGG TGGTCGTGGTTTGGTTTTAATTTGTGGGGCCAGGGC ACCCTGGTCACCGTCTCGAGCGGACAGCCGAAA	131
VL nucleotide sequence (excl. signal sequence)	GACCCTGTGTTGACCCAGACTCCACCCTCGGTGTCTG CAGCTGTGGGAGGCACAGTCACCATCAAGTGCCAGT CCAGTCAGAGTGTATAATAACAACGAATTATCCTGG TATCAGCAGAAACCAGGGCAGCCTCCCAAACCTTCTGA TCTATGATGCATCCACTCTGGCATCTGGGGTCCATC GCGGTTCAAAGGCAGTGGATCTGGGACACAGTTCACT CTCACCATCAGCGCGTGCAGTGTGACGATGCTGCC ACTTATTACTGTCAAGGCATTTATTATATTGGTGATTG GTATAGTGCTTTCGGCGGAGGGACCGAGGTGGTGGT CAAAGGTGATCCCGTG	132

Table 1G		
CDR Consensus sequences Group 1 – IMGT definition		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (IMGT definition)	GYTFTDHA	133
CDR-H2 amino acid sequence (IMGT definition)	FSPGNX ₁ DX ₂	134
CDR-H3 amino acid sequence (IMGT definition)	KRSLPGX ₆ X ₇ DX ₈	135
CDR-L1 amino acid sequence (IMGT definition)	X ₁₀ X ₁₁ X ₁₂ X ₁₃ X ₁₄ Y	136
CDR-L2 amino acid sequence (IMGT definition)	X ₁₇ X ₁₈ S	137
CDR-L3 amino acid sequence (IMGT definition)	X ₂₃ QX ₂₄ X ₂₅ X ₂₆ YPFT	138
X ₁ = G or D; X ₂ = I or V; X ₆ = P or D; X ₇ = M or F; X ₈ = C or Y; X ₁₀ = E or K; X ₁₁ = N or S; X ₁₂ = V or I; X ₁₃ = G, S, or N; X ₁₄ = I, E, or N; X ₁₇ = G or S; X ₁₈ = P or G; X ₂₃ = G or Q; X ₂₄ = S or H; X ₂₅ = Y or N; X ₂₆ = S or E		

Table 1H		
CDR Consensus sequences Group 1 – Kabat definition		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (Kabat definition)	DHAIH	139
CDR-H2 amino acid sequence (Kabat definition)	YFSPGNX ₁ DX ₂ X ₃ YX ₄ EKF ₅	140
CDR-H3 amino acid sequence (Kabat definition)	SLPGX ₆ X ₇ DX ₈	141
CDR-L1 amino acid sequence (Kabat definition)	X ₉ ASX ₁₀ X ₁₁ X ₁₂ X ₁₃ X ₁₄ YX ₁₅ X ₁₆	142
CDR-L2 amino acid sequence (Kabat definition)	X ₁₇ X ₁₈ SX ₁₉ X ₂₀ X ₂₁ X ₂₂	143
CDR-L3 amino acid sequence (Kabat definition)	X ₂₃ QX ₂₄ X ₂₅ X ₂₆ YPFT	144
X ₁ = G or D; X ₂ = I or V; X ₃ = K or R; X ₄ = N or S; X ₅ = G or D; X ₆ = P or D; X ₇ = M or F; X ₈ = C or Y; X ₉ = K or R; X ₁₀ = E or K; X ₁₁ = N or S; X ₁₂ = V or I; X ₁₃ = G, S, or N; X ₁₄ = I, E, or N; X ₁₅ = V or L; X ₁₆ = S, A, or V; X ₁₇ = G or S; X ₁₈ = P or G; X ₁₉ = N or T; X ₂₀ = R or L; X ₂₁ = Y, H, or Q; X ₂₂ = T or S; X ₂₃ = G or Q; X ₂₄ = S or H; X ₂₅ = Y or N; X ₂₆ = S or E		

Table 1I		
CDR Consensus sequences Group 1 – Chothia definition		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (Chothia definition)	GYTFTDH	145
CDR-H2 amino acid sequence (Chothia definition)	SPGNX ₁ D	146
CDR-H3 amino acid sequence (Chothia definition)	SLPGX ₆ X ₇ DX ₈	147
CDR-L1 amino acid sequence (Chothia definition)	X ₉ ASX ₁₀ X ₁₁ X ₁₂ X ₁₃ X ₁₄ YX ₁₅ X ₁₆	148
CDR-L2 amino acid sequence (Chothia definition)	X ₁₇ X ₁₈ SX ₁₉ X ₂₀ X ₂₁ X ₂₂	149

Table 1I		
CDR Consensus sequences Group 1 – Chothia definition		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (Chothia definition)	X ₂₃ QX ₂₄ X ₂₅ X ₂₆ YPFT	150
X ₁ = G or D; X ₆ = P or D; X ₇ = M or F; X ₈ = C or Y; X ₉ = K or R; X ₁₀ = E or K; X ₁₁ = N or S; X ₁₂ = V or I; X ₁₃ = G, S, or N; X ₁₄ = I, E, or N; X ₁₅ = V or L; X ₁₆ = S, A, or V; X ₁₇ = G or S; X ₁₈ = P or G; X ₁₉ = N or T; X ₂₀ = R or L; X ₂₁ = Y, H, or Q; X ₂₂ = T or S; X ₂₃ = G or Q; X ₂₄ = S or H; X ₂₅ = Y or N; X ₂₆ = S or E		

Table 1J		
CDR Consensus sequences Group 2 – IMGT definition		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (IMGT definition)	GX ₂₇ X ₂₈ FSX ₂₉ X ₃₀ X ₃₁ W	151
CDR-H2 amino acid sequence (IMGT definition)	X ₃₃ X ₃₄ X ₃₅ X ₃₆ X ₃₇ X ₃₈ X ₃₉ X ₄₀ X ₄₁	152
CDR-H3 amino acid sequence (IMGT definition)	ARX ₄₅ GYX ₄₆ X ₄₇ GX ₄₈ X ₄₉ GX ₅₀ X ₅₁ X ₅₂ X ₅₃ X ₅₄ VX ₅₅ X ₅₆ FNL	153
CDR-L1 amino acid sequence (IMGT definition)	QX ₅₈ X ₅₉ X ₆₀ X ₆₁ X ₆₂ X ₆₃ X ₆₄	154
CDR-L2 amino acid sequence (IMGT definition)	X ₆₆ X ₆₇ S	155
CDR-L3 amino acid sequence (IMGT definition)	QX ₇₀ X ₇₁ YX ₇₂ X ₇₃ X ₇₄ GX ₇₅ X ₇₆ X ₇₇ SX ₇₈ X ₇₉	156
X ₂₇ = I, V, or L; X ₂₈ = D or A; X ₂₉ = absent or G; X ₃₀ = S or I; X ₃₁ = Y or Q; X ₃₃ = I or M; X ₃₄ = Y or D; X ₃₅ = T or N; X ₃₆ = G or R; X ₃₇ = S, V, or absent; X ₃₈ = G, S, or absent; X ₃₉ = G, A, or absent; X ₄₀ = N, T, or absent; X ₄₁ = T, D, or absent; X ₄₅ = M or G; X ₄₆ = S, E, or G; X ₄₇ = A, D, or absent; X ₄₈ = Y or R; X ₄₉ = I, V, or absent; X ₅₀ = A, G, or absent; X ₅₁ = T, V, or absent; X ₅₂ = Y or absent; X ₅₃ = I, T, or absent; X ₅₄ = T, I, or L; X ₅₅ = G or absent; X ₅₆ = A or absent; X ₅₈ = S or T; X ₅₉ = I or V; X ₆₀ = S or Y; X ₆₁ = N or S; X ₆₂ = W, Y, or N; X ₆₃ = absent or N; X ₆₄ = absent or E; X ₆₆ = S, A, or D; X ₆₇ = A or T; X ₇₀ = C or G; X ₇₁ = T, S, or I; X ₇₂ = G or Y; X ₇₃ = S or I; X ₇₄ = S or absent; X ₇₅ = D or Y; X ₇₆ = S or W; X ₇₇ = G or Y; X ₇₈ = W or A; X ₇₉ = D, T, or absent		

Table 1K		
CDR Consensus sequences Group 2 – Kabat definition		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (Kabat definition)	X ₂₉ X ₃₀ X ₃₁ WX ₃₂ C	157
CDR-H2 amino acid sequence (Kabat definition)	CX ₃₃ X ₃₄ X ₃₅ X ₃₆ X ₃₇ X ₃₈ X ₃₉ X ₄₀ X ₄₁ X ₄₂ YAX ₄₃ WAX ₄₄ G	158
CDR-H3 amino acid sequence (Kabat definition)	X ₄₅ GYX ₄₆ X ₄₇ GX ₄₈ X ₄₉ GX ₅₀ X ₅₁ X ₅₂ X ₅₃ X ₅₄ VX ₅₅ X ₅₆ FNL	159
CDR-L1 amino acid sequence (Kabat definition)	QX ₅₇ SQX ₅₈ X ₅₉ X ₆₀ X ₆₁ X ₆₂ X ₆₃ X ₆₄ LX ₆₅	160
CDR-L2 amino acid sequence (Kabat definition)	X ₆₆ X ₆₇ SX ₆₈ LX ₆₉ S	161
CDR-L3 amino acid sequence (Kabat definition)	QX ₇₀ X ₇₁ YX ₇₂ X ₇₃ X ₇₄ GX ₇₅ X ₇₆ X ₇₇ SX ₇₈ X ₇₉	162
<p>X₂₉ = absent or G; X₃₀ = S or I; X₃₁ = Y or Q; X₃₂ = I or A; X₃₃ = I or M; X₃₄ = Y or D; X₃₅ = T or N; X₃₆ = G or R; X₃₇ = S, V, or absent; X₃₈ = G, S, or absent; X₃₉ = G, A, or absent; X₄₀ = N, T, or absent; X₄₁ = T, D, or absent; X₄₂ = Y or absent; X₄₃ = T or N; X₄₄ = K or R; X₄₅ = M or G; X₄₆ = S, E, or G; X₄₇ = A, D, or absent; X₄₈ = Y or R; X₄₉ = I, V, or absent; X₅₀ = A, G, or absent; X₅₁ = T, V, or absent; X₅₂ = Y or absent; X₅₃ = I, T, or absent; X₅₄ = T, I, or L; X₅₅ = G or absent; X₅₆ = A or absent; X₅₇ = A or S; X₅₈ = S or T; X₅₉ = I or V; X₆₀ = S or Y; X₆₁ = N or S; X₆₂ = W, Y, or N; X₆₃ = absent or N; X₆₄ = absent or E; X₆₅ = A or S; X₆₆ = S, A, or D; X₆₇ = A or T; X₆₈ = Y or T; X₆₉ = E or A; X₇₀ = C or G; X₇₁ = T, S, or I; X₇₂ = G or Y; X₇₃ = S or I; X₇₄ = S or absent; X₇₅ = D or Y; X₇₆ = S or W; X₇₇ = G or Y; X₇₈ = W or A; X₇₉ = D, T, or absent</p>		

Table 1L		
CDR Consensus sequences Group 2 – Chothia definition		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (Chothia definition)	GX ₂₇ X ₂₈ FSX ₂₉ X ₃₀ X ₃₁	163
CDR-H2 amino acid sequence (Chothia definition)	X ₃₄ X ₃₅ X ₃₆ X ₃₇ X ₃₈ X ₃₉ X ₄₀	164
CDR-H3 amino acid sequence (Chothia definition)	X ₄₅ GYX ₄₆ X ₄₇ GX ₄₈ X ₄₉ GX ₅₀ X ₅₁ X ₅₂ X ₅₃ X ₅₄ VX ₅₅ X ₅₆ FNL	165
CDR-L1 amino acid sequence (Chothia definition)	QX ₅₇ SQX ₅₈ X ₅₉ X ₆₀ X ₆₁ X ₆₂ X ₆₃ X ₆₄ LX ₆₅	166
CDR-L2 amino acid sequence (Chothia definition)	X ₆₆ X ₆₇ SX ₆₈ LX ₆₉ S	167
CDR-L3 amino acid sequence (Chothia definition)	QX ₇₀ X ₇₁ YX ₇₂ X ₇₃ X ₇₄ GX ₇₅ X ₇₆ X ₇₇ SX ₇₈ X ₇₉	168
<p>X₂₇ = I, V, or L; X₂₈ = D or A; X₂₉ = absent or G; X₃₀ = S or I; X₃₁ = Y or Q; X₃₄ = Y or D; X₃₅ = T or N; X₃₆ = G or R; X₃₇ = S, V, or absent; X₃₈ = G, S, or absent; X₃₉ = G, A, or absent; X₄₀ = N, T, or absent; X₄₅ = M or G; X₄₆ = S, E, or G; X₄₇ = A, D, or absent; X₄₈ = Y or R; X₄₉ = I, V, or absent; X₅₀ = A, G, or absent; X₅₁ = T, V, or absent; X₅₂ = Y or absent; X₅₃ = I, T, or absent; X₅₄ = T, I, or L; X₅₅ = G or absent; X₅₆ = A or absent; X₅₇ = A or S; X₅₈ = S or T; X₅₉ = I or V; X₆₀ = S or Y; X₆₁ = N or S; X₆₂ = W, Y, or N; X₆₃ = absent or N; X₆₄ = absent or E; X₆₅ = A or S; X₆₆ = S, A, or D; X₆₇ = A or T; X₆₈ = Y or T; X₆₉ = E or A; X₇₀ = C or G; X₇₁ = T, S, or I; X₇₂ = G or Y; X₇₃ = S or I; X₇₄ = S or absent; X₇₅ = D or Y; X₇₆ = S or W; X₇₇ = G or Y; X₇₈ = W or A; X₇₉ = D, T, or absent</p>		

Table 2A		
15C4.1D8.1G2 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	<u>GYTF</u> <u>TDHAIH</u> (IMGT) GYTF <u>TDHAIH</u> (Kabat) <u>GYTF</u> <u>TDHAIH</u> (Chothia)	169
CDR-H2 amino acid sequence (combined overlap)	<u>YFSPGN</u> <u>GDIKYNEKFKG</u> (IMGT) <u>YFSPGN</u> <u>GDIKYNEKFKG</u> (Kabat) Y <u>FSPGN</u> <u>GDIKYNEKFKG</u> (Chothia)	170
CDR-H3 amino acid sequence (combined overlap)	<u>KRSLP</u> <u>GPMD</u> <u>C</u> (IMGT) KR <u>SLP</u> <u>GPMD</u> <u>C</u> (Kabat) KR <u>SLP</u> <u>GPMD</u> <u>C</u> (Chothia)	171
CDR-L1 amino acid sequence (combined overlap)	K <u>ASENV</u> <u>GIYVS</u> (IMGT) <u>KASENV</u> <u>GIYVS</u> (Kabat) <u>KASENV</u> <u>GIYVS</u> (Chothia)	172
CDR-L2 amino acid sequence (combined overlap)	<u>GPSN</u> <u>RYT</u> (IMGT) <u>GPSN</u> <u>RYT</u> (Kabat) <u>GPSN</u> <u>RYT</u> (Chothia)	173
CDR-L3 amino acid sequence (combined overlap)	<u>GQSY</u> <u>SYPFT</u> (IMGT) <u>GQSY</u> <u>SYPFT</u> (Kabat) <u>GQSY</u> <u>SYPFT</u> (Chothia)	174

Table 2B		
8H3.2B9.2C7 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	<u>GYTF</u> <u>TDHAIH</u> (IMGT) GYTF <u>TDHAIH</u> (Kabat) <u>GYTF</u> <u>TDHAIH</u> (Chothia)	175

Table 2B		
8H3.2B9.2C7 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H2 amino acid sequence (combined overlap)	<u>YFSPGN</u> GDIKYNEKFKD (IMGT)	176
	<u>YFSPGN</u> GDIKYNEKFKD (Kabat)	
	Y <u>FSPGN</u> GDIKYNEKFKD (Chothia)	
CDR-H3 amino acid sequence (combined overlap)	<u>KRSLPG</u> DFDY (IMGT)	177
	KR <u>SLPG</u> DFDY (Kabat)	
	KR <u>SLPG</u> DFDY (Chothia)	
CDR-L1 amino acid sequence (combined overlap)	RAS <u>KSVSE</u> YLA (IMGT)	178
	<u>RASKSV</u> SEYLA (Kabat)	
	<u>RASKSV</u> SEYLA (Chothia)	
CDR-L2 amino acid sequence (combined overlap)	<u>SGSTL</u> LHS (IMGT)	179
	<u>SGSTL</u> LHS (Kabat)	
	<u>SGSTL</u> LHS (Chothia)	
CDR-L3 amino acid sequence (combined overlap)	<u>QQHNE</u> YPFT (IMGT)	180
	<u>QQHNE</u> YPFT (Kabat)	
	<u>QQHNE</u> YPFT (Chothia)	

Table 2C		
16E12.1D9.1B11 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	<u>GYTF</u> TDHAIH (IMGT)	181
	GYTF <u>TDH</u> AIH (Kabat)	
	<u>GYTF</u> TDHAIH (Chothia)	
CDR-H2 amino acid sequence (combined overlap)	<u>YFSPGN</u> DDVRYSEKFKG (IMGT)	182
	<u>YFSPGN</u> DDVRYSEKFKG (Kabat)	
	Y <u>FSPGN</u> DDVRYSEKFKG (Chothia)	
CDR-H3 amino acid sequence (combined overlap)	<u>KRSLPG</u> DFDY (IMGT)	183
	KR <u>SLPG</u> DFDY (Kabat)	
	KR <u>SLPG</u> DFDY (Chothia)	

Table 2C		
16E12.1D9.1B11 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-L1 amino acid sequence (combined overlap)	<u>RASKSINNYLV</u> (IMGT)	184
	<u>RASKSINNYLV</u> (Kabat)	
	<u>RASKSINNYLV</u> (Chothia)	
CDR-L2 amino acid sequence (combined overlap)	<u>SGSTLQT</u> (IMGT)	185
	<u>SGSTLQT</u> (Kabat)	
	<u>SGSTLQT</u> (Chothia)	
CDR-L3 amino acid sequence (combined overlap)	<u>QQHNEY PFT</u> (IMGT)	186
	<u>QQHNEY PFT</u> (Kabat)	
	<u>QQHNEY PFT</u> (Chothia)	

Table 2D		
14E9 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	<u>GIDFSSYWIC</u> (IMGT)	187
	GIDFSS <u>YWIC</u> (Kabat)	
	<u>GIDFSSYWIC</u> (Chothia)	
CDR-H2 amino acid sequence (combined overlap)	<u>CIYTGSGGNTYYATWAKG</u> (IMGT)	188
	<u>CIYTGSGGNTYYATWAKG</u> (Kabat)	
	<u>CIYTGSGGNTYYATWAKG</u> (Chothia)	
CDR-H3 amino acid sequence (combined overlap)	<u>ARMGYSAGYIGATYITVGAFNL</u> (IMGT)	189
	ARMGYSAGYIGATYITVGAFNL (Kabat)	
	<u>ARMGYSAGYIGATYITVGAFNL</u> (Chothia)	
CDR-L1 amino acid sequence (combined overlap)	QAS <u>Q</u> SISNWLA (IMGT)	190
	<u>QASQ</u> SISNWLA (Kabat)	
	<u>QASQ</u> SISNWLA (Chothia)	

Table 2D		
14E9 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-L2 amino acid sequence (combined overlap)	<u>SASYLES</u> (IMGT)	191
	<u>SASYLES</u> (Kabat)	
	<u>SASYLES</u> (Chothia)	
CDR-L3 amino acid sequence (combined overlap)	<u>QCTYGSSGDSGSWD</u> (IMGT)	192
	<u>QCTYGSSGDSGSWD</u> (Kabat)	
	<u>QCTYGSSGDSGSWD</u> (Chothia)	

Table 2E		
19H2 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	<u>GVAFSGSQWIC</u> (IMGT)	193
	GVAFSGSQWIC (Kabat)	
	<u>GVAFSGSQWIC</u> (Chothia)	
CDR-H2 amino acid sequence (combined overlap)	<u>CIYTGSSATDYANWARG</u> (IMGT)	194
	<u>CIYTGSSATDYANWARG</u> (Kabat)	
	<u>CIYTGSSATDYANWARG</u> (Chothia)	
CDR-H3 amino acid sequence (combined overlap)	<u>ARMGYEDGYVGGVYTIIVGAFNL</u> (IMGT)	195
	<u>ARMGYEDGYVGGVYTIIVGAFNL</u> (Kabat)	
	<u>ARMGYEDGYVGGVYTIIVGAFNL</u> (Chothia)	
CDR-L1 amino acid sequence (combined overlap)	<u>QASQTISSYLA</u> (IMGT)	196
	<u>QASQTISSYLA</u> (Kabat)	
	<u>QASQTISSYLA</u> (Chothia)	
CDR-L2 amino acid sequence (combined overlap)	<u>ATSYLES</u> (IMGT)	197
	<u>ATSYLES</u> (Kabat)	
	<u>ATSYLES</u> (Chothia)	

Table 2E		
19H2 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-L3 amino acid sequence (combined overlap)	<u>QCSYGSGYSGSWT</u> (IMGT)	198
	<u>QCSYGSGYSGSWT</u> (Kabat)	
	<u>QCSYGSGYSGSWT</u> (Chothia)	

Table 2F		
39A3 IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	<u>GLDFSGIYWAC</u> (IMGT)	199
	<u>GLDFSGIYWAC</u> (Kabat)	
	<u>GLDFSGIYWAC</u> (Chothia)	
CDR-H2 amino acid sequence (combined overlap)	<u>CMDNRVTYATWAKG</u> (IMGT)	200
	<u>CMDNRVTYATWAKG</u> (Kabat)	
	<u>CMDNRVTYATWAKG</u> (Chothia)	
CDR-H3 amino acid sequence (combined overlap)	<u>ARGGYGGRGLVFNL</u> (IMGT)	201
	<u>ARGGYGGRGLVFNL</u> (Kabat)	
	<u>ARGGYGGRGLVFNL</u> (Chothia)	
CDR-L1 amino acid sequence (combined overlap)	<u>QSSQSVYNNNELS</u> (IMGT)	202
	<u>QSSQSVYNNNELS</u> (Kabat)	
	<u>QSSQSVYNNNELS</u> (Chothia)	
CDR-L2 amino acid sequence (combined overlap)	<u>DASTLAS</u> (IMGT)	203
	<u>DASTLAS</u> (Kabat)	
	<u>DASTLAS</u> (Chothia)	
CDR-L3 amino acid sequence (combined overlap)	<u>QGIYYIGDWYSA</u> (IMGT)	204
	<u>QGIYYIGDWYSA</u> (Kabat)	
	<u>QGIYYIGDWYSA</u> (Chothia)	

Table 2G		
Group 1 Consensus - IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	GYTFTDHAIH	205
CDR-H2 amino acid sequence (combined overlap)	YFSPGNX ₁ DX ₂ X ₃ YX ₄ EKF ₅	206
CDR-H3 amino acid sequence (combined overlap)	KRSLPGX ₆ X ₇ DX ₈	207
CDR-L1 amino acid sequence (combined overlap)	X ₉ ASX ₁₀ X ₁₁ X ₁₂ X ₁₃ X ₁₄ YX ₁₅ X ₁₆	208
CDR-L2 amino acid sequence (combined overlap)	X ₁₇ X ₁₈ SX ₁₉ X ₂₀ X ₂₁ X ₂₂	209
CDR-L3 amino acid sequence (combined overlap)	X ₂₃ QX ₂₄ X ₂₅ X ₂₆ YPFT	210
X ₁ = G or D; X ₂ = I or V; X ₃ = K or R; X ₄ = N or S; X ₅ = G or D; X ₆ = P or D; X ₇ = M or F; X ₈ = C or Y; X ₉ = K or R; X ₁₀ = E or K; X ₁₁ = N or S; X ₁₂ = V or I; X ₁₃ = G, S, or N; X ₁₄ = I, E, or N; X ₁₅ = V or L; X ₁₆ = S, A, or V; X ₁₇ = G or S; X ₁₈ = P or G; X ₁₉ = N or T; X ₂₀ = R or L; X ₂₁ = Y, H, or Q; X ₂₂ = T or S; X ₂₃ = G or Q; X ₂₄ = S or H; X ₂₅ = Y or N; X ₂₆ = S or E		

Table 2H		
Group 2 - IMGT, Kabat, and Chothia CDR combined overlap sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (combined overlap)	GX ₂₇ X ₂₈ FSX ₂₉ X ₃₀ X ₃₁ WX ₃₂ C	211
CDR-H2 amino acid sequence (combined overlap)	CX ₃₃ X ₃₄ X ₃₅ X ₃₆ X ₃₇ X ₃₈ X ₃₉ X ₄₀ X ₄₁ X ₄₂ YAX ₄₃ WAX ₄₄ G	212
CDR-H3 amino acid sequence (combined overlap)	ARX ₄₅ GYX ₄₆ X ₄₇ GX ₄₈ X ₄₉ GX ₅₀ X ₅₁ X ₅₂ X ₅₃ X ₅₄ 4VX ₅₅ X ₅₆ FNL	213
CDR-L1 amino acid sequence (combined overlap)	QX ₅₇ SQX ₅₈ X ₅₉ X ₆₀ X ₆₁ X ₆₂ X ₆₃ X ₆₄ LX ₆₅	214
CDR-L2 amino acid sequence (combined overlap)	X ₆₆ X ₆₇ SX ₆₈ LX ₆₉ S	215
CDR-L3 amino acid sequence (combined overlap)	QX ₇₀ X ₇₁ YX ₇₂ X ₇₃ X ₇₄ GX ₇₅ X ₇₆ X ₇₇ SX ₇₈ X ₇₉	216
X ₂₇ = I, V, or L; X ₂₈ = D or A; X ₂₉ = absent or G; X ₃₀ = S or I; X ₃₁ = Y or Q; X ₃₂ = I or A; X ₃₃ = I or M; X ₃₄ = Y or D; X ₃₅ = T or N; X ₃₆ = G or R; X ₃₇ = S, V, or absent; X ₃₈ = G, S, or absent; X ₃₉ = G, A, or absent; X ₄₀ = N, T, or absent; X ₄₁ = T, D, or absent; X ₄₂ = Y or absent; X ₄₃ = T or N; X ₄₄ = K or R; X ₄₅ = M or G; X ₄₆ = S, E, or G; X ₄₇ = A, D, or absent; X ₄₈ = Y or R; X ₄₉ = I, V, or absent; X ₅₀ = A, G, or absent; X ₅₁ = T, V, or absent; X ₅₂ = Y or absent; X ₅₃ = I, T, or absent; X ₅₄ = T, I, or L; X ₅₅ = G or absent; X ₅₆ = A or absent; X ₅₇ = A or S; X ₅₈ = S or T; X ₅₉ = I or V; X ₆₀ = S or Y; X ₆₁ = N or S; X ₆₂ = W, Y, or N; X ₆₃ = absent or N; X ₆₄ = absent or E; X ₆₅ = A or S; X ₆₆ = S, A, or D; X ₆₇ = A or T; X ₆₈ = Y or T; X ₆₉ = E or A; X ₇₀ = C or G; X ₇₁ = T, S, or I; X ₇₂ = G or Y; X ₇₃ = S or I; X ₇₄ = S or absent; X ₇₅ = D or Y; X ₇₆ = S or W; X ₇₇ = G or Y; X ₇₈ = W or A; X ₇₉ = D, T, or absent		

Table 3A		
15C4.1D8.1G2 IMGT, Kabat, and Chothia CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	DH	217
CDR-H2 amino acid sequence (common sequence)	SPGNGD	218
CDR-H3 amino acid sequence (common sequence)	SLPGPMDC	219
CDR-L1 amino acid sequence (common sequence)	ENVGIY	220
CDR-L2 amino acid sequence (common sequence)	GPS	221
CDR-L3 amino acid sequence (common sequence)	GQSYSYPFT	222

Table 3B		
8H3.2B9.2C7 IMGT, Kabat, and Chothia CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	DH	223
CDR-H2 amino acid sequence (common sequence)	SPGNGD	224
CDR-H3 amino acid sequence (common sequence)	SLPGDFDY	225
CDR-L1 amino acid sequence (common sequence)	KSVSEY	226
CDR-L2 amino acid sequence (common sequence)	SGS	227
CDR-L3 amino acid sequence (common sequence)	QQHNEYPFT	228

Table 3C		
16E12.1D9.1B11 IMGT, Kabat, and Chothia CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	DH	229
CDR-H2 amino acid sequence (common sequence)	SPGNDD	230
CDR-H3 amino acid sequence (common sequence)	SLPGDFDY	231
CDR-L1 amino acid sequence (common sequence)	KSINNY	232
CDR-L2 amino acid sequence (common sequence)	SGS	233
CDR-L3 amino acid sequence (common sequence)	QQHNEYPFT	234

Table 3D		
14E9 IMGT, Kabat, and Chothia CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	SY	235
CDR-H2 amino acid sequence (common sequence)	YTGSGGN	236
CDR-H3 amino acid sequence (common sequence)	MGYSAGYIGATYITVGAFNL	237
CDR-L1 amino acid sequence (common sequence)	QISISNW	238
CDR-L2 amino acid sequence (common sequence)	SAS	239
CDR-L3 amino acid sequence (common sequence)	QCTYGSSGDSGSWD	240

Table 3E		
19H2 IMGT, Kabat, and Chothia CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	GSQ	241
CDR-H2 amino acid sequence (common sequence)	YTGSSAT	242
CDR-H3 amino acid sequence (common sequence)	MGYEDGYVGGVYTIVGAFNL	243
CDR-L1 amino acid sequence (common sequence)	QTISSY	244
CDR-L2 amino acid sequence (common sequence)	ATS	245
CDR-L3 amino acid sequence (common sequence)	QCSYGSGYSGSWT	246

Table 3F		
39A3 IMGT, Kabat, and Chothia CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	GIY	247
CDR-H2 amino acid sequence (common sequence)	DNR	248
CDR-H3 amino acid sequence (common sequence)	GGYGGRGLVFNL	249
CDR-L1 amino acid sequence (common sequence)	QSVYNNNE	250
CDR-L2 amino acid sequence (common sequence)	DAS	251
CDR-L3 amino acid sequence (common sequence)	QGIYYIGDWYSA	252

Table 3G		
Group 1 Consensus CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	DH	253
CDR-H2 amino acid sequence (common sequence)	SPGNX ₁ D	254
CDR-H3 amino acid sequence (common sequence)	SLPGX ₆ X ₇ DX ₈	255
CDR-L1 amino acid sequence (common sequence)	X ₁₀ X ₁₁ X ₁₂ X ₁₃ X ₁₄ Y	256
CDR-L2 amino acid sequence (common sequence)	X ₁₇ X ₁₈ S	257
CDR-L3 amino acid sequence (common sequence)	X ₂₃ QX ₂₄ X ₂₅ X ₂₆ YPFT	258
X ₁ = G or D; X ₆ = P or D; X ₇ = M or F; X ₈ = C or Y; X ₁₀ = E or K; X ₁₁ = N or S; X ₁₂ = V or I; X ₁₃ = G, S, or N; X ₁₄ = I, E, or N; X ₁₇ = G or S; X ₁₈ = P or G; X ₂₃ = G or Q; X ₂₄ = S or H; X ₂₅ = Y or N; X ₂₆ = S or E		

Table 3H		
Group 2 Consensus CDR common sequences		
Description	Sequence	SEQ ID NO:
CDR-H1 amino acid sequence (common sequence)	X ₂₉ X ₃₀ X ₃₁	259
CDR-H2 amino acid sequence (common sequence)	X ₃₄ X ₃₅ X ₃₆ X ₃₇ X ₃₈ X ₃₉ X ₄₀	260
CDR-H3 amino acid sequence (common sequence)	X ₄₅ GYX ₄₆ X ₄₇ GX ₄₈ X ₄₉ GX ₅₀ X ₅₁ X ₅₂ X ₅₃ X ₅₄ VX ₅₅ X ₅₆ FNL	261
CDR-L1 amino acid sequence (common sequence)	QX ₅₈ X ₅₉ X ₆₀ X ₆₁ X ₆₂ X ₆₃ X ₆₄	262
CDR-L2 amino acid sequence (common sequence)	X ₆₆ X ₆₇ S	263
CDR-L3 amino acid sequence (common sequence)	QX ₇₀ X ₇₁ YX ₇₂ X ₇₃ X ₇₄ GX ₇₅ X ₇₆ X ₇₇ SX ₇₈ X ₇₉	342
X ₂₉ = absent or G; X ₃₀ = S or I; X ₃₁ = Y or Q; X ₃₄ = Y or D; X ₃₅ = T or N; X ₃₆ = G or R; X ₃₇ = S, V, or absent; X ₃₈ = G, S, or absent; X ₃₉ = G, A, or absent; X ₄₀ = N, T, or absent; X ₄₅ = M or G; X ₄₆ = S, E, or G; X ₄₇ = A, D, or absent; X ₄₈ = Y or R; X ₄₉ = I, V, or absent; X ₅₀ = A, G, or absent; X ₅₁ = T, V, or absent; X ₅₂ = Y or absent; X ₅₃ = I, T, or absent; X ₅₄ = T, I, or L; X ₅₅ = G or absent; X ₅₆ = A or absent; X ₅₈ = S or T; X ₅₉ = I or V; X ₆₀ = S or Y; X ₆₁ = N or S; X ₆₂ = W, Y, or N; X ₆₃ = absent or N; X ₆₄ = absent or E; X ₆₆ = S, A, or D; X ₆₇ = A or T; X ₇₀ = C or G; X ₇₁ = T, S, or I; X ₇₂ = G or Y; X ₇₃ = S or I; X ₇₄ = S or absent; X ₇₅ = D or Y; X ₇₆ = S or W; X ₇₇ = G or Y; X ₇₈ = W or A; X ₇₉ = D, T, or absent		

Table 4A		
Humanized 8H3 Heavy Chain Sequences – Germline 1-3		
Description	Sequence	SEQ ID NO:
8H3-HV1-3-A	QVQLVQSGAEVKKPGASVKVSCKASGYTFTDHAHWW RQAPGQRLEWIGYFSPGNGDIKYNEKFKDRATLTADK SASTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGT LTVSS	264

Table 4A Humanized 8H3 Heavy Chain Sequences – Germline 1-3		
Description	Sequence	SEQ ID NO:
8H3-HV1-3-B	QVQLVQSGAEVKKPGASVKVSKASGYTFTDHAIHWV RQAPGQRLEWIGYFSPGNNDIKYSQKFKGRVTITADKS ASTAYMELSSLRSED TAVYYCKRSLPGDFDYWGQGT LTVSS	265
8H3-HV1-3-C	QVQLVQSGAEVKKPGASVKVSKASGYTFTDHAIHWV RQAPGQRLEWIGYFSPGNADTKYSQKFQGRVTITADK SASTAYMELSSLRSED TAVYYCKRSLPGDFDYWGQGT LTVSS	266

Table 4B Humanized 8H3 Heavy Chain Sequences – Germline 1-69		
Description	Sequence	SEQ ID NO:
8H3-HV1-69-A	QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAIHWV RQAPGQGLEWIGYFSPGNNDIKYNEKFKDRATLTADK STSTAYMELSSLRSED TAVYFCKRSLPGDFDYWGQGT LTVSS	267
8H3-HV1-69-B	QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAIHWV RQAPGQGLEWIGYFSPGNNDIKYNQKFKGRVTITADK STSTAYMELSSLRSED TAVYYCKRSLPGDFDYWGQGT LTVSS	268
8H3-HV1-69-C	QVQLVQSGAEVKKPGSSVKVSKASGYTFSDHAIHWV RQAPGQGLEWIGYFSPGNADINYAQKFQGRVTITADK STSTAYMELSSLRSED TAVYYCKRSLPGDFDYWGQGT LTVSS	269

Table 4C Humanized 8H3 Heavy Chain Sequences – Germline 5		
Description	Sequence	SEQ ID NO:
8H3-HV5-A	EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWVR QMPGKGLEWIGYFSPGNNDIKYNEKFKDQATLSADKSI STAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LTVSS	270
8H3-HV5-B	EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWV RQMPGKGLEWIGYFSPGNNDIKYNEKFKGQVTISADK SISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGT LTVSS	271
8H3-HV5-C	EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWV RQMPGKGLEWIGYFSPGNADIRYSEKFQGGVTISADK SISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGT LTVSS	272

Table 4D Humanized 8H3 Heavy Chain Sequences – Germline 7		
Description	Sequence	SEQ ID NO:
8H3-HV7-A	QVQLVQSGSELKKPGASVKVSKASGYTFTDHAIHWV RQAPGQGLEWIGYFSPGNNDIKYNEKFKDRAVLSADK SVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LTVSS	273
8H3-HV7-B	QVQLVQSGSELKKPGASVKVSKASGYTFTDHAIHWV RQAPGQGLEWIGYISTGNGNDIKYNQKFTGRAVLSLTKS	274

Table 4D		
Humanized 8H3 Heavy Chain Sequences – Germline 7		
Description	Sequence	SEQ ID NO:
	VSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTV TVSS	
8H3-HV7-C	QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWV RQAPGQGLEWIGYISTGNANITYAQGFTGRAVLSDKS VSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTV TVSS	275

Table 4E		
Humanized 8H3 Light Chain Sequences – Germline 1		
Description	Sequence	SEQ ID NO:
8H3-KV1-A	DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQE KPGKANKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISS LQPEDFATYFCQQHNEYPTFGQGKLEIK	276
8H3-KV1-B	DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQ KPGKAPKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISS LQPEDFATYYCQQHNEYPTFGQGKLEIK	277
8H3-KV1-C	DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQK PGKAPKLLIYASTLHSGVPSRFSGSGSGTEFTLTISSL QPEDFATYYCQQHNEYPTFGQGKLEIK	278

Table 4F		
Humanized 8H3 Light Chain Sequences – Germline 3		
Description	Sequence	SEQ ID NO:
8H3-KV3-A	EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQE KPGQANRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISS LQSEDFAVYFCQQHNEYPTFGQGKLEIK	279
8H3-KV3-B	EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQ QKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTIS SLQSEDFAVYYCQQHNEYPTFGQGKLEIK	280
8H3-KV3-C	EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQ QKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTIS SLQSEDFAVYYCQQHNEYPTFGQGKLEIK	281

Table 4G		
Humanized 8H3 Light Chain Sequences – Germline 6		
Description	Sequence	SEQ ID NO:
8H3-KV6-A	EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQE KPDQSNKLLIYSGSTLHSGVPSRFSGSGSGTDFTLTIN SLEAEDAATYFCQQHNEYPTFGQGKLEIK	282
8H3-KV6-B	EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQ KPDQSPKLLIYSGSTLHSGVPSRFSGSGSGTDFTLTINS LEAEDAATYYCQQHNEYPTFGQGKLEIK	283
8H3-KV6-C	EIVLTQSPDFQSVTPKEKVTITCRASKSISYLAWYQQK PDQSPKLLIYSGSTLFSGVPSRFSGSGSGTDFTLTINSL EAEDAATYYCQQHNEYPTFGQGKLEIK	284

[0013] In certain aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises CDRs comprising the amino acid sequences of any of the CDR

combinations set forth in Tables 1A-3H. In certain embodiments, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises a CDR-H1 comprising the amino acid sequence of SEQ ID NO:253, a CDR-H2 comprising the amino acid sequence of SEQ ID NO:254, a CDR-H3 comprising the amino acid sequence of SEQ ID NO:255, a CDR-L1 comprising the amino acid sequence of SEQ ID NO:256, a CDR-L2 comprising the amino acid sequence of SEQ ID NO:257, and a CDR-L3 comprising the amino acid sequence of SEQ ID NO:258. In some embodiments, CDR-H1 comprises the amino acid sequence of SEQ ID NO:253. In some embodiments, CDR-H2 comprises the amino acid sequence of SEQ ID NO:254. In some embodiments, CDR-H3 comprises the amino acid sequence of SEQ ID NO:255. In some embodiments, CDR-L1 comprises the amino acid sequence of SEQ ID NO:256. In some embodiments, CDR-L2 comprises the amino acid sequence of SEQ ID NO:257. In some embodiments, CDR-L3 comprises the amino acid sequence of SEQ ID NO:258.

[0014] In certain embodiments, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises a CDR-H1 comprising the amino acid sequence of SEQ ID NO:259, a CDR-H2 comprising the amino acid sequence of SEQ ID NO:260, a CDR-H3 comprising the amino acid sequence of SEQ ID NO:261, a CDR-L1 comprising the amino acid sequence of SEQ ID NO:262, a CDR-L2 comprising the amino acid sequence of SEQ ID NO:263, and a CDR-L3 comprising the amino acid sequence of SEQ ID NO:342. In some embodiments, CDR-H1 comprises the amino acid sequence of SEQ ID NO:259. In some embodiments, CDR-H2 comprises the amino acid sequence of SEQ ID NO:260. In some embodiments, CDR-H3 comprises the amino acid sequence of SEQ ID NO:261. In some embodiments, CDR-L1 comprises the amino acid sequence of SEQ ID NO:262. In some embodiments, CDR-L2 comprises the amino acid sequence of SEQ ID NO:263. In some embodiments, CDR-L3 comprises the amino acid sequence of SEQ ID NO:342.

[0015] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:3-5 and light chain CDRs of SEQ ID NOS:6-8. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:9-11 and light chain CDRs of SEQ ID NOS:12-14. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:15-17 and light chain CDRs of SEQ ID NOS:18-20. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:169-171 and light chain CDRs of SEQ ID NOS:172-174.

[0016] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:25-27 and light chain CDRs of SEQ ID NOS:28-30. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of

the disclosure comprises heavy chain CDRs of SEQ ID NOS:31-33 and light chain CDRs of SEQ ID NOS:32-34. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:35-37 and light chain CDRs of SEQ ID NOS:38-40. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:175-177 and light chain CDRs of SEQ ID NOS:178-180.

[0017] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:47-49 and light chain CDRs of SEQ ID NOS:50-52. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:53-55 and light chain CDRs of SEQ ID NOS:56-58. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:59-61 and light chain CDRs of SEQ ID NOS:62-64. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:181-183 and light chain CDRs of SEQ ID NOS:184-186.

[0018] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:69-71 and light chain CDRs of SEQ ID NOS:72-74. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:75-77 and light chain CDRs of SEQ ID NOS:78-80. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:81-83 and light chain CDRs of SEQ ID NOS:84-86. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:187-189 and light chain CDRs of SEQ ID NOS:190-192.

[0019] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:91-93 and light chain CDRs of SEQ ID NOS:94-96. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:97-99 and light chain CDRs of SEQ ID NOS:100-102. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:103-105 and light chain CDRs of SEQ ID NOS:106-108. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:193-195 and light chain CDRs of SEQ ID NOS:196-198.

[0020] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:113-115 and light chain CDRs of SEQ ID NOS:116-118. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:119-121 and light chain CDRs of

SEQ ID NOS:122-124. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:125-127 and light chain CDRs of SEQ ID NOS:128-130. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:199-201 and light chain CDRs of SEQ ID NOS:202-204.

[0021] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:133-135 and light chain CDRs of SEQ ID NOS:136-138. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:139-141 and light chain CDRs of SEQ ID NOS:142-144. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:145-147 and light chain CDRs of SEQ ID NOS:148-150.

[0022] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:151-153 and light chain CDRs of SEQ ID NOS:154-156. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:157-159 and light chain CDRs of SEQ ID NOS:160-162. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:163-165 and light chain CDRs of SEQ ID NOS:166-168.

[0023] In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:205-207 and light chain CDRs of SEQ ID NOS:208-210. In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy chain CDRs of SEQ ID NOS:211-213 and light chain CDRs of SEQ ID NOS:214-216.

[0024] In certain embodiments, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises a CDR-H1 comprising the amino acid sequence of SEQ ID NO:133, 139, 145, 169, 175, 181, 205, 217, 223, 229, or 253; a CDR-H2 comprising the amino acid sequence of SEQ ID NO:134, 140, 146, 170, 176, 182, 206, 218, 224, 230, or 254; a CDR-H3 comprising the amino acid sequence of SEQ ID NO:135, 141, 147, 171, 177, 183, 207, 219, 225, 231, or 255; a CDR-L1 comprising the amino acid sequence of SEQ ID NO:136, 142, 148, 172, 178, 184, 208, 220, 226, 232, or 256; a CDR-L2 comprising the amino acid sequence of SEQ ID NO:137, 143, 149, 173, 179, 185, 209, 221, 227, 233, or 257; and a CDR-L3 comprising the amino acid sequence of SEQ ID NO:138, 144, 150, 174, 180, 186, 210, 222, 228, 234, or 258.

[0025] In certain embodiments, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises a CDR-H1 comprising the amino acid sequence of SEQ ID NO:151, 157, 163, 187, 193, 199, 211, 235, 241, 247, or 259; a CDR-H2 comprising the amino acid sequence of SEQ ID NO:152, 158, 164, 188, 194, 200, 212, 236, 242, 248, or 260; a CDR-H3 comprising

the amino acid sequence of SEQ ID NO:153, 159, 165, 189, 195, 201, 213, 237, 243, 249, or 261; a CDR-L1 comprising the amino acid sequence of SEQ ID NO:154, 160, 166, 190, 196, 202, 214, 238, 244, 250, or 262; a CDR-L2 comprising the amino acid sequence of SEQ ID NO:155, 161, 167, 191, 197, 203, 215, 239, 245, 251, or 263; and a CDR-L3 comprising the amino acid sequence of SEQ ID NO:156, 162, 168, 192, 198, 204, 216, 240, 246, 252, or 342.

[0026] The antibodies and antigen-binding fragments of the disclosure can be murine, chimeric, humanized or human.

[0027] In further aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising heavy and light chain variable regions of SEQ ID NOS:1 and 2, respectively. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS:1 and 2, respectively.

[0028] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising heavy and light chain variable regions of SEQ ID NOS:23 and 24, respectively. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS:23 and 24, respectively.

[0029] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising heavy and light chain variable regions of SEQ ID NOS:45 and 46, respectively. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS:45 and 46, respectively.

[0030] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising heavy and light chain variable regions of SEQ ID NOS:67 and 68, respectively. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS:67 and 68, respectively.

[0031] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising heavy and light chain variable regions of SEQ ID NOS:89 and 90, respectively. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light

chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS:89 and 90, respectively.

[0032] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising heavy and light chain variable regions of SEQ ID NOS:111 and 112, respectively. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS:111 and 112, respectively.

[0033] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising a heavy chain variable region of any one of SEQ ID NOS:133-144 and a light chain variable region of any one of SEQ ID NOS:145-153. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having a heavy variable region having at least 95%, 98%, 99%, or 99.5% sequence identity of any one of SEQ ID NOS:133-134 and a light variable region having at least 95%, 98%, 99%, or 99.5% sequence identity of any one of SEQ ID NOS:145-153.

[0034] In yet other aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with an antibody or antigen binding fragment comprising a heavy chain variable region of any one of SEQ ID NOS:264-275 and a light chain variable region of any one of SEQ ID NOS:276-284. In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having a heavy variable region having at least 95%, 98%, 99%, or 99.5% sequence identity of any one of SEQ ID NOS:264-275 and a light variable region having at least 95%, 98%, 99%, or 99.5% sequence identity of any one of SEQ ID NOS:276-284.

[0035] In yet other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure is a single-chain variable fragment (scFv). An exemplary scFv comprises the heavy chain variable fragment N-terminal to the light chain variable fragment. In some embodiments, the scFv heavy chain variable fragment and light chain variable fragment are covalently bound to a linker sequence of 4-15 amino acids. The scFv can be in the form of a bi-specific T-cell engager or within a chimeric antigen receptor (CAR).

[0036] The anti-glyco-cMET antibodies and antigen-binding fragments can be in the form of a multimer of a single-chain variable fragment, a bispecific single-chain variable fragment and a multimer of a bispecific single-chain variable fragment. In some embodiments, the multimer of a single chain variable fragment is selected a divalent single-chain variable fragment, a tribody or a tetrabody. In some of these embodiments, the multimer of a bispecific single-chain variable fragment is a bispecific T-cell engager.

[0037] Other aspects of the disclosure are drawn to nucleic acids encoding the anti-glyco-cMET antibodies and antibody-binding fragments of the disclosure. In some embodiments, the

portion of the nucleic acid nucleic acid encoding an anti-glyco-cMET antibody or antigen-binding fragment is codon-optimized for expression in a human cell. In certain aspects, the disclosure provides an anti-glyco-cMET antibody or antigen binding fragment having heavy and light chain variable regions encoded by a heavy chain nucleotide sequence having at least 95%, 98%, 99%, or 99.5% sequence identity to SEQ ID NO:21, 43, or 65 and a light chain nucleotide sequence having at least 95%, 98%, 99%, or 99.5% sequence identity to SEQ ID NO:22, 44 or 66. In other aspects, the disclosure provides an anti-glyco-cMET antibody or antigen binding fragment having heavy and light chain variable regions encoded by a heavy chain nucleotide sequence having at least 95%, 98%, 99%, or 99.5% sequence identity to SEQ ID NO:87, 109, or 131 and a light chain nucleotide sequence having at least 95%, 98%, 99%, or 99.5% sequence identity to SEQ ID NO:88, 110 or 132. Vectors (*e.g.*, a viral vector such as a lentiviral vector) and host cells comprising the nucleic acids are also within the scope of the disclosure. The heavy and light chains coding sequences can be present on a single vector or on separate vectors.

[0038] In yet another aspect of the disclosure is a pharmaceutical composition comprising an anti-glyco-cMET antibody, antigen-binding fragment, nucleic acid (or pair of nucleic acids), vector (or pair of vectors) or host cell according to the disclosure, and a physiologically suitable buffer, adjuvant or diluent.

[0039] Still another aspect of the disclosure is a method of making a chimeric antigen receptor comprising incubating a cell comprising a nucleic acid or a vector according to the disclosure, under conditions suitable for expression of the coding region and collecting the chimeric antigen receptor.

[0040] Another aspect of the disclosure is a method of detecting cancer comprising contacting a biological sample (*e.g.*, a cell, tissue sample, or extracellular vesicle) with an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure and detecting whether the antibody is bound to the biological sample (*e.g.*, cell, tissue sample, or extracellular vesicle).

[0041] Yet another aspect of the disclosure is an anti-glyco-cMET antibody or antigen-binding fragment according to the disclosure of the disclosure for use in detecting cancer.

[0042] Yet another aspect of the disclosure is a method of treating cancer comprising administering a prophylactically or therapeutically effective amount of an anti-glyco-cMET antibody, antigen-binding fragment, nucleic acid, vector, host cell or pharmaceutical composition according to the disclosure to a subject in need thereof.

[0043] Yet another aspect of the disclosure is an anti-glyco-cMET antibody, antigen-binding fragment, nucleic acid, vector, host cell or pharmaceutical composition according to the disclosure for use in the treatment of cancer.

[0044] Yet another aspect of the disclosure is use of an anti-glyco-cMET antibody, antigen-binding fragment, nucleic acid, vector, host cell or pharmaceutical composition according to the disclosure for the manufacture of a medicament for the treatment of cancer.

[0045] Glyco-cMET peptides are also provided herein. The peptides can be 13-30 amino acids in length and comprise amino acids 1-11, 1-12, 1-13, 1-14, 1-15, 1-16, 1-17, 1-18, 1-19, 1-20, 2-11, 2-12, 2-13, 2-14, 2-15, 2-16, 2-17, 2-18, 2-19, 2-20, 3-11, 3-12, 3-13, 3-14, 3-15, 3-16, 3-17, 3-18, 3-19, 3-20, 4-11, 4-12, 4-13, 4-14, 4-15, 4-16, 4-17, 4-18, 4-19, 4-20, 5-11, 5-12, 5-13, 5-14, 5-15, 5-16, 5-17, 5-18, 5-19, 5-20, 6-11, 6-12, 6-13, 6-14, 6-15, 6-16, 6-17, 6-18, 6-19, 6-20, 7-11, 7-12, 7-13, 7-14, 7-15, 7-16, 7-17, 7-18, 7-19, 7-20, 8-11, 8-12, 8-13, 8-14, 8-15, 8-16, 8-17, 8-18, 8-19, 8-20, 9-11, 9-12, 9-13, 9-14, 9-15, 9-16, 9-17, 9-18, 9-19, or 9-20 of SEQ ID NO:285 (PTKSFISGG**ST**ITGVGKLN, glycosylated with GalNAc on the serine and threonine residues shown in bold underlined text). The glyco-cMET peptides are describe in Section 5.10 and numbered embodiments 894 to 920. The peptides can be included in a composition, as described in Section 5.10.1 and numbered embodiments 921 and 922. The glyco-cMET peptides can be used in methods for producing antibodies in an animal and/or eliciting an immune response in an animal. Methods for using the glyco-cMET peptides are described in Section 5.10.2 and numbered embodiments 923 to 926.

4. BRIEF DESCRIPTION OF THE FIGURES

[0046] FIG. 1: ELISA of 14E9, 19H2, and 39A3 rabbit antibodies against 50 ng of Tn-glycosylated cMET and Syndecan2 peptides.

[0047] FIGS. 2A-2B-5: Flow cytometry analysis of cMET mouse antibodies on A549 COSMC-KO and A549 cells. FIG. 2A: Representative histogram of 15C4.1D8.1G2, 8H3.2B9.2C7, and 16E12.1D9.1B11, anti-Golgi, mouse IgG isotype control, and anti-cMET antibodies on A549 COSMC-KO and A549 cells. FIGS. 2B1-2B-4: Titration of 15C4.1D8.1G2, 8H3.2B9.2C7, and 16E12.1D9.1B11 on cell surface antigens found on A549 COSMC-KO and A549 cells. FIG. 2B-5: legend for FIG. 2B-1 to 2B-4.

[0048] FIGS. 3A-3B-5: Flow cytometry analysis of cMET rabbit antibodies on A549 COSMC-KO and A549 cells. FIG. 3A: Representative histograms for staining of 14E9, 19H2, and 39A3, anti-Golgi, mouse IgG isotype control, and anti-cMET antibodies on A549 COSMC-KO and A549 cells. FIG. 3B-1-3B-4: Titration of 14E9, 19H2, and 39A3 on cell surface antigens found on A549 COSMC-KO and A549 cells. FIG. 3B-5: legend for FIG. 3B-1 to 3B-4.

[0049] FIGS. 4A-4C: Immunofluorescence staining of cMET mouse and rabbit antibodies. FIG. 4A-4B: Immunofluorescence staining of 15C4.1D8.1G2, 8H3.2B9.2C7, 16E12.1D9.1B11, anti-cMET, and anti-Tn antibodies on A549 and A549 COSMC-KO cells (FIG. 4A) and/or T47D and T47D COSMC-KO cells (FIG. 4B). FIG. 4C: Immunofluorescence staining of 14E9, 19H2, 39A3, anti-cMET, and anti-Tn antibodies on A549 COSMC-KO and A549 cells.

[0050] FIGS. 5A-5B: Immunohistochemistry of cMET mouse and rabbit antibodies. FIG. 5A: Staining of 15C4.1D8.1G2, 8H3.2B9.2C7, 16E12.1D9.1B11, 14E9, 19H2, 39A3, and IgG control antibodies on colon cancer and normal tissues (TMA-T051b Biomax). FIG. 5B: Statistics of positive and negative stained tissues.

[0051] FIGS. 6A-1-6B-2: Immunohistochemistry of cMET mouse antibodies. FIG. 6A-1: Staining of 8H3.2B9.2C7 (“GO-8H3”) antibody on ovarian (TMA-OV1502), pancreas (TMA-PA2082), lung (TMA-LC121b), and cholangiocarcinoma cancer (TMA-GA802a). Statistics shown in FIG. 6A-2. Positive samples had ~70% of cancer cells that had strong cellular surface stain. About 10-20% of analyzed cancer tissue had specific cellular surface stain ~70% of cancer cells. FIG. 6B-1: Staining of 8H3.2B9.2C7 (“GO-8H3”) on normal tissue (TMA-FDA999x). Statistics shown in FIG. 6B-2. No specific cellular surface staining was observed on normal tissue.

[0052] FIGS. 7A-7C: Cell killing assay of cMET CARTs. FIG. 7A: Killing of cMET CARTs (8H3.2B9.2C7) on A673 COSMC-KO and A673 target cells with a titration of ratios of T cells to target cells (1, 5, and 10). FIG. 7B: Summary of time for cMET-CARTs to kill 50% of the target cells A673 COSMC-KO (KT50). N/A = 50% of cells were not killed. FIG. 7C: Killing of cMET CARTs (8H3.2B9.2C7) on various cell lines with low (less than 500 receptors per cell) and high cMET-Tn expression (2×10^5 receptors per cell). cMET-CART demonstrated cytotoxicity in cells with low levels of cMET-Tn expression (A549-wt cells). Experiments were conducted with a titration of ratios of T cells to target cells (1, 5, and 10).

[0053] FIG. 8A-8B: In vivo activity of cMET-CART (8H3) in solid tumor mouse models. FIG. 8A: A549 solid tumor model (Lung cancer cell line) established by flank injection (CDx). The tumor volume at CART injection was 88 mm³. Mice were treated with 2nd generation 8H3-CAR-T by IT injection (2 doses at 1×10^7 cells). Tumor volume was measured by caliper. FIG. 8B: Lung cancer solid tumor model (PDx) established by flank injection (Champions model CTG-2823). The tumor volume at CART injection was 200 mm³ and CART was delivered by IV injection (4 doses at 1×10^7 cells).

[0054] FIGS. 9A-9C: Exemplary cMET-CART constructs 15C4-CART (FIG. 9A), 16E12-CART (FIG. 9B), and 8H3-CART (FIG. 9C). Testing of the constructs is described in Example 5.

[0055] FIG. 10: Cytotoxicity of hu8H3-CART on A673 (Tn+) and (Tn-) cells at an E:T ratio of 2:1.

5. DETAILED DESCRIPTION

5.1 Antibodies

[0056] The disclosure provides novel antibodies that are directed to a glycoform of cMET present on tumor cells. These are exemplified by the antibodies 15C4.1D8.1G2 (hereinafter, “15C4”), 8H3.2B9.2C7 (hereinafter, “8H3”), 16E12.1D9.1B11 (hereinafter, “16E12”), 14E9,

19H2, and 39A3. 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 were identified in a screen for antibodies that bind to a glycosylated peptide present in cMET: PTKSFISGGSTITGVGKLN (SEQ ID NO:285), glycosylated with GalNAc on the serine and threonine residues shown in bold underlined text so as to mimic the glycosylation pattern of cMET present on tumor cells.

[0057] The anti-glyco-cMET antibodies of the disclosure, exemplified by antibodies 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3, are useful as tools in cancer diagnosis and therapy.

[0058] Thus, in certain aspects, the disclosure provides antibodies and antigen binding fragments that bind to a glycoform of cMET present on tumor cells (referred to herein as “glyco-cMET”), and preferably to the peptide PTKSFISGGSTITGVGKLN (SEQ ID NO:285), glycosylated with GalNAc on the serine and threonine residues shown in bold underlined text.

[0059] The anti-glyco-cMET antibodies of the disclosure may be polyclonal, monoclonal, genetically engineered, and/or otherwise modified in nature, including but not limited to chimeric antibodies, humanized antibodies, human antibodies, primatized antibodies, single chain antibodies, bispecific antibodies, dual-variable domain antibodies, *etc.* In various embodiments, the antibodies comprise all or a portion of a constant region of an antibody. In some embodiments, the constant region is an isotype selected from: IgA (*e.g.*, IgA₁ or IgA₂), IgD, IgE, IgG (*e.g.*, IgG₁, IgG₂, IgG₃ or IgG₄), and IgM. In specific embodiments, the anti-glyco-cMET antibodies of the disclosure comprise an IgG₁ constant region isotype.

[0060] The term “monoclonal antibody” as used herein is not limited to antibodies produced through hybridoma technology. A monoclonal antibody is derived from a single clone, including any eukaryotic, prokaryotic, or phage clone, by any means available or known in the art. Monoclonal antibodies useful with the present disclosure can be prepared using a wide variety of techniques known in the art including the use of hybridoma, recombinant, and phage display technologies, or a combination thereof. In many uses of the present disclosure, including *in vivo* use of the anti-glyco-cMET antibodies in humans, chimeric, primatized, humanized, or human antibodies can suitably be used.

[0061] The term “chimeric” antibody as used herein refers to an antibody having variable sequences derived from a non-human immunoglobulin, such as a rat or a mouse antibody, and human immunoglobulin constant regions, typically chosen from a human immunoglobulin template. Methods for producing chimeric antibodies are known in the art. See, *e.g.*, Morrison, 1985, *Science* 229(4719):1202-7; Oi *et al.*, 1986, *BioTechniques* 4:214-221; Gillies *et al.*, 1985, *J. Immunol. Methods* 125:191-202; U.S. Pat. Nos. 5,807,715; 4,816,567; and 4,816,397, which are incorporated herein by reference in their entireties.

[0062] “Humanized” forms of non-human (*e.g.*, murine) antibodies are chimeric immunoglobulins that contain minimal sequences derived from non-human immunoglobulin. In general, a humanized antibody will comprise substantially all of at least one, and typically two,

variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework regions (FR) are those of a human immunoglobulin sequence. The humanized antibody can also comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin consensus sequence. Methods of antibody humanization are known in the art. See, *e.g.*, Riechmann *et al.*, 1988, *Nature* 332:323-7; U.S. Pat. Nos. 5,530,101; 5,585,089; 5,693,761; 5,693,762; and 6,180,370 to Queen *et al.*; EP239400; PCT publication WO 91/09967; U.S. Pat. No. 5,225,539; EP592106; EP519596; Padlan, 1991, *Mol. Immunol.*, 28:489-498; Studnicka *et al.*, 1994, *Prot. Eng.* 7:805-814; Roguska *et al.*, 1994, *Proc. Natl. Acad. Sci.* 91:969-973; and U.S. Pat. No. 5,565,332, all of which are hereby incorporated by reference in their entireties.

[0063] Exemplary humanized sequences are described in numbered embodiments 8 to 115. The variable region sequences for exemplary humanized antibodies and antigen-binding fragments thereof of the disclosure are set forth in Tables 4A-4G.

[0064] "Human antibodies" include antibodies having the amino acid sequence of a human immunoglobulin and include antibodies isolated from human immunoglobulin libraries or from animals transgenic for one or more human immunoglobulin and that do not express endogenous immunoglobulins. Human antibodies can be made by a variety of methods known in the art including phage display methods using antibody libraries derived from human immunoglobulin sequences. See U.S. Pat. Nos. 4,444,887 and 4,716,111; and PCT publications WO 98/46645; WO 98/50433; WO 98/24893; WO 98/16654; WO 96/34096; WO 96/33735; and WO 91/10741, each of which is incorporated herein by reference in its entirety. Human antibodies can also be produced using transgenic mice which are incapable of expressing functional endogenous immunoglobulins but which can express human immunoglobulin genes. See, *e.g.*, PCT publications WO 98/24893; WO 92/01047; WO 96/34096; WO 96/33735; U.S. Pat. Nos. 5,413,923; 5,625,126; 5,633,425; 5,569,825; 5,661,016; 5,545,806; 5,814,318; 5,885,793; 5,916,771; and 5,939,598, which are incorporated by reference herein in their entireties. Fully human antibodies that recognize a selected epitope can be generated using a technique referred to as "guided selection." In this approach, a selected non-human monoclonal antibody, *e.g.*, a mouse antibody, is used to guide the selection of a completely human antibody recognizing the same epitope (see, Jespers *et al.*, 1988, *Biotechnology* 12:899-903).

[0065] "Primatized antibodies" comprise monkey variable regions and human constant regions. Methods for producing primatized antibodies are known in the art. See, *e.g.*, U.S. Pat. Nos. 5,658,570; 5,681,722; and 5,693,780, which are incorporated herein by reference in their entireties.

[0066] Anti-glyco-cMET antibodies of the disclosure include both full-length (intact) antibody molecules, as well as antigen-binding fragments that are capable of binding glyco-cMET.

Examples of antigen-binding fragments include by way of example and not limitation, Fab, Fab', F(ab')₂, Fv fragments, single chain Fv fragments and single domain fragments.

[0067] A Fab fragment contains the constant domain of the light chain (CL) and the first constant domain (CH1) of the heavy chain. Fab' fragments differ from Fab fragments by the addition of a few residues at the carboxyl terminus of the heavy chain CH1 domain including one or more cysteines from the antibody hinge region. F(ab') fragments are produced by cleavage of the disulfide bond at the hinge cysteines of the F(ab')₂ pepsin digestion product. Additional chemical couplings of antibody fragments are known to those of ordinary skill in the art. Fab and F(ab')₁ fragments lack the Fc fragment of intact antibody, clear more rapidly from the circulation of animals, and may have less non-specific tissue binding than an intact antibody (see, e.g., Wahl *et al.*, 1983, J. Nucl. Med. 24:316).

[0068] An "Fv" fragment is the minimum fragment of an antibody that contains a complete target recognition and binding site. This region consists of a dimer of one heavy and one light chain variable domain in a tight, non-covalent association (V_H-V_L dimer). It is in this configuration that the three CDRs of each variable domain interact to define a target binding site on the surface of the V_H-V_L dimer. Often, the six CDRs confer target binding specificity to the antibody. However, in some instances even a single variable domain (or half of an Fv comprising only three CDRs specific for a target) can have the ability to recognize and bind target, although at a lower affinity than the entire binding site.

[0069] "Single-chain Fv" or "scFv" antigen-binding fragments comprise the V_H and V_L domains of an antibody, where these domains are present in a single polypeptide chain. Generally, the Fv polypeptide further comprises a polypeptide linker between the V_H and V_L domains which enables the scFv to form the desired structure for target binding.

[0070] "Single domain antibodies" are composed of single V_H or V_L domains which exhibit sufficient affinity to glyco-cMET. In a specific embodiment, the single domain antibody is a camelized antibody (see, e.g., Riechmann, 1999, Journal of Immunological Methods 231:25-38).

[0071] The anti-glyco-cMET antibodies of the disclosure may also be bispecific and other multiple specific antibodies. Bispecific antibodies are monoclonal, often human or humanized, antibodies that have binding specificities for two different epitopes on the same or different antigen. In the present disclosure, one of the binding specificities can be directed towards glyco-cMET, the other can be for any other antigen, e.g., for a cell-surface protein, receptor, receptor subunit, tissue-specific antigen, virally derived protein, virally encoded envelope protein, bacterially derived protein, or bacterial surface protein, *etc.* In certain embodiments, the bispecific and other multispecific anti-glyco-cMET antibodies and antigen binding fragments specifically bind to a second cMET epitope, an epitope on another protein co-expressed on cancer cells with cMET, or an epitope on another protein presented on a different cell, such as

an activated T cell. Bispecific antibodies of the disclosure include IgG format bispecific antibodies and single chain-based bispecific antibodies.

[0072] IgG format bispecific antibodies of the disclosure can be any of the various types of IgG format bispecific antibodies known in the art, such as quadroma bispecific antibodies, “knobs-in-holes” bispecific antibodies, CrossMab bispecific antibodies (*i.e.*, bispecific domain-exchanged antibodies), charge paired bispecific antibodies, common light chain bispecific antibodies, one-arm single-chain Fab-immunoglobulin gamma bispecific antibodies, disulfide stabilized Fv bispecific antibodies, DuetMabs, controlled Fab-arm exchange bispecific antibodies, strand-exchange engineered domain body bispecific antibodies, two-arm leucine zipper heterodimeric monoclonal bispecific antibodies, $\kappa\lambda$ -body bispecific antibodies, dual variable domain bispecific antibodies, and cross-over dual variable domain bispecific antibodies. See, *e.g.*, Köhler and Milstein, 1975, *Nature* 256:495-497; Milstein and Cuello, 1983, *Nature* 305:537-40; Ridgway *et al.*, 1996, *Protein Eng.* 9:617-621; Schaefer *et al.*, 2011, *Proc Natl Acad Sci USA* 108:11187-92; Gunasekaran *et al.*, 2010, *J Biol Chem* 285:19637-46; Fischer *et al.*, 2015 *Nature Commun* 6:6113; Schanzer *et al.*, 2014, *J Biol Chem* 289:18693-706; Metz *et al.*, 2012 *Protein Eng Des Sel* 25:571-80; Mazor *et al.*, 2015 *MAbs* 7:377-89; Labrijn *et al.*, 2013 *Proc Natl Acad Sci USA* 110:5145-50; Davis *et al.*, 2010 *Protein Eng Des Sel* 23:195-202; Wranik *et al.*, 2012, *J Biol Chem* 287:43331-9; Gu *et al.*, 2015, *PLoS One* 10(5):e0124135; Steinmetz *et al.*, 2016, *MAbs* 8(5):867-78; Klein *et al.*, 2016, *mAbs*, 8(6):1010-1020; Liu *et al.*, 2017, *Front. Immunol.* 8:38; and Yang *et al.*, 2017, *Int. J. Mol. Sci.* 18:48, which are incorporated herein by reference in their entireties.

[0073] In some embodiments, the bispecific antibodies of the disclosure are domain exchanged antibodies referred to in the scientific and patent literature as CrossMabs. See, *e.g.*, Schaefer *et al.*, 2011, *Proc Natl Acad Sci USA* 108:11187-92. The CrossMab technology is described in detail in WO 2009/080251, WO 2009/080252, WO 2009/080253, WO 2009/080254, WO 2013/026833, WO 2016/020309, and Schaefer *et al.*, 2011, *Proc Natl Acad Sci USA* 108:11187-92, which are incorporated herein by reference in their entireties. Briefly, the CrossMab technology is based on a domain crossover between heavy and light chains within one Fab-arm of a bispecific IgG, which promotes correct chain association. A CrossMab bispecific antibody of the disclosure can be a “CrossMab^{FAB}” antibody, in which the heavy and light chains of the Fab portion of one arm of a bispecific IgG antibody are exchanged. In other embodiments, a CrossMab bispecific antibody of the disclosure can be a “CrossMab^{VH-VL}” antibody, in which the only the variable domains of the heavy and light chains of the Fab portion of one arm of a bispecific IgG antibody are exchanged. In yet other embodiments, a CrossMab bispecific antibody of the disclosure can be a “CrossMab^{CH1-CL}” antibody, in which only the constant domains of the heavy and light chains of the Fab portion of one arm of a bispecific IgG antibody are exchanged. CrossMab^{CH1-CL} antibodies, in contrast to CrossMab^{FAB} and

CrossMab^{VH-VL}, do not have predicted side products and, therefore, in some embodiments CrossMab^{CH1-CL} bispecific antibodies are preferred. See, Klein *et al.*, 2016, *mAbs*, 8(6):1010-1020.

[0074] In some embodiments, the bispecific antibodies of the disclosure are controlled Fab-arm exchange bispecific antibodies. Methods for making Fab-arm exchange bispecific antibodies are described in PCT Publication No. WO2011/131746 and Labrijn *et al.*, 2014 *Nat Protoc.* 9(10):2450-63, incorporated herein by reference in their entireties. Briefly, controlled Fab-arm exchange bispecific antibodies can be made by separately expressing two parental IgG1s containing single matching point mutations in the CH3 domain, mixing the parental IgG1s under redox conditions *in vitro* to enable recombination of half-molecules, and removing the reductant to allow reoxidation of interchain disulfide bonds, thereby forming the bispecific antibodies.

[0075] In some embodiments, the bispecific antibodies of the disclosure are “bottle opener,” “mAb-Fv,” “mAb-scFv,” “central-scFv,” “central-Fv,” “one-armed central-scFv” or “dual scFv” format bispecific antibodies. Bispecific antibodies of these formats are described in PCT Publication No. WO 2016/182751, the contents of which are incorporated herein by reference in their entireties. Each of these formats relies on the self-assembling nature of Fc domains of antibody heavy chains, whereby two Fc subunit containing “monomers” assemble into a Fc domain containing “dimer.”

[0076] In the bottle opener format, the first monomer comprises a scFv covalently linked to the N-terminus of a Fc subunit, optionally via a linker, and the second monomer comprises a heavy chain (comprising a VH, CH1, and second Fc subunit). A bottle opener format bispecific antibody further comprises a light chain capable of pairing with the second monomer to form a Fab.

[0077] The mAb-Fv bispecific antibody format relies upon an “extra” VH domain attached to the C-terminus of one heavy chain monomer and an “extra” VL domain attached to the other heavy chain monomer, forming a third antigen binding domain. In some embodiments, a mAb-Fv bispecific antibody comprises a first monomer comprising a first VH domain, CH1 domain and a first Fc subunit, with a VL domain covalently attached to the C-terminus. The second monomer comprises a VH domain, a CH1 domain a second Fc subunit, and a VH covalently attached to the C-terminus of the second monomer. The two C-terminally attached variable domains make up a Fv. The mAb-Fv further comprises two light chains, which when associated with the first and second monomers form Fabs.

[0078] The mAb-scFv bispecific format relies on the use of a C-terminal attachment of a scFv to one of the monomers of a mAb, thus forming a third antigen binding domain. Thus, the first monomer comprises a first heavy chain (comprising a VH, CH1 and a first Fc subunit), with a C-terminally covalently attached scFv. mAb-scFv bispecific antibodies further comprise a second

monomer (comprising a VH, CH1, and first Fc subunit) and two light chains, which when associated with the first and second monomers form Fabs.

[0079] The central-scFv bispecific format relies on the use of an inserted scFv domain in a mAb, thus forming a third antigen binding domain. The scFv domain is inserted between the Fc subunit and the CH1 domain of one of the monomers, thus providing a third antigen binding domain. Thus, the first monomer can comprise a VH domain, a CH1 domain (and optional hinge) and a first Fc subunit, with a scFv covalently attached between the C-terminus of the CH1 domain and the N-terminus of the first Fc subunit using optional domain linkers. The other monomer can be a standard Fab side monomer. Central-scFv bispecific antibodies further comprise two light chains, which when associated with the first and second monomers form Fabs.

[0080] The central-Fv bispecific format relies on the use of an inserted Fv domain thus forming a third antigen binding domain. Each monomer can contain a component of the Fv (e.g. one monomer comprises a variable heavy domain and the other a variable light domain). Thus, one monomer can comprise a VH domain, a CH1 domain, a first Fc subunit and a VL domain covalently attached between the C-terminus of the CH1 domain and the N-terminus of the first Fc subunit, optionally using domain linkers. The other monomer can comprise a VH domain, a CH1 domain, a second Fc subunit and an additional VH domain covalently attached between the C-terminus of the CH1 domain and the N-terminus of the second Fc domain, optionally using domain linkers. Central-Fv bispecific antibodies further comprise two light chains, which when associated with the first and second monomers form Fabs.

[0081] The one-armed central-scFv bispecific format comprises one monomer comprising just an Fc subunit, while the other monomer comprises an inserted scFv domain thus forming a second antigen binding domain. Thus, one monomer can comprise a VH domain, a CH1 domain and a first Fc subunit, with a scFv covalently attached between the C-terminus of the CH1 domain and the N-terminus of the first Fc subunit, optionally using domain linkers. The second monomer can comprise an Fc domain. This embodiment further utilizes a light chain comprising a variable light domain and a constant light domain, that associates with the first monomer to form a Fab.

[0082] The dual scFv bispecific format comprises a first monomer comprising a scFv covalently attached to the N-terminus of a first Fc subunit, optionally via a linker, and second monomer comprising a scFv covalently attached to the N-terminus of a second Fc subunit, optionally via a linker.

[0083] Bispecific antibodies of the disclosure can comprise an Fc domain composed of a first and a second subunit. In one embodiment, the Fc domain is an IgG Fc domain. In a particular embodiment, the Fc domain is an IgG₁ Fc domain. In another embodiment the Fc domain is an IgG₄ Fc domain. In a more specific embodiment, the Fc domain is an IgG₄ Fc domain

comprising an amino acid substitution at position S228 (Kabat EU index numbering), particularly the amino acid substitution S228P. Unless otherwise specified herein, numbering of amino acid residues in an Fc domain or constant region is according to the EU numbering system, also called the EU index, as described in Kabat *et al.*, 1991, Sequences of Proteins of Immunological Interest, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, MD. This amino acid substitution reduces *in vivo* Fab arm exchange of IgG₄ antibodies (see Stubenrauch *et al.*, 2010, Drug Metabolism and Disposition 38:84-91). In a further particular embodiment, the Fc domain is a human Fc domain. In an even more particular embodiment, the Fc domain is a human IgG₁ Fc domain. An exemplary sequence of a human IgG₁ Fc region is:

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DKTHTCPPCPAPELLGGPSVFLFPPKPKDTLMISRTPEVTCVWVDVSHEDPEVKFNWYVDGVE  
VHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREP  
Q (SEQ ID NO:287).
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[0084] In particular embodiments, the Fc domain comprises a modification promoting the association of the first and the second subunit of the Fc domain. The site of most extensive protein-protein interaction between the two subunits of a human IgG Fc domain is in the CH3 domain. Thus, in one embodiment said modification is in the CH3 domain of the Fc domain.

[0085] In a specific embodiment said modification promoting the association of the first and the second subunit of the Fc domain is a so-called “knob-into-hole” modification, comprising a “knob” modification in one of the two subunits of the Fc domain and a “hole” modification in the other one of the two subunits of the Fc domain. The knob-into-hole technology is described *e.g.* in US 5,731,168; US 7,695,936; Ridgway *et al.*, 1996, Prot Eng 9:617-621, and Carter, J, 2001, Immunol Meth 248:7-15. Generally, the method involves introducing a protuberance (“knob”) at the interface of a first polypeptide and a corresponding cavity (“hole”) in the interface of a second polypeptide, such that the protuberance can be positioned in the cavity so as to promote heterodimer formation and hinder homodimer formation. Protuberances are constructed by replacing small amino acid side chains from the interface of the first polypeptide with larger side chains (*e.g.* tyrosine or tryptophan). Compensatory cavities of identical or similar size to the protuberances are created in the interface of the second polypeptide by replacing large amino acid side chains with smaller ones (*e.g.* alanine or threonine).

[0086] Accordingly, in some embodiments, an amino acid residue in the CH3 domain of the first subunit of the Fc domain is replaced with an amino acid residue having a larger side chain volume, thereby generating a protuberance within the CH3 domain of the first subunit which is positionable in a cavity within the CH3 domain of the second subunit, and an amino acid residue in the CH3 domain of the second subunit of the Fc domain is replaced with an amino acid residue having a smaller side chain volume, thereby generating a cavity within the CH3 domain of the second subunit within which the protuberance within the CH3 domain of the first

subunit is positionable. Preferably said amino acid residue having a larger side chain volume is selected from the group consisting of arginine (R), phenylalanine (F), tyrosine (Y), and tryptophan (W). Preferably said amino acid residue having a smaller side chain volume is selected from the group consisting of alanine (A), serine (S), threonine (T), and valine (V). The protuberance and cavity can be made by altering the nucleic acid encoding the polypeptides, e.g. by site-specific mutagenesis, or by peptide synthesis.

[0087] In a specific such embodiment, in the first subunit of the Fc domain the threonine residue at position 366 is replaced with a tryptophan residue (T366W), and in the second subunit of the Fc domain the tyrosine residue at position 407 is replaced with a valine residue (Y407V) and optionally the threonine residue at position 366 is replaced with a serine residue (T366S) and the leucine residue at position 368 is replaced with an alanine residue (L368A) (numbering according to Kabat EU index). In a further embodiment, in the first subunit of the Fc domain additionally the serine residue at position 354 is replaced with a cysteine residue (S354C) or the glutamic acid residue at position 356 is replaced with a cysteine residue (E356C) (particularly the serine residue at position 354 is replaced with a cysteine residue), and in the second subunit of the Fc domain additionally the tyrosine residue at position 349 is replaced by a cysteine residue (Y349C) (numbering according to Kabat EU index). In a particular embodiment, the first subunit of the Fc domain comprises the amino acid substitutions S354C and T366W, and the second subunit of the Fc domain comprises the amino acid substitutions Y349C, T366S, L368A and Y407V (numbering according to Kabat EU index).

[0088] In some embodiments, electrostatic steering (e.g., as described in Gunasekaran *et al.*, 2010, J Biol Chem 285(25):19637-46) can be used to promote the association of the first and the second subunit of the Fc domain.

[0089] In some embodiments, the Fc domain comprises one or more amino acid substitutions that reduces binding to an Fc receptor and/or effector function.

[0090] In a particular embodiment the Fc receptor is an Fcγ receptor. In one embodiment the Fc receptor is a human Fc receptor. In one embodiment the Fc receptor is an activating Fc receptor. In a specific embodiment the Fc receptor is an activating human Fcγ receptor, more specifically human FcγRIIIa, FcγRI or FcγRIIa, most specifically human FcγRIIIa. In one embodiment the effector function is one or more selected from the group of complement dependent cytotoxicity (CDC), antibody-dependent cell-mediated cytotoxicity (ADCC), antibody-dependent cellular phagocytosis (ADCP), and cytokine secretion. In a particular embodiment, the effector function is ADCC.

[0091] Typically, the same one or more amino acid substitution is present in each of the two subunits of the Fc domain. In one embodiment, the one or more amino acid substitution reduces the binding affinity of the Fc domain to an Fc receptor. In one embodiment, the one or

more amino acid substitution reduces the binding affinity of the Fc domain to an Fc receptor by at least 2-fold, at least 5-fold, or at least 10-fold.

[0092] In one embodiment, the Fc domain comprises an amino acid substitution at a position selected from the group of E233, L234, L235, N297, P331 and P329 (numberings according to Kabat EU index). In a more specific embodiment, the Fc domain comprises an amino acid substitution at a position selected from the group of L234, L235 and P329 (numberings according to Kabat EU index). In some embodiments, the Fc domain comprises the amino acid substitutions L234A and L235A (numberings according to Kabat EU index). In one such embodiment, the Fc domain is an IgG₁ Fc domain, particularly a human IgG₁ Fc domain. In one embodiment, the Fc domain comprises an amino acid substitution at position P329. In a more specific embodiment, the amino acid substitution is P329A or P329G, particularly P329G (numberings according to Kabat EU index). In one embodiment, the Fc domain comprises an amino acid substitution at position P329 and a further amino acid substitution at a position selected from E233, L234, L235, N297 and P331 (numberings according to Kabat EU index). In a more specific embodiment, the further amino acid substitution is E233P, L234A, L235A, L235E, N297A, N297D or P331S. In particular embodiments, the Fc domain comprises amino acid substitutions at positions P329, L234 and L235 (numberings according to Kabat EU index). In more particular embodiments, the Fc domain comprises the amino acid mutations L234A, L235A and P329G (which can be referred to using the shorthand terms “P329G LALA”, “PGLALA” or “LALAPG”). Specifically, in particular embodiments, each subunit of the Fc domain comprises the amino acid substitutions L234A, L235A and P329G (Kabat EU index numbering), *i.e.* in each of the first and the second subunit of the Fc domain the leucine residue at position 234 is replaced with an alanine residue (L234A), the leucine residue at position 235 is replaced with an alanine residue (L235A) and the proline residue at position 329 is replaced by a glycine residue (P329G) (numbering according to Kabat EU index). In one such embodiment, the Fc domain is an IgG₁ Fc domain, particularly a human IgG₁ Fc domain.

[0093] Single chain-based bispecific antibodies of the disclosure can be any of the various types of single chain-based bispecific antibodies known in the art, such as bispecific T-cell engagers (BiTEs), diabodies, tandem diabodies (tandabs), dual-affinity retargeting molecules (DARTs), and bispecific killer cell engagers. See, *e.g.*, Löffler *et al.*, 2000, *Blood* 95:2098–103; Holliger *et al.*, 1993, *Proc Natl Acad Sci USA*, 90:6444–8; Kipriyanov *et al.*, 1999, *Mol Biol* 293:41–56; Johnson *et al.*, 2010, *Mol Biol* 399:436–49; Wiernik *et al.*, 2013, *Clin Cancer Res* 19:3844–55; Liu *et al.*, 2017, *Front. Immunol.* 8:38; and Yang *et al.*, 2017, *Int. J. Mol. Sci.* 18:48, which are incorporated herein by reference in their entireties.

[0094] In some embodiments, the bispecific antibodies of the disclosure are bispecific T-cell engagers (BiTEs). BiTEs are single polypeptide chain molecules having two antigen-binding domains, one of which binds to a T-cell antigen and the second of which binds to an antigen

present on the surface of a target (see, PCT Publication WO 05/061547; Baeuerle *et al.*, 2008, *Drugs of the Future* 33: 137-147; Bargou, *et al.*, 2008, *Science* 321:974-977, incorporated herein by reference in their entireties). Thus, the BiTEs of the disclosure have an antigen binding domain that binds to a T-cell antigen, and a second antigen binding domain that is directed towards glyco-cMET.

[0095] In some embodiments, the bispecific antibodies of the disclosure are dual-affinity retargeting molecules (DARTs). DARTs comprise at least two polypeptide chains that associate (especially through a covalent interaction) to form at least two epitope binding sites, which may recognize the same or different epitopes. Each of the polypeptide chains of a DART comprise an immunoglobulin light chain variable region and an immunoglobulin heavy chain variable region, but these regions do not interact to form an epitope binding site. Rather, the immunoglobulin heavy chain variable region of one (*e.g.*, the first) of the DART polypeptide chains interacts with the immunoglobulin light chain variable region of a different (*e.g.*, the second) DART™ polypeptide chain to form an epitope binding site. Similarly, the immunoglobulin light chain variable region of one (*e.g.*, the first) of the DART polypeptide chains interacts with the immunoglobulin heavy chain variable region of a different (*e.g.*, the second) DART polypeptide chain to form an epitope binding site. DARTs may be monospecific, bispecific, trispecific, etc., thus being able to simultaneously bind one, two, three or more different epitopes (which may be of the same or of different antigens). DARTs may additionally be monovalent, bivalent, trivalent, tetravalent, pentavalent, hexavalent, etc., thus being able to simultaneously bind one, two, three, four, five, six or more molecules. These two attributes of DARTs (*i.e.*, degree of specificity and valency may be combined, for example to produce bispecific antibodies (*i.e.*, capable of binding two epitopes) that are tetravalent (*i.e.*, capable of binding four sets of epitopes), *etc.* DART molecules are disclosed in PCT Publications WO 2006/113665, WO 2008/157379, and WO 2010/080538, which are incorporated herein by reference in their entireties.

[0096] In some embodiments of the bispecific antibodies of the disclosure, one of the binding specificities is directed towards glyco-cMET, and the other is directed to an antigen expressed on immune effector cells. The term “immune effector cell” or “effector cell” as used herein refers to a cell within the natural repertoire of cells in the mammalian immune system which can be activated to affect the viability of a target cell. Immune effector cells include cells of the lymphoid lineage such as natural killer (NK) cells, T cells including cytotoxic T cells, or B cells, but also cells of the myeloid lineage can be regarded as immune effector cells, such as monocytes or macrophages, dendritic cells and neutrophilic granulocytes. Hence, said effector cell is preferably an NK cell, a T cell, a B cell, a monocyte, a macrophage, a dendritic cell or a neutrophilic granulocyte. Recruitment of effector cells to aberrant cells means that immune effector cells are brought in close vicinity to the aberrant target cells such that the effector cells

can directly kill, or indirectly initiate the killing of the aberrant cells that they are recruited to. In order to avoid non specific interactions it is preferred that the bispecific antibodies of the disclosure specifically recognize antigens on immune effector cells that are at least over-expressed by these immune effector cells compared to other cells in the body. Target antigens present on immune effector cells may include CD3, CD8, CD16, CD25, CD28, CD64, CD89, NKG2D and NKp46. Preferably, the antigen on immune effector cells is CD3 expressed on T cells.

[0097] As used herein, “CD3” refers to any native CD3 from any vertebrate source, including mammals such as primates (*e.g.* humans), non-human primates (*e.g.* cynomolgus monkeys) and rodents (*e.g.* mice and rats), unless otherwise indicated. The term encompasses “full-length,” unprocessed CD3 as well as any form of CD3 that results from processing in the cell. The term also encompasses naturally occurring variants of CD3, *e.g.*, splice variants or allelic variants. The most preferred antigen on an immune effector cell is the CD3 epsilon chain. This antigen has been shown to be very effective in recruiting T cells to aberrant cells. Hence, a bispecific antibody of the disclosure preferably specifically recognizes CD3 epsilon. The amino acid sequence of human CD3 epsilon is shown in UniProt (uniprot.org) accession no. P07766 (version 144), or NCBI (ncbi.nlm.nih.gov/) RefSeq NP_000724.1. The amino acid sequence of cynomolgus [*Macaca fascicularis*] CD3 epsilon is shown in NCBI GenBank no. BAB71849.1. For human therapeutic use, bispecific antibodies in which the CD3-binding domain specifically binds to human CD3 (*e.g.*, the human CD3 epsilon chain) are used. For preclinical testing in non-human animals and cell lines, bispecific antibodies in which the CD3-binding domain specifically binds to the CD3 in the species utilized for the preclinical testing (*e.g.*, cynomolgus CD3 for primate testing) can be used.

[0098] As used herein, a binding domain that “specifically binds to” or “specifically recognizes” a target antigen from a particular species does not preclude the binding to or recognition of the antigen from other species, and thus encompasses antibodies in which one or more of the binding domains have inter-species cross-reactivity. For example, a CD3-binding domain that “specifically binds to” or “specifically recognizes” human CD3 may also bind to or recognize cynomolgus CD3, and vice versa.

[0099] In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody H2C (described in PCT publication no. WO2008/119567) for binding an epitope of CD3. In other embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody V9 (described in Rodrigues *et al.*, 1992, Int J Cancer Suppl 7:45-50 and U.S. Pat. No. 6,054,297) for binding an epitope of CD3. In yet other embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody FN18 (described in Nooij *et al.*, 1986, Eur J Immunol 19:981-984) for binding an epitope of CD3. In yet other embodiments,

a bispecific antibody of the disclosure can compete with monoclonal antibody SP34 (described in Pessano *et al.*, 1985, EMBO J 4:337-340) for binding an epitope of CD3.

[0100] In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody mAb1 (described in U.S. Pat. No. 10,730,944) for binding an epitope of CD8. In other embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody YTS169 (described in US2015/ 0191543) for binding an epitope of CD8. In other embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibodies 4C9 5F4 (described in WO1987/005912) for binding an epitope of CD8.

[0101] In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody 3G8_ (described in WO2006/064136) for binding an epitope of CD16. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody VEP13 (described in Ziegler-Heitbrock *et al.*, 1984, Clin.Exp. Immunol. 58:470-477) for binding an epitope of CD16. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody B73.1 (described in Perussia *et al.*, 1983, J. Immunol.130(5):2142-2148) for binding an epitope of CD16.

[0102] In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody daclizumab and its variants (described in WO2014/145000) for binding an epitope of CD25. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibodies AB1, AB7, AB11, or AB12 (described in WO2004/045512) for binding an epitope of CD25. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibodies ALD25H1, ALD25H2, or ALD25H4 (described in WO2020/234399) for binding an epitope of CD25.

[0103] In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody FR104 (described in WO2017/103003) for binding an epitope of CD28. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody hCD28.3 (described in WO2011/101791) for binding an epitope of CD28.

[0104] In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibodies MS or 21 F2 (described in WO2009/077483) for binding an epitope of NKG2D. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibodies 5C5, 320, 230, 013, 296 or 395 (described in WO2021/009146) for binding an epitope of NKG2D. In some embodiments, a bispecific antibody of the disclosure can compete with monoclonal antibody KYK-2.0 (described in WO2010/017103) for binding an epitope of NKG2D.

[0105] The anti-glyco-cMET antibodies of the disclosure include derivatized antibodies. For example, but not by way of limitation, derivatized antibodies are typically modified by glycosylation, acetylation, pegylation, phosphorylation, amidation, derivatization by known

protecting/blocking groups, proteolytic cleavage, linkage to a cellular ligand or other protein. Any of numerous chemical modifications can be carried out by known techniques, including, but not limited to, specific chemical cleavage, acetylation, formylation, metabolic synthesis of tunicamycin, *etc.* Additionally, the derivative can contain one or more non-natural amino acids, *e.g.*, using ambrx technology (see, *e.g.*, Wolfson, 2006, *Chem. Biol.* 13(10):1011-2).

[0106] The anti-glyco-cMET antibodies or binding fragments may be antibodies or fragments whose sequences have been modified to alter at least one constant region-mediated biological effector function. For example, in some embodiments, an anti-glyco-cMET antibody may be modified to reduce at least one constant region-mediated biological effector function relative to the unmodified antibody, *e.g.*, reduced binding to the Fc receptor (FcγR). FcγR binding can be reduced by mutating the immunoglobulin constant region segment of the antibody at particular regions necessary for FcγR interactions (see, *e.g.*, Canfield and Morrison, 1991, *J. Exp. Med.* 173:1483-1491; and Lund *et al.*, 1991, *J. Immunol.* 147:2657-2662). Reduction in FcγR binding ability of the antibody can also reduce other effector functions which rely on FcγR interactions, such as opsonization, phagocytosis and antigen-dependent cellular cytotoxicity ("ADCC").

[0107] The anti-glyco-cMET antibody or binding fragments described herein include antibodies and/or binding fragments that have been modified to acquire or improve at least one constant region-mediated biological effector function relative to an unmodified antibody, *e.g.*, to enhance FcγR interactions (see, *e.g.*, US 2006/0134709). For example, an anti-glyco-cMET antibody of the disclosure can have a constant region that binds FcγRIIA, FcγRIIB and/or FcγRIIIA with greater affinity than the corresponding wild type constant region.

[0108] Thus, antibodies of the disclosure may have alterations in biological activity that result in increased or decreased opsonization, phagocytosis, or ADCC. Such alterations are known in the art. For example, modifications in antibodies that reduce ADCC activity are described in U.S. Pat. No. 5,834,597. An exemplary ADCC lowering variant corresponds to "mutant 3" (shown in FIG. 4 of U.S. Pat. No. 5,834,597) in which residue 236 is deleted and residues 234, 235 and 237 (using EU numbering) are substituted with alanines. Another exemplary ADCC lowering variant comprises amino acid mutations L234A, L235A and P329G (which can be referred to using the shorthand term "P329G LALA"). The "P329G LALA" combination of amino acid substitutions almost completely abolishes Fcγ receptor (as well as complement) binding of a human IgG₁ Fc domain, as described in PCT publication no. WO 2012/130831, incorporated herein by reference in its entirety. WO 2012/130831 also describes methods of preparing such mutant Fc domains and methods for determining its properties such as Fc receptor binding or effector functions.

[0109] In some embodiments, the anti-glyco-cMET antibodies of the disclosure have low levels of, or lack, fucose. Antibodies lacking fucose have been correlated with enhanced ADCC activity, especially at low doses of antibody. See Shields *et al.*, 2002, *J. Biol. Chem.* 277:26733-

26740; Shinkawa *et al.*, 2003, J. Biol. Chem. 278:3466-73. Methods of preparing fucose-less antibodies include growth in rat myeloma YB2/0 cells (ATCC CRL 1662). YB2/0 cells express low levels of FUT8 mRNA, which encodes α -1, 6-fucosyltransferase, an enzyme necessary for fucosylation of polypeptides.

[0110] In some embodiments, the anti-glyco-cMET antibodies or binding fragments include bisected oligosaccharides, *e.g.*, in which a biantennary oligosaccharide attached to an Fc domain is bisected by GlcNAc. Such variants may have reduced fucosylation and/or improved ADCC function as described above. Examples of such antibody variants are described, *e.g.*, in Umana *et al.*, 1999, Nat Biotechnol 17:176-180; Ferrara *et al.*, 2006, Biotechnol Bioeng 93: 851-861; WO 99/54342; WO 2004/065540; and WO 2003/011878.

[0111] In yet another aspect, the anti-glyco-cMET antibodies or binding fragments include modifications that increase or decrease their binding affinities to the fetal Fc receptor, FcRn, for example, by mutating the immunoglobulin constant region segment at particular regions involved in FcRn interactions (see, *e.g.*, WO 2005/123780). In particular embodiments, an anti-glyco-cMET antibody of the IgG class is mutated such that at least one of amino acid residues 250, 314, and 428 of the heavy chain constant region is substituted alone, or in any combinations thereof, such as at positions 250 and 428, or at positions 250 and 314, or at positions 314 and 428, or at positions 250, 314, and 428, with positions 250 and 428 a specific combination. For position 250, the substituting amino acid residue can be any amino acid residue other than threonine, including, but not limited to, alanine, cysteine, aspartic acid, glutamic acid, phenylalanine, glycine, histidine, isoleucine, lysine, leucine, methionine, asparagine, proline, glutamine, arginine, serine, valine, tryptophan, or tyrosine. For position 314, the substituting amino acid residue can be any amino acid residue other than leucine, including, but not limited to, alanine, cysteine, aspartic acid, glutamic acid, phenylalanine, glycine, histidine, isoleucine, lysine, methionine, asparagine, proline, glutamine, arginine, serine, threonine, valine, tryptophan, or tyrosine. For position 428, the substituting amino acid residues can be any amino acid residue other than methionine, including, but not limited to, alanine, cysteine, aspartic acid, glutamic acid, phenylalanine, glycine, histidine, isoleucine, lysine, leucine, asparagine, proline, glutamine, arginine, serine, threonine, valine, tryptophan, or tyrosine. Specific combinations of suitable amino acid substitutions are identified in Table 1 of U.S. Pat. No. 7,217,797, which is incorporated herein by reference. Such mutations increase binding to FcRn, which protects the antibody from degradation and increases its half-life.

[0112] In yet other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure has one or more amino acids inserted into one or more of its hypervariable regions, for example as described in Jung and Pluckthun, 1997, Protein Engineering 10:9, 959-966; Yazaki *et al.*, 2004, Protein Eng. Des Sel. 17(5):481-9. Epub 2004 Aug. 17; and U.S. Pat. App. No. 2007/0280931.

[0113] In yet other aspects, particularly useful for diagnostic applications, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure is attached to a detectable moiety. Detectable moieties include a radioactive moiety, a colorimetric molecule, a fluorescent moiety, a chemiluminescent moiety, an antigen, an enzyme, a detectable bead (such as a magnetic or electrodense (e.g., gold) bead), or a molecule that binds to another molecule (e.g., biotin or streptavidin)).

[0114] Radioisotopes or radionuclides may include ^3H , ^{14}C , ^{15}N , ^{35}S , ^{90}Y , ^{99}Tc , ^{111}In , ^{125}I , ^{131}I .

[0115] Fluorescent labels may include rhodamine, lanthanide phosphors, fluorescein and its derivatives, fluorochrome, GFP (GFP for "Green Fluorescent Protein"), dansyl, umbelliferone, phycoerythrin, phycocyanin, allophycocyanin, o-phthaldehyde, and fluorescamine.

[0116] Enzymatic labels may include horseradish peroxidase, β galactosidase, luciferase, alkaline phosphatase, glucose-6-phosphate dehydrogenase ("G6PDH"), alpha-D-galactosidase, glucose oxidase, glucose amylase, carbonic anhydrase, acetylcholinesterase, lysozyme, malate dehydrogenase and peroxidase.

[0117] Chemiluminescent labels or chemiluminescers, such as isoluminol, luminol and the dioxetanes.

[0118] Other detectable moieties include molecules such as biotin, digoxigenin or 5-bromodeoxyuridine.

[0119] In yet other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure may be used in a detection system to detect a biomarker in a sample, such as, e.g., a patient-derived biological sample. The biomarker may be a protein biomarker (e.g., a tumor-associated glycoform of cMET, for example a glycoform of cMET comprising the amino acid sequence PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285), glycosylated with GalNAc on the serine and threonine residues shown in bold underlined text) present on the surface of or within, e.g., a cancer cell (e.g., from a tissue biopsy or a circulating tumor cell) or a cancer-derived extracellular vesicle).

[0120] Extracellular vesicles (EVs) are lipid membranous vesicles released from almost all cell types. EVs carry complex molecular cargoes, such as proteins, RNAs (e.g., mRNA and noncoding RNAs (microRNA, transfer RNA, circular RNA and long noncoding RNA)), and DNA fragments. The molecular contents of EVs largely reflect the cell of origin and thus show cell-type specificity. In particular, cancer-derived EVs contain and present on their surfaces cancer-specific molecules expressed by parental cancer cells (see, e.g., Yáñez-Mó *et al.*, 2015, *J Extracell Vesicles*. 4:27066; and Li *et al.*, 2015, *Cell Res*. 25:981-984)

[0121] In one embodiment, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure is used in a method of detecting a biomarker in a sample comprising EVs (e.g., a liquid biopsy). In such embodiments, the biomarker is recognized by the anti-glyco-cMET

antibody or antigen-binding fragment of the disclosure. The biomarker may be present on the surface of EVs. Exemplary methods of detecting the biomarker include, but are not limited to, immunoassays, such as immunoprecipitation; Western blot; ELISA; immunohistochemistry; immunocytochemistry; flow cytometry; and immuno-PCR. In some embodiments, an immunoassay can be a chemiluminescent immunoassay. In some embodiments, an immunoassay can be a high-throughput and/or automated immunoassay platform.¶

[0122] In some embodiments, the method of detecting a biomarker in a sample comprises contacting a sample with an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure. In some embodiments, such methods further comprise contacting the sample with one or more detection labels. In some embodiments, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure is labeled with one or more detection labels.

[0123] In some embodiments, a capture assay is performed to selectively capture EVs from a sample such as a liquid biopsy sample exemplary examples of capture assays for EVs are described in US2021/0214806, which is hereby incorporated by reference in its entirety. In some embodiments, a capture assay is performed to selectively capture EVs of a certain size range, and/or certain characteristic(s), for example, EVs associated with cancer (e.g., a tumor-associated glycoform of cMET, for example a glycoform of cMET comprising the amino acid sequence PTKSFISGGSTITGVGKLN (SEQ ID NO:285), glycosylated with GalNAc on the serine and threonine residues shown in bold underlined text), glycosylated with GalNAc on the threonine residue shown in bold underlined text). In some such embodiments, prior to performing the capture assay, a sample may be pre-processed to remove non-EVs, including but not limited to, e.g., soluble proteins and interfering entities such as, e.g., cell debris. In some embodiments, EVs are purified from a sample using size exclusion chromatography.

[0124] In some embodiments, the method for detecting a biomarker comprises analyzing individual EVs (e.g., a single EV assay). For example, such an assay may involve (i) a capture assay such as an antibody capture assay and (ii) one or more detection assays for at least one or more additional biomarkers, wherein the capture assay is performed prior to the detection assay. See, e.g., US2021/0214806.¶

[0125] In some embodiments, a capture assay comprises a step of contacting a sample with at least one capture agent comprising an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure. The capture agent may be immobilized on a solid substrate. The solid substrate may be provided in a form that is suitable for capturing EVs and does not interfere with downstream handling, processing, and/or detection. For example, in some embodiments, a solid substrate may be or comprise a bead (e.g., a magnetic bead). In some embodiments, a solid substrate may be or comprise a surface. For example, in some embodiments, such a surface may be a capture surface of an assay chamber (e.g., a tube, a well, a microwell, a plate, a filter, a membrane, a matrix, etc.). In some embodiments, a capture agent is or

comprises a magnetic bead comprising a capture moiety (e.g., an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure) conjugated thereto. See, e.g., US2021/0214806.

[0126] In certain aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with 15C4, or an antibody or antigen binding fragment comprising heavy and light chain variable regions of 15C4 (SEQ ID NOS:1 and 2, respectively).

[0127] In certain aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with 8H3 or an antibody or antigen binding fragment comprising heavy chain variable region of murine or humanized 8H3 (e.g., SEQ ID NO:23 (murine) and SEQ ID NOS: 264-275 (exemplary humanized sequences)) and a light chain variable region of murine or humanized 8H3 (e.g., SEQ ID NO:24 (murine) and SEQ ID NO:276-284 (exemplary humanized sequences)).

[0128] In certain aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with 16E12, or an antibody or antigen binding fragment comprising heavy and light chain variable regions of 16E12 (SEQ ID NOS:45 and 46, respectively).

[0129] In certain aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with 14E9, or an antibody or antigen binding fragment comprising heavy and light chain variable regions of 14E9 (SEQ ID NOS:67 and 68, respectively).

[0130] In certain aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with 19H2, or an antibody or antigen binding fragment comprising heavy and light chain variable regions of 19H2 (SEQ ID NOS:89 and 90, respectively).

[0131] In certain aspects, an anti-glyco-cMET antibody or antigen binding fragment of the disclosure competes with 39A3, or an antibody or antigen binding fragment comprising heavy and light chain variable regions of 39A3 (SEQ ID NOS:111 and 112, respectively).

[0132] Competition can be assayed on cells that express the glyco-cMET epitope bound by 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3, or on a glycosylated cMET peptide containing the epitope bound by 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3, e.g., the peptide PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285), glycosylated with GalNAc on the serine and threonine residues shown in bold and underlined text. Cells that do not express the epitope or unglycosylated peptides can be used as controls.

[0133] Cells on which a competition assay can be carried out include but are not limited to lung, breast, renal, and liver cell lines (e.g., breast cancer cell line T47D; adenocarcinomic human alveolar basal epithelial cell line A549) and recombinant cells that are engineered to express the glyco-cMET epitope. In one non-limiting example, T47D cells, which express cMET but are inherently Tn-negative, are engineered to express the cMET Tn-antigen by knockout of the COSMC chaperone. Wildtype T47D cells expressing the unglycosylated form of cMET can be used as a negative control. In another non-limiting example, A549 cells, which express cMET

but are inherently Tn-negative, are engineered to express the cMET Tn-antigen by knockout of the COSMC chaperone. Wildtype A549 cells expressing the unglycosylated form of cMET can be used as a negative control

[0134] Assays for competition include, but are not limited to, a radioactive material labeled immunoassay (RIA), an enzyme-linked immunosorbent assay (ELISA), a sandwich ELISA, fluorescence activated cell sorting (FACS) assays, surface plasmon resonance (e.g., Biacore) assays, and bio-layer interferometry (BLI) assays. In some embodiments, antibody competition assays can be carried out using BLI (e.g., using an Octet-HTX system (Molecular Devices)). Antibody competition or epitope binning of monoclonal antibodies can be assessed in tandem against their specific antigen using BLI. In a BLI assay, the antigen can be immobilized onto a biosensor and presented to two competing antibodies in consecutive steps. The binding to non-overlapping epitopes occurs if saturation with the first antibody does not block the binding of the second antibody. In some embodiments, antibody competition assays can be carried out using surface plasmon resonance (e.g., using a Biacore system (Cytiva)). In a surface plasmon resonance assay, one or more antibodies can be immobilized onto a biosensor and presented with an analyte (e.g., the glyco-cMET peptide of SEQ ID NO:285 or a negative control analyte such as an unglycosylated cMET peptide of SEQ ID NO:286). In some embodiments, the antibodies are contacted with a saturating concentration of the analyte, for example a concentration of at least about 0.5 μM . In some embodiments the saturating concentration is about 1 μM , about 1.5 μM , or about 2 μM . When comparing the binding affinities of two antibodies, the affinities of both antibodies are preferably measured using the same concentration of both antibodies, e.g., measured using a 1 μM concentration of each antibody.

[0135] In conducting an antibody competition assay between a reference antibody and a test antibody (irrespective of species or isotype), one may first label the reference with a detectable label, such as a fluorophore, biotin or an enzymatic (or even radioactive) label to enable subsequent identification. In this case, cells expressing glyco-cMET are incubated with unlabeled test antibody, labeled reference antibody is added, and the intensity of the bound label is measured. If the test antibody competes with the labeled reference antibody by binding to an overlapping epitope, the intensity will be decreased relative to a control reaction carried out without test antibody.

[0136] In a specific embodiment of this assay, the concentration of labeled reference antibody that yields 80% of maximal binding (“ $\text{conc}_{80\%}$ ”) under the assay conditions (e.g., a specified density of cells) is first determined, and a competition assay carried out with 10 x $\text{conc}_{80\%}$ of unlabeled test antibody and $\text{conc}_{80\%}$ of labeled reference antibody.

[0137] The inhibition can be expressed as an inhibition constant, or K_i , which is calculated according to the following formula:

$$K_i = \text{IC}_{50} / (1 + [\text{reference Ab concentration}] / K_d),$$

where IC_{50} is the concentration of test antibody that yields a 50% reduction in binding of the reference antibody and K_d is the dissociation constant of the reference antibody, a measure of its affinity for glyco-cMET. Antibodies that compete with anti-glyco-cMET antibodies disclosed herein can have a K_i from 10 pM to 10 nM under assay conditions described herein.

[0138] In various embodiments, a test antibody is considered to compete with a reference antibody if it decreases binding of the reference antibody by at least about 20% or more, for example, by at least about 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95% or even more, or by a percentage ranging between any of the foregoing values, at a reference antibody concentration that is 80% of maximal binding under the specific assay conditions used, and a test antibody concentration that is 10-fold higher than the reference antibody concentration.

[0139] In one example of a competition assay, the glycosylated cMET peptide of SEQ ID NO:285 is adhered onto a solid surface, e.g., a microwell plate, by contacting the plate with a solution of the peptide (e.g., at a concentration of 1 μ g/mL in PBS over night at 4 °C). The plate is washed (e.g., 0.1% Tween 20 in PBS) and blocked (e.g., in Superblock, Thermo Scientific, Rockford, IL). A mixture of sub-saturating amount of biotinylated 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 (e.g., at a concentration of 80 ng/mL) and unlabeled 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 (the "reference" antibody) or competing anti-glyco-cMET antibody (the "test" antibody) in serial dilution (e.g., at a concentration of 2.8 μ g/mL, 8.3 μ g/mL, or 25 μ g/mL) in ELISA buffer (e.g., 1% BSA and 0.1% Tween 20 in PBS) is added to wells and plates are incubated for 1 hour with gentle shaking. The plate is washed, 1 μ g/mL HRP-conjugated Streptavidin diluted in ELISA buffer is added to each well and the plates incubated for 1 hour. Plates are washed and bound antibodies were detected by addition of substrate (e.g., TMB, Biofx Laboratories Inc., Owings Mills, MD). The reaction is terminated by addition of stop buffer (e.g., Bio FX Stop Reagents, Biofx Laboratories Inc., Owings Mills, MD) and the absorbance is measured at 650 nm using microplate reader (e.g., VERSAmax, Molecular Devices, Sunnyvale, CA).

[0140] Variations on this competition assay can also be used to test competition between 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 and another anti-glyco-cMET antibody. For example, in certain aspects, the anti-glyco-cMET antibody is used as a reference antibody and 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 is used as a test antibody. Additionally, instead of a glycosylated cMET peptide of SEQ ID NO:285, membrane-bound glyco-cMET expressed on cell surface (for example on the surface of one of the cell types mentioned above) in culture can be used. Generally, about 10^4 to 10^6 transfectants, e.g., about 10^5 transfectants, are used. Other formats for competition assays are known in the art and can be employed.

[0141] In various embodiments, an anti-glyco-cMET antibody of the disclosure reduces the binding of labeled 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 by at least 40%, by at least 50%, by at least 60%, by at least 70%, by at least 80%, by at least 90%, or by a percentage ranging

between any of the foregoing values (e.g., an anti-glyco-cMET antibody of the disclosure reduces the binding of labeled 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 by 50% to 70%) when the anti-glyco-cMET antibody is used at a concentration of 0.08 µg/mL, 0.4 µg/mL, 2 µg/mL, 10 µg/mL, 50 µg/mL, 100 µg/mL or at a concentration ranging between any of the foregoing values (e.g., at a concentration ranging from 2 µg/mL to 10 µg/mL).

[0142] In other embodiments, 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 reduces the binding of a labeled anti-glyco-cMET antibody of the disclosure by at least 40%, by at least 50%, by at least 60%, by at least 70%, by at least 80%, by at least 90%, or by a percentage ranging between any of the foregoing values (e.g., 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 reduces the binding of a labeled an anti-glyco-cMET antibody of the disclosure by 50% to 70%) when 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 is used at a concentration of 0.4 µg/mL, 2 µg/mL, 10 µg/mL, 50 µg/mL, 250 µg/mL or at a concentration ranging between any of the foregoing values (e.g., at a concentration ranging from 2 µg/mL to 10 µg/mL).

[0143] In the foregoing assays, the 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 antibody can be replaced by any antibody or antigen-binding fragment comprising the CDRs or the heavy and light chain variable regions of 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3, such as a humanized or chimeric counterpart of 3C7, 13C3, or 13G2. Exemplary humanized heavy and light chain variable regions of 8H3 are provided in Tables 4A-4G.

[0144] In certain aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure has an epitope which is the same or similar to the epitope of 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3. The epitope of an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure can be characterized, for example, by performing alanine scanning. A library of glycopeptides, each varying from the cMET glycopeptide (SEQ ID NO:285) by an alanine point mutation at one amino acid position of SEQ ID NO:285 (or, where the cMET peptide has an alanine, by a glycine point mutation). By measuring an antibody or antigen binding fragment's binding to each of the peptides by ELISA, the antibody or antigen binding fragment's epitope can be mapped.

[0145] In certain aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain variable sequences (or encoded by the nucleotide sequences) set forth in Tables 1A-1F (murine) and 4A-4G (humanized). In other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure comprises heavy and/or light chain CDR sequences (or encoded by the nucleotide sequences) set forth in Tables 1A-3H. The framework sequences for such anti-glyco-cMET antibody and antigen-binding fragment can be the native murine framework sequences of the VH and VL sequences set forth in Tables 1A-1F or can be non-native (e.g., humanized or human) framework sequences. Humanized framework sequences of the VH and VL sequences of 8H3 are set forth in Tables 4A-4G.

[0146] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS: 1 and 2, respectively.

[0147] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS: 23 and 24, respectively.

[0148] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS: 45 and 46, respectively.

[0149] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS: 67 and 68, respectively.

[0150] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS: 89 and 90, respectively.

[0151] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having heavy and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of SEQ ID NOS: 111 and 112, respectively.

[0152] In yet other aspects, the disclosure provides an anti-cMET antibody or antigen binding fragment having a heavy chain variable region having at least 95%, 98%, 99%, or 99.5% sequence identity of one of SEQ ID NOS:264-275 and light chain variable regions having at least 95%, 98%, 99%, or 99.5% sequence identity of one of SEQ ID NOS:276-284.

[0153] In yet other aspects, an anti-glyco-cMET antibody or antigen-binding fragment of the disclosure is a single-chain variable fragment (scFv). An exemplary scFv comprises the heavy chain variable fragment N-terminal to the light chain variable fragment. Another exemplary scFv comprises the light chain variable fragment N-terminal to the heavy chain variable fragment. In some embodiments, the scFv heavy chain variable fragment and light chain variable fragment are covalently bound to a linker sequence of 4-15 amino acids. The scFv can be in the form of a bi-specific T-cell engager or within a chimeric antigen receptor (CAR).

5.1.1. Antibody Specificity

[0154] In some embodiments, the anti-glyco-cMET antibodies of the disclosure specifically bind to the cMET glycoprotein PTKSFISGG**ST**ITGVGKLN (SEQ ID NO:285), glycosylated with GalNAc on the serine and threonine residues shown in bold underlined text. In certain embodiments, the anti-glyco-cMET antibodies of the disclosure specifically binds to the cMET glycoprotein, and does not specifically bind to one or more of: the unglycosylated cMET peptide

PTKSFISGGSTITGVGKNLN (SEQ ID NO:286) (the “unglycosylated cMET peptide”); the MUC1 tandem repeat (VTSAPDTRPAPGSTAPPAHG)₃ (SEQ ID NO:288) that has been glycosylated *in vitro* using purified recombinant human glycosyltransferases GalNAc-T1, GalNAc-T2, and GalNAc-T4 (“the first MUC1 glycopeptide”); the MUC1 peptide TAPPAHGVT**TS**APD**TR**PAPG**ST**APPAHGVT (SEQ ID NO:289) that has been glycosylated *in vitro* with GalNAc on the serine and threonine residues shown with bold and underlined text (the “second MUC1 glycopeptide”); the podoplanin peptide ERG**IK**PPLEELSGK (SEQ ID NO:290) that has been glycosylated *in vitro* with GalNAc on the threonine residue shown with bold and underlined text (the “PDPN glycopeptide”); the CD44v6 peptide GYRQ**IP**KEDSH**ST**TGTAAA (SEQ ID NO:345) that has been glycosylated *in vitro* with GalNAc on the threonine and serine residues shown with bold and underlined text (the “CD44v6 glycopeptide”); the MUC4 peptide CTIPSTAMHTR**ST**AAPILP (SEQ ID NO:291) that has been glycosylated *in vitro* with GalNAc on the serine and threonine residues shown with bold and underlined text (the “MUC4 glycopeptide”); and the LAMP1 peptide CEQDRP**SPT**IAPPAPPSPSP (SEQ ID NO:292) that has been glycosylated *in vitro* with GalNAc on the serine and threonine residues shown with bold and underlined text (the “LAMP1 glycopeptide”).

[0155] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the unglycosylated cMET peptide.

[0156] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the first MUC1 glycopeptide.

[0157] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the second MUC1 glycopeptide.

[0158] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the PDPN glycopeptide.

[0159] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the CD44v6 glycopeptide.

[0160] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the MUC4 glycopeptide.

[0161] In some embodiments, an anti-glyco-cMET antibody of the disclosure has a binding affinity to the cMET glycopeptide which is at least 3 times, at least 5 times, at least 10 times, at least 20 times, at least 50 times, at least 100 times, or at least 1000 times the binding affinity of the anti-glyco-cMET antibody to the LAMP1 glycopeptide.

[0162] Assays for determining affinity, including relative affinity, include but are not limited to a radioactive material labeled immunoassay (RIA), an enzyme-linked immunosorbent assay (ELISA), a sandwich ELISA, fluorescence activated cell sorting (FACS) assays, surface plasmon resonance (e.g., Biacore) assays, and bio-layer interferometry (BLI) assays. In some embodiments, affinity is measured by surface plasmon resonance (e.g., Biacore). In other embodiments, affinity

[0163] Exemplary anti-glyco-cMET antibody and fragments thereof are described in numbered embodiments 1 to 657.

5.2 Antibody-Drug Conjugates

[0164] Another aspect of the disclosure concerns antibody drug conjugates (ADCs) including the anti-glyco-cMET antibodies and antigen-binding fragments of the disclosure. The ADCs generally comprise an anti-glyco-cMET antibody and/or binding fragment as described herein having one or more cytotoxic and/or cytostatic agents linked thereto by way of one or more linkers. In specific embodiments, the ADCs are compounds according to structural formula (I):



or salts thereof, where each "D" represents, independently of the others, a cytotoxic and/or cytostatic agent ("drug"); each "L" represents, independently of the others, a linker; "Ab" represents an anti-glyco-cMET antigen binding domain, such as an anti-glyco-cMET antibody or binding fragment described herein; each "XY" represents a linkage formed between a functional group R^x on the linker and a "complementary" functional group R^y on the antibody, and n represents the number of drugs linked to, or drug-to-antibody ratio (DAR), of the ADC.

[0165] Specific embodiments of the various antibodies (Ab) that can comprise the ADCs include the various embodiments of anti-glyco-cMET antibodies and/or binding fragments described above.

[0166] In some specific embodiments of the ADCs and/or salts of structural formula (I), each D is the same and/or each L is the same.

[0167] Specific embodiments of cytotoxic and/or cytostatic agents (D) and linkers (L) that can comprise the anti-glyco-cMET ADCs of the disclosure, as well as the number of cytotoxic and/or cytostatic agents linked to the ADCs, are described in more detail below.

5.2.1. Cytotoxic and/or Cytostatic Agents

[0168] The cytotoxic and/or cytostatic agents may be any agents known to inhibit the growth and/or replication of and/or kill cells, and in particular cancer and/or tumor cells. Numerous agents having cytotoxic and/or cytostatic properties are known in the literature. Non-limiting examples of classes of cytotoxic and/or cytostatic agents include, by way of example and not limitation, radionuclides, alkylating agents, topoisomerase I inhibitors, topoisomerase II inhibitors, DNA intercalating agents (*e.g.*, groove binding agents such as minor groove binders), RNA/DNA antimetabolites, cell cycle modulators, kinase inhibitors, protein synthesis inhibitors, histone deacetylase inhibitors, mitochondria inhibitors, and antimetabolic agents.

[0169] Specific non-limiting examples of agents within certain of these various classes are provided below.

[0170] Alkylating Agents: asaley ((L-Leucine, N-[N-acetyl-4-[bis-(2-chloroethyl)amino]-DL-phenylalanyl]-, ethylester; NSC 167780; CAS Registry No. 3577897)); AZQ ((1,4-cyclohexadiene-1,4-dicarbamic acid, 2,5-bis(1-aziridinyl)-3,6-dioxo-, diethyl ester; NSC 182986; CAS Registry No. 57998682)); BCNU ((N,N'-Bis(2-chloroethyl)-N-nitrosourea; NSC 409962; CAS Registry No. 154938)); busulfan (1,4-butanediol dimethanesulfonate; NSC 750; CAS Registry No. 55981); (carboxyphthalato)platinum (NSC 27164; CAS Registry No. 65296813); CBDCA ((cis-(1,1-cyclobutanedicarboxylato)diammineplatinum(II)); NSC 241240; CAS Registry No. 41575944)); CCNU ((N-(2-chloroethyl)-N'-cyclohexyl-N-nitrosourea; NSC 79037; CAS Registry No. 13010474)); CHIP (iproplatin; NSC 256927); chlorambucil (NSC 3088; CAS Registry No. 305033); chlorozotocin ((2-[[[(2-chloroethyl) nitrosoamino]carbonyl]amino]-2-deoxy-D-glucopyranose; NSC 178248; CAS Registry No. 54749905)); cis-platinum (cisplatin; NSC 119875; CAS Registry No. 15663271); clomesone (NSC 338947; CAS Registry No. 88343720); cyanomorpholinodoxorubicin (NCS 357704; CAS Registry No. 88254073); cyclodisone (NSC 348948; CAS Registry No. 99591738); dianhydrogalactitol (5,6-diepoxydulcitol; NSC 132313; CAS Registry No. 23261203); fluorodopan ((5-[(2-chloroethyl)-(2-fluoroethyl)amino]-6-methyl-uracil; NSC 73754; CAS Registry No. 834913); hepsulfam (NSC 329680; CAS Registry No. 96892578); hycanthone (NSC 142982; CAS Registry No. 23255938); melphalan (NSC 8806; CAS Registry No. 3223072); methyl CCNU ((1-(2-chloroethyl)-3-(trans-4-methylcyclohexane)-1-nitrosourea; NSC 95441; 13909096); mitomycin C (NSC 26980; CAS Registry No. 50077); mitozolamide (NSC 353451; CAS Registry No. 85622953); nitrogen mustard ((bis(2-chloroethyl)methylamine hydrochloride; NSC 762; CAS Registry No. 55867); PCNU ((1-(2-chloroethyl)-3-(2,6-dioxo-3-piperidyl)-1-nitrosourea; NSC 95466; CAS Registry No. 13909029)); piperazine alkylator ((1-(2-chloroethyl)-4-(3-

chloropropyl)-piperazine dihydrochloride; NSC 344007)); piperazinedione (NSC 135758; CAS Registry No. 41109802); pipobroman ((N,N-bis(3-bromopropionyl) piperazine; NSC 25154; CAS Registry No. 54911)); porfiromycin (N-methylmitomycin C; NSC 56410; CAS Registry No. 801525); spirohydantoin mustard (NSC 172112; CAS Registry No. 56605164); teroxirone (triglycidylisocyanurate; NSC 296934; CAS Registry No. 2451629); tetraplatin (NSC 363812; CAS Registry No. 62816982); thio-tepa (N,N',N''-tri-1,2-ethanediythio phosphoramidate; NSC 6396; CAS Registry No. 52244); triethylenemelamine (NSC 9706; CAS Registry No. 51183); uracil nitrogen mustard (desmethyltopan; NSC 34462; CAS Registry No. 66751); Yoshi-864 ((bis(3-mesyloxy propyl)amine hydrochloride; NSC 102627; CAS Registry No. 3458228).

[0171] Topoisomerase I Inhibitors: camptothecin (NSC 94600; CAS Registry No. 7689-03-4); various camptothecin derivatives and analogs (for example, NSC 100880, NSC 603071, NSC 107124, NSC 643833, NSC 629971, NSC 295500, NSC 249910, NSC 606985, NSC 74028, NSC 176323, NSC 295501, NSC 606172, NSC 606173, NSC 610458, NSC 618939, NSC 610457, NSC 610459, NSC 606499, NSC 610456, NSC 364830, and NSC 606497); morpholinisoxorubicin (NSC 354646; CAS Registry No. 89196043); SN-38 (NSC 673596; CAS Registry No. 86639-52-3).

[0172] Topoisomerase II Inhibitors: doxorubicin (NSC 123127; CAS Registry No. 25316409); amonafide (benzisoquinolinedione; NSC 308847; CAS Registry No. 69408817); m-AMSA ((4'-(9-acridinylamino)-3'-methoxymethanesulfonanilide; NSC 249992; CAS Registry No. 51264143)); anthrapyrazole derivative ((NSC 355644); etoposide (VP-16; NSC 141540; CAS Registry No. 33419420); pyrazoloacridine ((pyrazolo[3,4,5-kl]acridine-2(6H)-propanamine, 9-methoxy-N, N-dimethyl-5-nitro-, monomethanesulfonate; NSC 366140; CAS Registry No. 99009219); bisantrene hydrochloride (NSC 337766; CAS Registry No. 71439684); daunorubicin (NSC 821151; CAS Registry No. 23541506); deoxydoxorubicin (NSC 267469; CAS Registry No. 63950061); mitoxantrone (NSC 301739; CAS Registry No. 70476823); menogaril (NSC 269148; CAS Registry No. 71628961); N,N-dibenzyl daunomycin (NSC 268242; CAS Registry No. 70878512); oxanthrazole (NSC 349174; CAS Registry No. 105118125); rubidazone (NSC 164011; CAS Registry No. 36508711); teniposide (VM-26; NSC 122819; CAS Registry No. 29767202).

[0173] DNA Intercalating Agents: anthramycin (CAS Registry No. 4803274); chicamycin A (CAS Registry No. 89675376); tomaymycin (CAS Registry No. 35050556); DC-81 (CAS Registry No. 81307246); sibiromycin (CAS Registry No. 12684332); pyrrolobenzodiazepine derivative (CAS Registry No. 945490095); SGD-1882 ((S)-2-(4-aminophenyl)-7-methoxy-8-(3-4(S)-7-methoxy-2-(4-methoxyphenyl)-- 5-oxo-5,11a-dihydro-1H-benzo[e]pyrrolo[1,2-a][1,4]diazepin-8-yl)oxy)propox- y)-1H-benzo[e]pyrrolo[1,2-a][1,4]diazepin-5(11aH)-one); SG2000 (SJG-136; (11aS,11a'S)-8,8'-(propane-1,3-diylbis(oxy))bis(7-methoxy-2-methylene-

2,3- -dihydro-1H-benzo[e]pyrrolo[1,2-a][1,4]diazepin-5(11aH)-one); NSC 694501; CAS Registry No. 232931576).

[0174] RNA/DNA Antimetabolites: L-alanosine (NSC 153353; CAS Registry No. 59163416); 5-azacytidine (NSC 102816; CAS Registry No. 320672); 5-fluorouracil (NSC 19893; CAS Registry No. 51218); acivicin (NSC 163501; CAS Registry No. 42228922); aminopterin derivative N-[2-chloro-5-[[[(2,4-diamino-5-methyl-6-quinazoliny)methyl]amino]benzoyl-]L-aspartic acid (NSC 132483); aminopterin derivative N-[4-[[[(2,4-diamino-5-ethyl-6-quinazoliny)methyl]amino]benzoyl]L-aspartic acid (NSC 184692); aminopterin derivative N-[2-chloro-4-[[[(2,4-diamino-6-pteridiny)methyl]amino]benzoyl]L-aspartic acid monohydrate (NSC 134033); an antifo ((N^α-(4-amino-4-deoxypteroyl)-N⁷-hemiphthaloyl-L-ornithin- e; NSC 623017)); Baker's soluble antifol (NSC 139105; CAS Registry No. 41191042); dichlorallyl lawsone ((2-(3,3-dichloroallyl)-3-hydroxy-1,4-naphthoquinone; NSC 126771; CAS Registry No. 36417160); brequinar (NSC 368390; CAS Registry No. 96201886); ftorafur ((pro-drug; 5-fluoro-1-(tetrahydro-2-furyl)-uracil; NSC 148958; CAS Registry No. 37076689); 5,6-dihydro-5-azacytidine (NSC 264880; CAS Registry No. 62402317); methotrexate (NSC 740; CAS Registry No. 59052); methotrexate derivative (N-[[4-[[[(2,4-diamino-6-pteridiny)methyl]methylamino]-1-naphthaleny]carbonyl]L-glutamic acid; NSC 174121); PALA ((N-(phosphonoacetyl)-L-aspartate; NSC 224131; CAS Registry No. 603425565); pyrazofurin (NSC 143095; CAS Registry No. 30868305); trimetrexate (NSC 352122; CAS Registry No. 82952645).

[0175] DNA Antimetabolites: 3-HP (NSC 95678; CAS Registry No. 3814797); 2'-deoxy-5-fluorouridine (NSC 27640; CAS Registry No. 50919); 5-HP (NSC 107392; CAS Registry No. 19494894); α-TGDR (α-2'-deoxy-6-thioguanosine; NSC 71851 CAS Registry No. 2133815); aphidicolin glycinate (NSC 303812; CAS Registry No. 92802822); ara C (cytosine arabinoside; NSC 63878; CAS Registry No. 69749); 5-aza-2'-deoxycytidine (NSC 127716; CAS Registry No. 2353335); β-TGDR (β-2'-deoxy-6-thioguanosine; NSC 71261; CAS Registry No. 789617); cyclocytidine (NSC 145668; CAS Registry No. 10212256); guanazole (NSC 1895; CAS Registry No. 1455772); hydroxyurea (NSC 32065; CAS Registry No. 127071); inosine glycodialdehyde (NSC 118994; CAS Registry No. 23590990); macbecin II (NSC 330500; CAS Registry No. 73341738); pyrazoloimidazole (NSC 51143; CAS Registry No. 6714290); thioguanine (NSC 752; CAS Registry No. 154427); thiopurine (NSC 755; CAS Registry No. 50442).

[0176] Cell Cycle Modulators: silibinin (CAS Registry No. 22888-70-6); epigallocatechin gallate (EGCG; CAS Registry No. 989515); procyanidin derivatives (e.g., procyanidin A1 [CAS Registry No. 103883030], procyanidin B1 [CAS Registry No. 20315257], procyanidin B4 [CAS Registry No. 29106512], arecatannin B1 [CAS Registry No. 79763283]); isoflavones (e.g., genistein [4%5,7-trihydroxyisoflavone; CAS Registry No. 446720], daidzein [4',7-

dihydroxyisoflavone, CAS Registry No. 486668]; indole-3-carbinol (CAS Registry No. 700061); quercetin (NSC 9219; CAS Registry No. 117395); estramustine (NSC 89201; CAS Registry No. 2998574); nocodazole (CAS Registry No. 31430189); podophyllotoxin (CAS Registry No. 518285); vinorelbine tartrate (NSC 608210; CAS Registry No. 125317397); cryptophycin (NSC 667642; CAS Registry No. 124689652).

[0177] Kinase Inhibitors: afatinib (CAS Registry No. 850140726); axitinib (CAS Registry No. 319460850); ARRY-438162 (binimetinib) (CAS Registry No. 606143899); bosutinib (CAS Registry No. 380843754); cabozantinib (CAS Registry No. 1140909483); ceritinib (CAS Registry No. 1032900256); crizotinib (CAS Registry No. 877399525); dabrafenib (CAS Registry No. 1195765457); dasatinib (NSC 732517; CAS Registry No. 302962498); erlotinib (NSC 718781; CAS Registry No. 183319699); everolimus (NSC 733504; CAS Registry No. 159351696); fostamatinib (NSC 745942; CAS Registry No. 901119355); gefitinib (NSC 715055; CAS Registry No. 184475352); ibrutinib (CAS Registry No. 936563961); imatinib (NSC 716051; CAS Registry No. 220127571); lapatinib (CAS Registry No. 388082788); lenvatinib (CAS Registry No. 857890392); mubritinib (CAS 366017096); nilotinib (CAS Registry No. 923288953); nintedanib (CAS Registry No. 656247175); palbociclib (CAS Registry No. 571190302); pazopanib (NSC 737754; CAS Registry No. 635702646); pegaptanib (CAS Registry No. 222716861); ponatinib (CAS Registry No. 1114544318); rapamycin (NSC 226080; CAS Registry No. 53123889); regorafenib (CAS Registry No. 755037037); AP 23573 (ridaforolimus) (CAS Registry No. 572924540); INCB018424 (ruxolitinib) (CAS Registry No. 1092939177); ARRY-142886 (selumetinib) (NSC 741078; CAS Registry No. 606143-52-6); sirolimus (NSC 226080; CAS Registry No. 53123889); sorafenib (NSC 724772; CAS Registry No. 475207591); sunitinib (NSC 736511; CAS Registry No. 341031547); tofacitinib (CAS Registry No. 477600752); temsirolimus (NSC 683864; CAS Registry No. 163635043); trametinib (CAS Registry No. 871700173); vandetanib (CAS Registry No. 443913733); vemurafenib (CAS Registry No. 918504651); SU6656 (CAS Registry No. 330161870); CEP-701 (lesaurtinib) (CAS Registry No. 111358884); XL019 (CAS Registry No. 945755566); PD-325901 (CAS Registry No. 391210109); PD-98059 (CAS Registry No. 167869218); ATP-competitive TORC1/TORC2 inhibitors including PI-103 (CAS Registry No. 371935749), PP242 (CAS Registry No. 1092351671), PP30 (CAS Registry No. 1092788094), Torin 1 (CAS Registry No. 1222998368), LY294002 (CAS Registry No. 154447366), XL-147 (CAS Registry No. 934526893), CAL-120 (CAS Registry No. 870281348), ETP-45658 (CAS Registry No. 1198357797), PX 866 (CAS Registry No. 502632668), GDC-0941 (CAS Registry No. 957054307), BGT226 (CAS Registry No. 1245537681), BEZ235 (CAS Registry No. 915019657), XL-765 (CAS Registry No. 934493762).

[0178] Protein Synthesis Inhibitors: acriflavine (CAS Registry No. 65589700); amikacin (NSC 177001; CAS Registry No. 39831555); arbekacin (CAS Registry No. 51025855); astromicin

(CAS Registry No. 55779061); azithromycin (NSC 643732; CAS Registry No. 83905015); bekanamycin (CAS Registry No. 4696768); chlortetracycline (NSC 13252; CAS Registry No. 64722); clarithromycin (NSC 643733; CAS Registry No. 81103119); clindamycin (CAS Registry No. 18323449); clomocycline (CAS Registry No. 1181540); cycloheximide (CAS Registry No. 66819); dactinomycin (NSC 3053; CAS Registry No. 50760); dalfopristin (CAS Registry No. 112362502); demeclocycline (CAS Registry No. 127333); dibekacin (CAS Registry No. 34493986); dihydrostreptomycin (CAS Registry No. 128461); dirithromycin (CAS Registry No. 62013041); doxycycline (CAS Registry No. 17086281); emetine (NSC 33669; CAS Registry No. 483181); erythromycin (NSC 55929; CAS Registry No. 114078); flurithromycin (CAS Registry No. 83664208); framycetin (neomycin B; CAS Registry No. 119040); gentamycin (NSC 82261; CAS Registry No. 1403663); glycylicyclines, such as tigecycline (CAS Registry No. 220620097); hygromycin B (CAS Registry No. 31282049); isepamicin (CAS Registry No. 67814760); josamycin (NSC 122223; CAS Registry No. 16846245); kanamycin (CAS Registry No. 8063078); ketolides such as telithromycin (CAS Registry No. 191114484), cethromycin (CAS Registry No. 205110481), and solithromycin (CAS Registry No. 760981837); lincomycin (CAS Registry No. 154212); lymecycline (CAS Registry No. 992212); meclocycline (NSC 78502; CAS Registry No. 2013583); metacycline (rondomycin; NSC 356463; CAS Registry No. 914001); midecamycin (CAS Registry No. 35457808); minocycline (NSC 141993; CAS Registry No. 10118908); miocamycin (CAS Registry No. 55881077); neomycin (CAS Registry No. 119040); netilmicin (CAS Registry No. 56391561); oleandomycin (CAS Registry No. 3922905); oxazolidinones, such as eperezolid (CAS Registry No. 165800044), linezolid (CAS Registry No. 165800033), posizolid (CAS Registry No. 252260029), radezolid (CAS Registry No. 869884786), ranbezolid (CAS Registry No. 392659380), sutezolid (CAS Registry No. 168828588), tedizolid (CAS Registry No. 856867555); oxytetracycline (NSC 9169; CAS Registry No. 2058460); paromomycin (CAS Registry No. 7542372); penimepicycline (CAS Registry No. 4599604); peptidyl transferase inhibitors, e.g., chloramphenicol (NSC 3069; CAS Registry No. 56757) and derivatives such as azidamfenicol (CAS Registry No. 13838089), florfenicol (CAS Registry No. 73231342), and thiamphenicol (CAS Registry No. 15318453), and pleuromutilins such as retapamulin (CAS Registry No. 224452668), tiamulin (CAS Registry No. 55297955), valnemulin (CAS Registry No. 101312929); pirlimycin (CAS Registry No. 79548735); puromycin (NSC 3055; CAS Registry No. 53792); quinupristin (CAS Registry No. 120138503); ribostamycin (CAS Registry No. 53797356); rokitamycin (CAS Registry No. 74014510); rolitetracycline (CAS Registry No. 751973); roxithromycin (CAS Registry No. 80214831); sisomicin (CAS Registry No. 32385118); spectinomycin (CAS Registry No. 1695778); spiramycin (CAS Registry No. 8025818); streptogramins such as pristinamycin (CAS Registry No. 270076603), quinupristin/dalfopristin (CAS Registry No. 126602899), and virginiamycin (CAS Registry No. 11006761); streptomycin (CAS Registry No. 57921); tetracycline (NSC 108579; CAS Registry No. 60548); tobramycin (CAS Registry No.

32986564); troleandomycin (CAS Registry No. 2751099); tylosin (CAS Registry No. 1401690); verdamicin (CAS Registry No. 49863481).

[0179] Histone Deacetylase Inhibitors: abexinostat (CAS Registry No. 783355602); belinostat (NSC 726630; CAS Registry No. 414864009); chidamide (CAS Registry No. 743420022); entinostat (CAS Registry No. 209783802); givinostat (CAS Registry No. 732302997); mocetinostat (CAS Registry No. 726169739); panobinostat (CAS Registry No. 404950807); quisinostat (CAS Registry No. 875320299); resminostat (CAS Registry No. 864814880); romidepsin (CAS Registry No. 128517077); sulforaphane (CAS Registry No. 4478937); thioureaidobutyronitrile (Kevetrin™; CAS Registry No. 6659890); valproic acid (NSC 93819; CAS Registry No. 99661); vorinostat (NSC 701852; CAS Registry No. 149647789); ACY-1215 (rocilinostat; CAS Registry No. 1316214524); CUDC-101 (CAS Registry No. 1012054599); CHR-2845 (tefinostat; CAS Registry No. 914382608); CHR-3996 (CAS Registry No. 1235859138); 4SC-202 (CAS Registry No. 910462430); CG200745 (CAS Registry No. 936221339); SB939 (pracinostat; CAS Registry No. 929016966).

[0180] Mitochondria Inhibitors: pancratistatin (NSC 349156; CAS Registry No. 96281311); rhodamine-123 (CAS Registry No. 63669709); edelfosine (NSC 324368; CAS Registry No. 70641519); d-alpha-tocopherol succinate (NSC 173849; CAS Registry No. 4345033); compound 11β (CAS Registry No. 865070377); aspirin (NSC 406186; CAS Registry No. 50782); ellipticine (CAS Registry No. 519233); berberine (CAS Registry No. 633658); cerulenin (CAS Registry No. 17397896); GX015-070 (Obatoclox®; 1H-Indole, 2-(2-((3,5-dimethyl-1H-pyrrol-2-yl)methylene)-3-methoxy-2H-pyrrol-5-yl)-; NSC 729280; CAS Registry No. 803712676); celastrol (tripterine; CAS Registry No. 34157830); metformin (NSC 91485; CAS Registry No. 1115704); Brilliant green (NSC 5011; CAS Registry No. 633034); ME-344 (CAS Registry No. 1374524556).

[0181] Antimitotic Agents: allocolchicine (NSC 406042); auristatins, such as MMAE (monomethyl auristatin E; CAS Registry No. 474645-27-7) and MMAF (monomethyl auristatin F; CAS Registry No. 745017-94-1; halichondrin B (NSC 609395); colchicine (NSC 757; CAS Registry No. 64868); cholchicine derivative (N-benzoyl-deacetyl benzamide; NSC 33410; CAS Registry No. 63989753); dolastatin 10 (NSC 376128; CAS Registry No. 110417-88-4); maytansine (NSC 153858; CAS Registry No. 35846-53-8); rhozoxin (NSC 332598; CAS Registry No. 90996546); taxol (NSC 125973; CAS Registry No. 33069624); taxol derivative ((2'-N-[3-(dimethylamino)propyl]glutaramate taxol; NSC 608832); thiocolchicine (3-demethylthiocolchicine; NSC 361792); trityl cysteine (NSC 49842; CAS Registry No. 2799077); vinblastine sulfate (NSC 49842; CAS Registry No. 143679); vincristine sulfate (NSC 67574; CAS Registry No. 2068782).

[0182] Any of these agents that include or that may be modified to include a site of attachment to an antibody may be included in the ADCs disclosed herein.

[0183] In a specific embodiment, the cytotoxic and/or cytostatic agent is an antimetabolic agent.

[0184] In another specific embodiment, the cytotoxic and/or cytostatic agent is an auristatin, for example, monomethyl auristatin E ("MMAE") or monomethyl auristatin F ("MMAF").

5.2.2. Linkers

[0185] In the anti-glyco-cMET ADCs of the disclosure, the cytotoxic and/or cytostatic agents are linked to the antibody by way of linkers. The linker linking a cytotoxic and/or cytostatic agent to the antibody of an ADC may be short, long, hydrophobic, hydrophilic, flexible or rigid, or may be composed of segments that each independently have one or more of the above-mentioned properties such that the linker may include segments having different properties. The linkers may be polyvalent such that they covalently link more than one agent to a single site on the antibody, or monovalent such that covalently they link a single agent to a single site on the antibody.

[0186] As will be appreciated by skilled artisans, the linkers link cytotoxic and/or cytostatic agents to the antibody by forming a covalent linkage to the cytotoxic and/or cytostatic agent at one location and a covalent linkage to antibody at another. The covalent linkages are formed by reaction between functional groups on the linker and functional groups on the agents and antibody. As used herein, the expression "linker" is intended to include (i) unconjugated forms of the linker that include a functional group capable of covalently linking the linker to a cytotoxic and/or cytostatic agent and a functional group capable of covalently linking the linker to an antibody; (ii) partially conjugated forms of the linker that includes a functional group capable of covalently linking the linker to an antibody and that is covalently linked to a cytotoxic and/or cytostatic agent, or vice versa; and (iii) fully conjugated forms of the linker that is covalently linked to both a cytotoxic and/or cytostatic agent and an antibody. In some specific embodiments of linkers and anti-glyco-cMET ADCs of the disclosure, as well as synthons used to conjugate linker-agents to antibodies, moieties comprising the functional groups on the linker and covalent linkages formed between the linker and antibody are specifically illustrated as R_x and XY, respectively.

[0187] The linkers are preferably, but need not be, chemically stable to conditions outside the cell, and may be designed to cleave, immolate and/or otherwise specifically degrade inside the cell. Alternatively, linkers that are not designed to specifically cleave or degrade inside the cell may be used. Choice of stable versus unstable linker may depend upon the toxicity of the cytotoxic and/or cytostatic agent. For agents that are toxic to normal cells, stable linkers are preferred. Agents that are selective or targeted and have lower toxicity to normal cells may utilize, chemical stability of the linker to the extracellular milieu is less important. A wide variety of linkers useful for linking drugs to antibodies in the context of ADCs are known in the art. Any of these linkers, as well as other linkers, may be used to link the cytotoxic and/or cytostatic agents to the antibody of the anti-glyco-cMET ADCs of the disclosure.

[0188] Exemplary polyvalent linkers that may be used to link many cytotoxic and/or cytostatic agents to a single antibody molecule are described, for example, in WO 2009/073445; WO 2010/068795; WO 2010/138719; WO 2011/120053; WO 2011/171020; WO 2013/096901; WO 2014/008375; WO 2014/093379; WO 2014/093394; WO 2014/093640, the content of which are incorporated herein by reference in their entireties. For example, the Fleximer linker technology developed by Mersana *et al.* has the potential to enable high-DAR ADCs with good physicochemical properties. As shown below, the Mersana technology is based on incorporating drug molecules into a solubilizing poly-acetal backbone via a sequence of ester bonds. The methodology renders highly-loaded ADCs (DAR up to 20) while maintaining good physicochemical properties.

[0189] Additional examples of dendritic type linkers can be found in US 2006/116422; US 2005/271615; de Groot *et al.* (2003) *Angew. Chem. Int. Ed.* 42:4490-4494; Amir *et al.* (2003) *Angew. Chem. Int. Ed.* 42:4494-4499; Shamis *et al.* (2004) *J. Am. Chem. Soc.* 126:1726-1731; Sun *et al.* (2002) *Bioorganic & Medicinal Chemistry Letters* 12:2213-2215; Sun *et al.* (2003) *Bioorganic & Medicinal Chemistry* 11:1761-1768; King *et al.* (2002) *Tetrahedron Letters* 43:1987-1990, each of which is incorporated herein by reference.

[0190] Exemplary monovalent linkers that may be used are described, for example, in Nolting, 2013, *Antibody-Drug Conjugates, Methods in Molecular Biology* 1045:71-100; Kitson *et al.*, 2013, *CROs/CMOs--Chemica Oggi--Chemistry Today* 31(4):30-38; Ducry *et al.*, 2010, *Bioconjugate Chem.* 21:5-13; Zhao *et al.*, 2011, *J. Med. Chem.* 54:3606-3623; U.S. Pat. No. 7,223,837; U.S. Pat. No. 8,568,728; U.S. Pat. No. 8,535,678; and WO2004010957, each of which is incorporated herein by reference.

[0191] By way of example and not limitation, some cleavable and noncleavable linkers that may be included in the anti-glyco-cMET ADCs of the disclosure are described below.

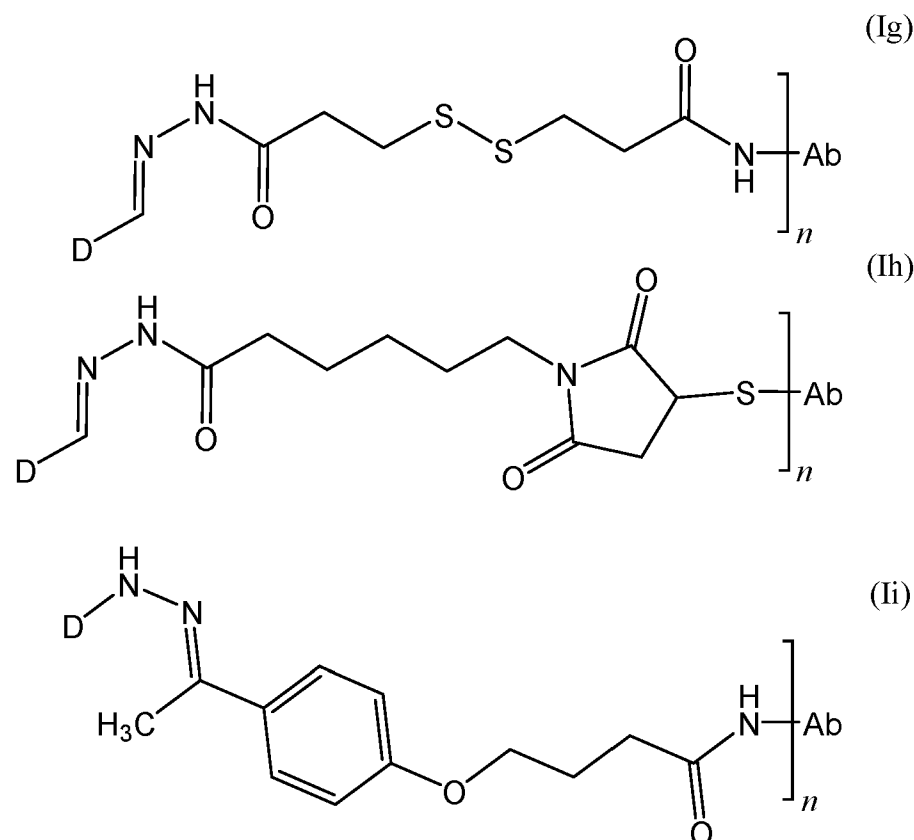
5.2.3. Cleavable Linkers

[0192] In certain embodiments, the linker selected is cleavable *in vivo*. Cleavable linkers may include chemically or enzymatically unstable or degradable linkages. Cleavable linkers generally rely on processes inside the cell to liberate the drug, such as reduction in the cytoplasm, exposure to acidic conditions in the lysosome, or cleavage by specific proteases or other enzymes within the cell. Cleavable linkers generally incorporate one or more chemical bonds that are either chemically or enzymatically cleavable while the remainder of the linker is noncleavable. In certain embodiments, a linker comprises a chemically labile group such as hydrazone and/or disulfide groups. Linkers comprising chemically labile groups exploit differential properties between the plasma and some cytoplasmic compartments. The intracellular conditions to facilitate drug release for hydrazone containing linkers are the acidic environment of endosomes and lysosomes, while the disulfide containing linkers are reduced in the cytosol, which contains high thiol concentrations, e.g., glutathione. In certain embodiments,

the plasma stability of a linker comprising a chemically labile group may be increased by introducing steric hindrance using substituents near the chemically labile group.

[0193] Acid-labile groups, such as hydrazone, remain intact during systemic circulation in the blood's neutral pH environment (pH 7.3-7.5) and undergo hydrolysis and release the drug once the ADC is internalized into mildly acidic endosomal (pH 5.0-6.5) and lysosomal (pH 4.5-5.0) compartments of the cell. This pH dependent release mechanism has been associated with nonspecific release of the drug. To increase the stability of the hydrazone group of the linker, the linker may be varied by chemical modification, e.g., substitution, allowing tuning to achieve more efficient release in the lysosome with a minimized loss in circulation.

[0194] Hydrazone-containing linkers may contain additional cleavage sites, such as additional acid-labile cleavage sites and/or enzymatically labile cleavage sites. ADCs including exemplary hydrazone-containing linkers include the following structures:



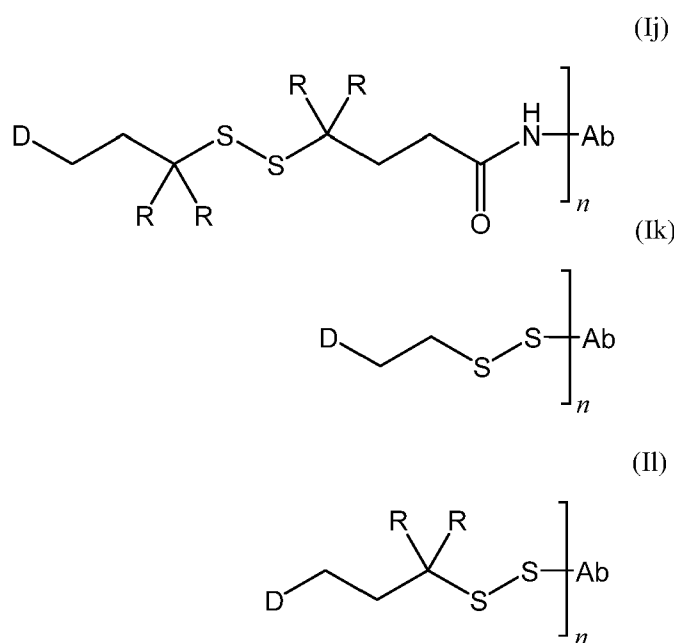
wherein D and Ab represent the cytotoxic and/or cytostatic agent (drug) and Ab, respectively, and n represents the number of drug-linkers linked to the antibody. In certain linkers such as linker (Ig), the linker comprises two cleavable groups--a disulfide and a hydrazone moiety. For such linkers, effective release of the unmodified free drug requires acidic pH or disulfide reduction and acidic pH. Linkers such as (Ih) and (Ii) have been shown to be effective with a single hydrazone cleavage site.

[0195] Additional linkers which remain intact during systemic circulation and undergo hydrolysis and release the drug when the ADC is internalized into acidic cellular compartments include carbonates. Such linkers can be useful in cases where the cytotoxic and/or cytostatic agent can be covalently attached through an oxygen.

[0196] Other acid-labile groups that may be included in linkers include cis-aconityl-containing linkers. cis-Aconityl chemistry uses a carboxylic acid juxtaposed to an amide bond to accelerate amide hydrolysis under acidic conditions.

[0197] Cleavable linkers may also include a disulfide group. Disulfides are thermodynamically stable at physiological pH and are designed to release the drug upon internalization inside cells, wherein the cytosol provides a significantly more reducing environment compared to the extracellular environment. Scission of disulfide bonds generally requires the presence of a cytoplasmic thiol cofactor, such as (reduced) glutathione (GSH), such that disulfide-containing linkers are reasonably stable in circulation, selectively releasing the drug in the cytosol. The intracellular enzyme protein disulfide isomerase, or similar enzymes capable of cleaving disulfide bonds, may also contribute to the preferential cleavage of disulfide bonds inside cells. GSH is reported to be present in cells in the concentration range of 0.5-10 mM compared with a significantly lower concentration of GSH or cysteine, the most abundant low-molecular weight thiol, in circulation at approximately 5 Tumor cells, where irregular blood flow leads to a hypoxic state, result in enhanced activity of reductive enzymes and therefore even higher glutathione concentrations. In certain embodiments, the *in vivo* stability of a disulfide-containing linker may be enhanced by chemical modification of the linker, e.g., use of steric hindrance adjacent to the disulfide bond.

[0198] ADCs including exemplary disulfide-containing linkers include the following structures:



wherein D and Ab represent the drug and antibody, respectively, n represents the number of drug-linkers linked to the antibody and R is independently selected at each occurrence from hydrogen or alkyl, for example. In certain embodiments, increasing steric hindrance adjacent to the disulfide bond increases the stability of the linker. Structures such as (Ij) and (II) show increased *in vivo* stability when one or more R groups is selected from a lower alkyl such as methyl.

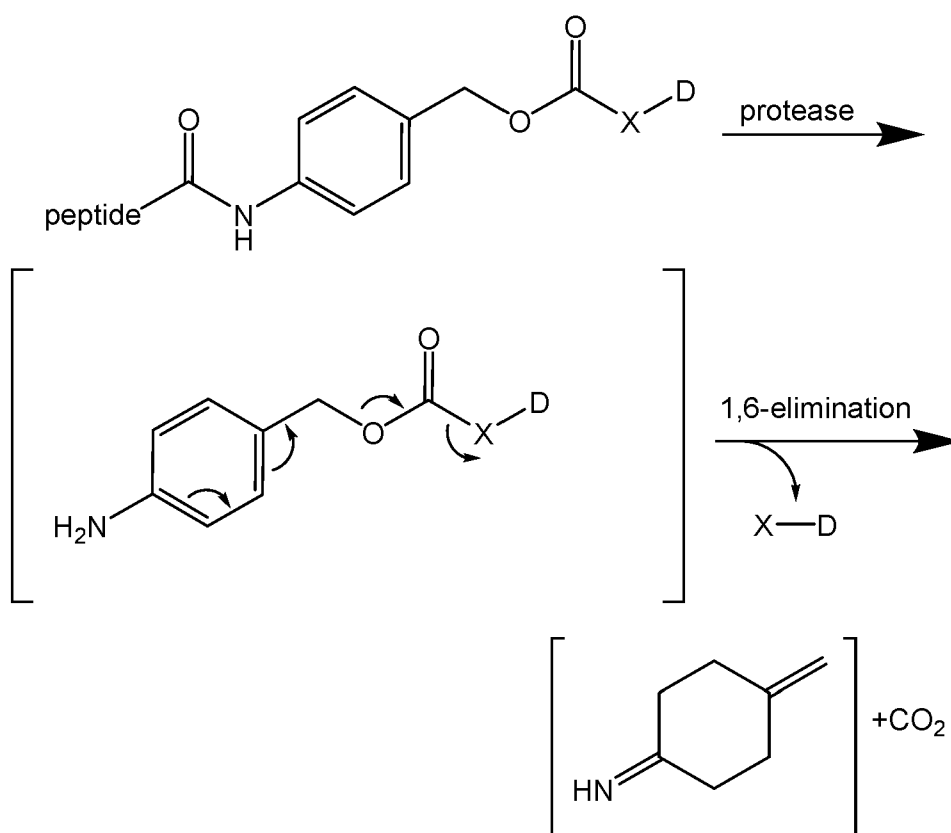
[0199] Another type of cleavable linker that may be used is a linker that is specifically cleaved by an enzyme. Such linkers are typically peptide-based or include peptidic regions that act as substrates for enzymes. Peptide based linkers tend to be more stable in plasma and extracellular milieu than chemically labile linkers. Peptide bonds generally have good serum stability, as lysosomal proteolytic enzymes have very low activity in blood due to endogenous inhibitors and the unfavorably high pH value of blood compared to lysosomes. Release of a drug from an antibody occurs specifically due to the action of lysosomal proteases, e.g., cathepsin and plasmin. These proteases may be present at elevated levels in certain tumor cells.

[0200] In exemplary embodiments, the cleavable peptide is selected from tetrapeptides such as Gly-Phe-Leu-Gly (SEQ ID NO:343), Ala-Leu-Ala-Leu (SEQ ID NO:344) or dipeptides such as Val-Cit, Val-Ala, Met-(D)Lys, Asn-(D)Lys, Val-(D)Asp, Phe-Lys, Ile-Val, Asp-Val, His-Val, NorVal-(D)Asp, Ala-(D)Asp 5, Met-Lys, Asn-Lys, Ile-Pro, Me3Lys-Pro, PhenylGly-(D)Lys, Met-(D)Lys, Asn-(D)Lys, Pro-(D)Lys, Met-(D)Lys, Asn-(D)Lys, AM Met-(D)Lys, Asn-(D)Lys, AW Met-(D)Lys, and Asn-(D)Lys. In certain embodiments, dipeptides are preferred over longer polypeptides due to hydrophobicity of the longer peptides.

[0201] A variety of dipeptide-based cleavable linkers useful for linking drugs such as doxorubicin, mitomycin, camptothecin, pyrrolobenzodiazepine, tallysomycin and auristatin/auristatin family members to antibodies have been described (see, Dubowchik *et al.*, 1998, J. Org. Chem. 67:1866-1872; Dubowchik *et al.*, 1998, Bioorg. Med. Chem. Lett. 8(21):3341-3346; Walker *et al.*, 2002, Bioorg. Med. Chem. Lett. 12:217-219; Walker *et al.*, 2004, Bioorg. Med. Chem. Lett. 14:4323-4327; Sutherland *et al.*, 2013, Blood 122: 1455-1463; and Francisco *et al.*, 2003, Blood 102:1458-1465, of each of which is incorporated herein by reference). All of these dipeptide linkers, or modified versions of these dipeptide linkers, may be used in the anti-glyco-cMET ADCs of the disclosure. Other dipeptide linkers that may be used include those found in ADCs such as Seattle Genetics' Brentuximab Vendotin SGN-35 (Adcetris™), Seattle Genetics SGN-75 (anti-CD-70, Val-Cit-monomethyl auristatin F (MMAF), Seattle Genetics SGN-CD33A (anti-CD-33, Val-Ala-(SGD-1882)), Celldex Therapeutics glembatumumab (CDX-011) (anti-NMB, Val-Cit-monomethyl auristatin E (MMAE), and Cytogen PSMA-ADC (PSMA-ADC-1301) (anti-PSMA, Val-Cit-MMAE).

[0202] Enzymatically cleavable linkers may include a self-immolative spacer to spatially separate the drug from the site of enzymatic cleavage. The direct attachment of a drug to a peptide linker can result in proteolytic release of an amino acid adduct of the drug, thereby impairing its activity. The use of a self-immolative spacer allows for the elimination of the fully active, chemically unmodified drug upon amide bond hydrolysis.

[0203] One self-immolative spacer is the bifunctional para-aminobenzyl alcohol group, which is linked to the peptide through the amino group, forming an amide bond, while amine containing drugs may be attached through carbamate functionalities to the benzylic hydroxyl group of the linker (PABC). The resulting prodrugs are activated upon protease-mediated cleavage, leading to a 1,6-elimination reaction releasing the unmodified drug, carbon dioxide, and remnants of the linker group. The following scheme depicts the fragmentation of p-amidobenzyl ether and release of the drug:

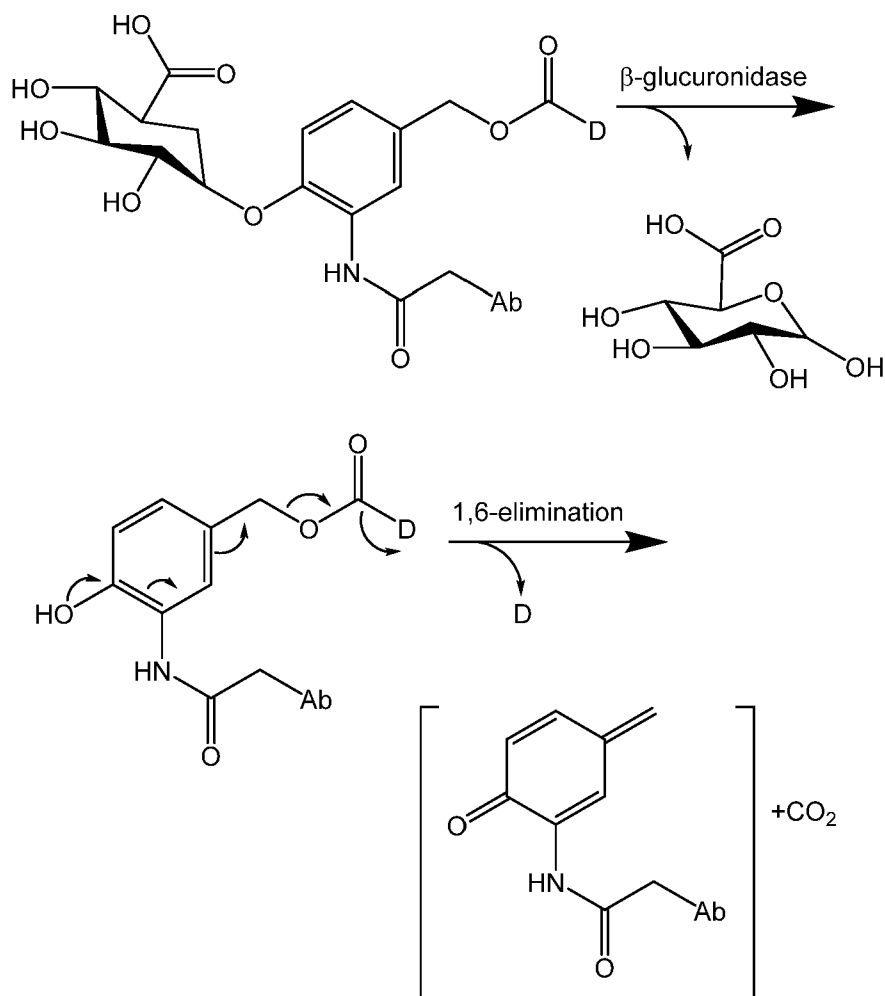


wherein X-D represents the unmodified drug.

[0204] Heterocyclic variants of this self-immolative group have also been described. See for example, U.S. Pat. No. 7,989,434, incorporated herein by reference.

[0205] In some embodiments, the enzymatically cleavable linker is a β -glucuronic acid-based linker. Facile release of the drug may be realized through cleavage of the β -glucuronide glycosidic bond by the lysosomal enzyme β -glucuronidase. This enzyme is present abundantly within lysosomes and is overexpressed in some tumor types, while the enzyme activity outside

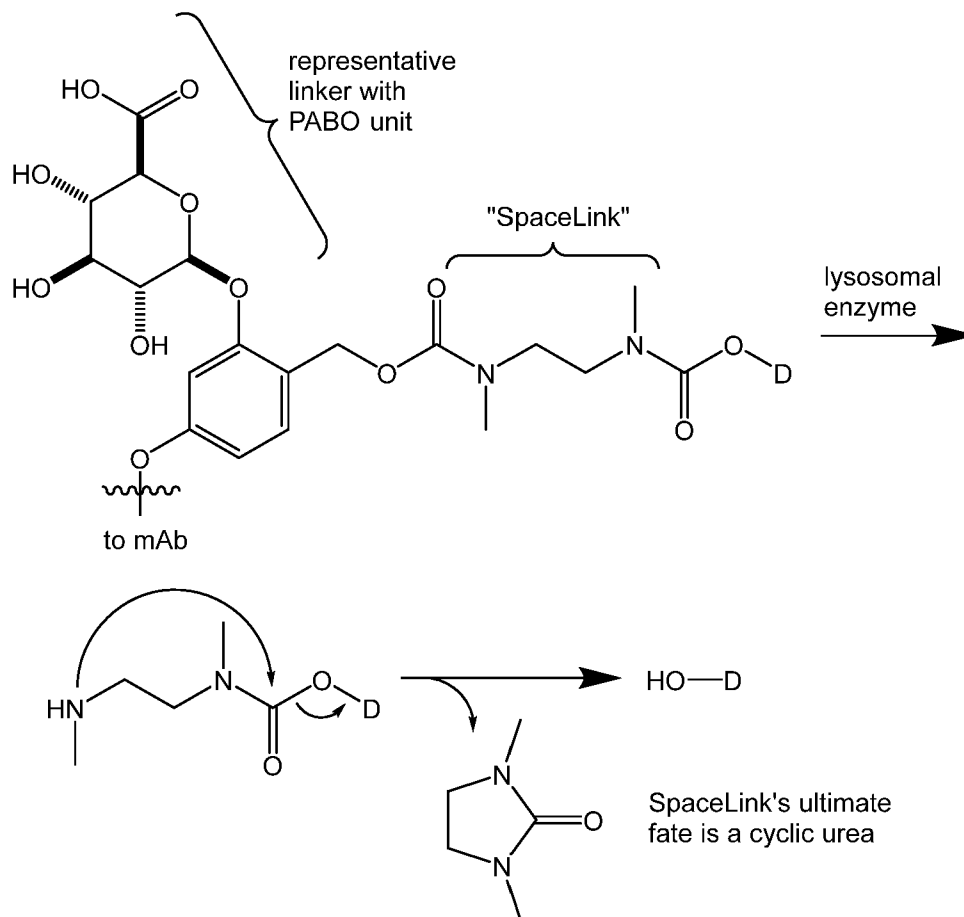
cells is low. β -Glucuronic acid-based linkers may be used to circumvent the tendency of an ADC to undergo aggregation due to the hydrophilic nature of β -glucuronides. In some embodiments, β -glucuronic acid-based linkers are preferred as linkers for ADCs linked to hydrophobic drugs. The following scheme depicts the release of the drug from an ADC containing a β -glucuronic acid-based linker:



[0206] A variety of cleavable β -glucuronic acid-based linkers useful for linking drugs such as auristatins, camptothecin and doxorubicin analogues, CBI minor-groove binders, and psymberin to antibodies have been described (see, see Nolting, Chapter 5 "Linker Technology in Antibody-Drug Conjugates," In: *Antibody-Drug Conjugates: Methods in Molecular Biology*, vol. 1045, pp. 71-100, Laurent Ducry (Ed.), Springer Science & Business Media, LLC, 2013; Jeffrey *et al.*, 2006, *Bioconjug. Chem.* 17:831-840; Jeffrey *et al.*, 2007, *Bioorg. Med. Chem. Lett.* 17:2278-2280; and Jiang *et al.*, 2005, *J. Am. Chem. Soc.* 127:11254-11255, each of which is incorporated herein by reference). All of these β -glucuronic acid-based linkers may be used in the anti-glyco-cMET ADCs of the disclosure.

[0207] Additionally, cytotoxic and/or cytostatic agents containing a phenol group can be covalently bonded to a linker through the phenolic oxygen. One such linker, described in WO

2007/089149, relies on a methodology in which a diamino-ethane "SpaceLink" is used in conjunction with traditional "PABO"-based self-immolative groups to deliver phenols. The cleavage of the linker is depicted schematically below, where D represents a cytotoxic and/or cytostatic agent having a phenolic hydroxyl group.

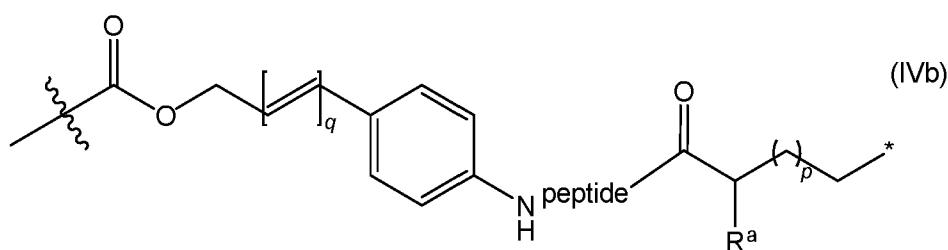
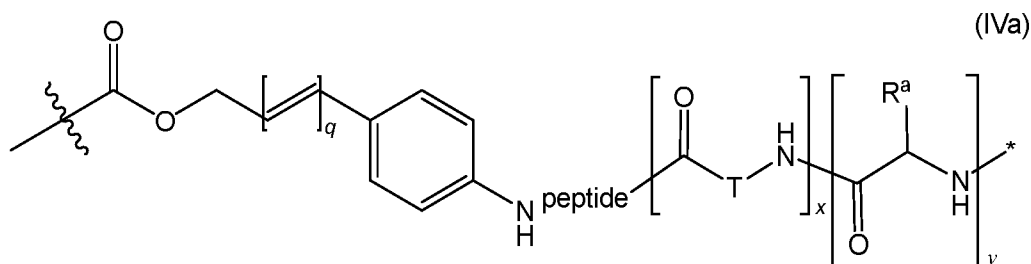


[0208] Cleavable linkers may include noncleavable portions or segments, and/or cleavable segments or portions may be included in an otherwise non-cleavable linker to render it cleavable. By way of example only, polyethylene glycol (PEG) and related polymers may include cleavable groups in the polymer backbone. For example, a polyethylene glycol or polymer linker may include one or more cleavable groups such as a disulfide, a hydrazone or a dipeptide.

[0209] Other degradable linkages that may be included in linkers include ester linkages formed by the reaction of PEG carboxylic acids or activated PEG carboxylic acids with alcohol groups on a biologically active agent, wherein such ester groups generally hydrolyze under physiological conditions to release the biologically active agent. Hydrolytically degradable linkages include, but are not limited to, carbonate linkages; imine linkages resulting from reaction of an amine and an aldehyde; phosphate ester linkages formed by reacting an alcohol with a phosphate group; acetal linkages that are the reaction product of an aldehyde and an

alcohol; orthoester linkages that are the reaction product of a formate and an alcohol; and oligonucleotide linkages formed by a phosphoramidite group, including but not limited to, at the end of a polymer, and a 5' hydroxyl group of an oligonucleotide.

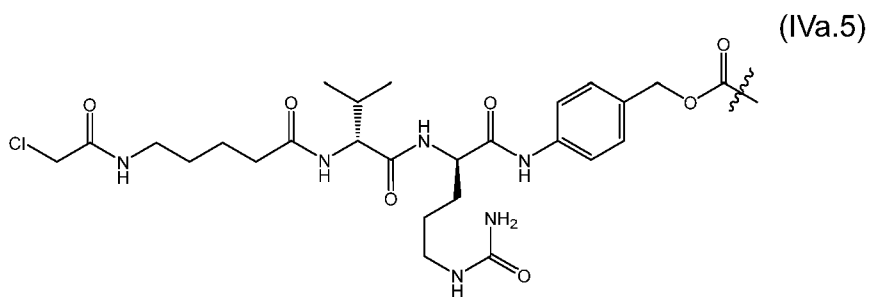
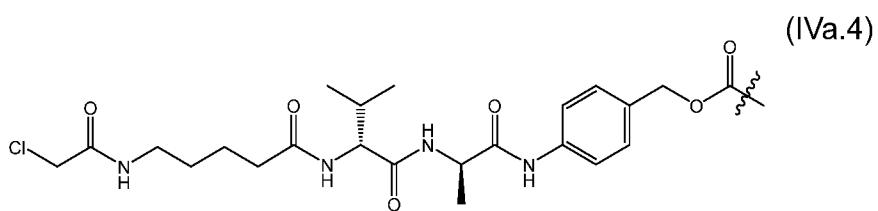
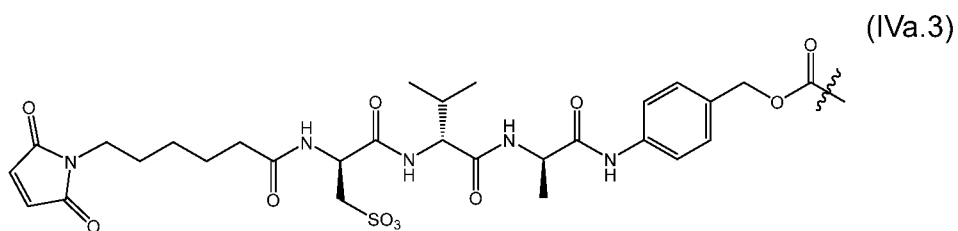
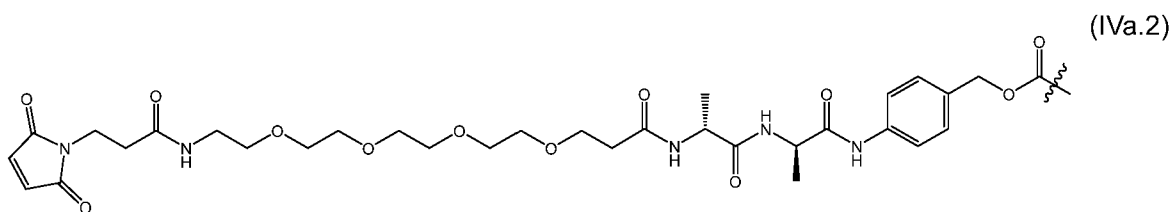
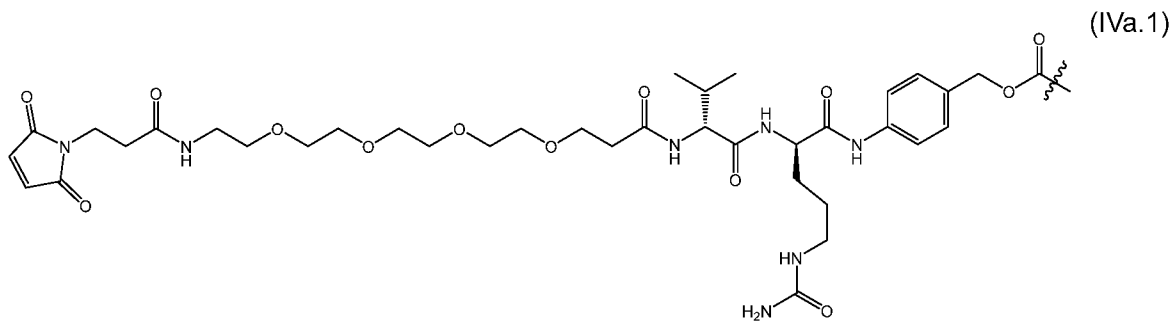
[0210] In certain embodiments, the linker comprises an enzymatically cleavable peptide moiety, for example, a linker comprising structural formula (IVa) or (IVb):

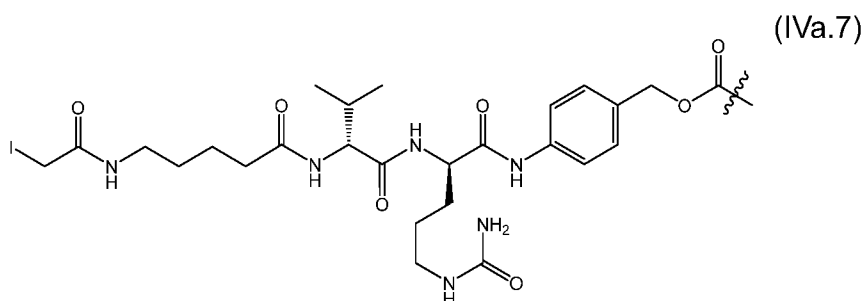
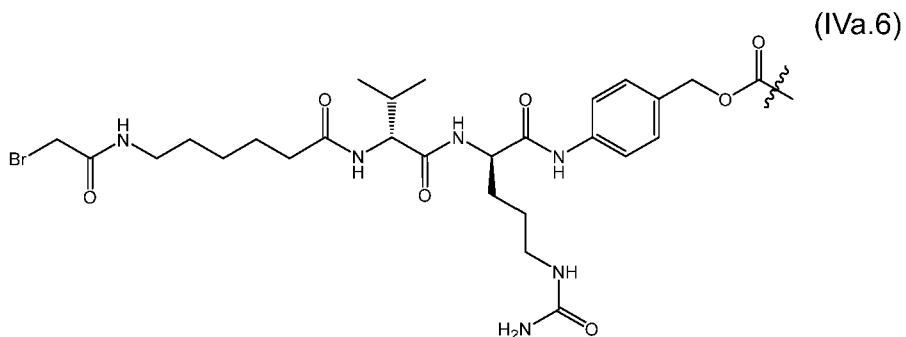


or a salt thereof, wherein: peptide represents a peptide (illustrated C→N and not showing the carboxy and amino “termini”) cleavable by a lysosomal enzyme; T represents a polymer comprising one or more ethylene glycol units or an alkylene chain, or combinations thereof; R^a is selected from hydrogen, alkyl, sulfonate and methyl sulfonate; p is an integer ranging from 0 to 5; q is 0 or 1; x is 0 or 1; y is 0 or 1; represents the point of attachment of the linker to a cytotoxic and/or cytostatic agent; and * represents the point of attachment to the remainder of the linker.

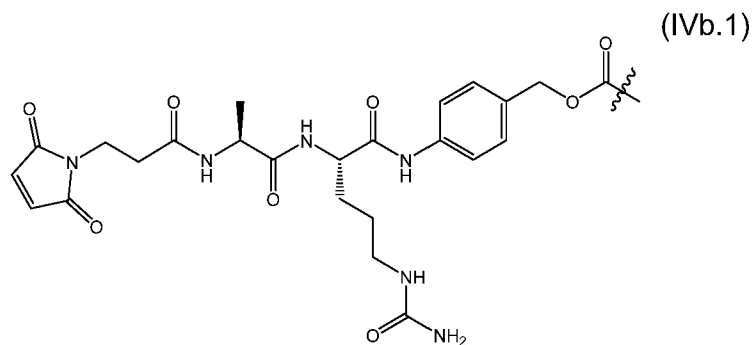
[0211] In certain embodiments, the peptide is selected from a tripeptide or a dipeptide. In particular embodiments, the dipeptide is selected from: Val-Cit; Cit-Val; Ala-Ala; Ala-Cit; Cit-Ala; Asn-Cit; Cit-Asn; Cit-Cit; Val-Glu; Glu-Val; Ser-Cit; Cit-Ser; Lys-Cit; Cit-Lys; Asp-Cit; Cit-Asp; Ala-Val; Val-Ala; Phe-Lys; Val-Lys; Ala-Lys; Phe-Cit; Leu-Cit; Ile-Cit; Phe-Arg; and Trp-Cit. In certain embodiments, the dipeptide is selected from: Cit-Val; and Ala-Val.

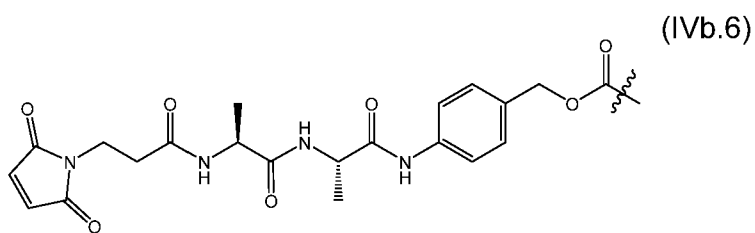
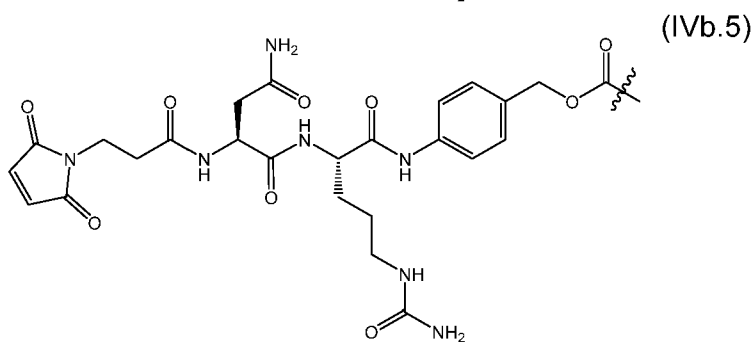
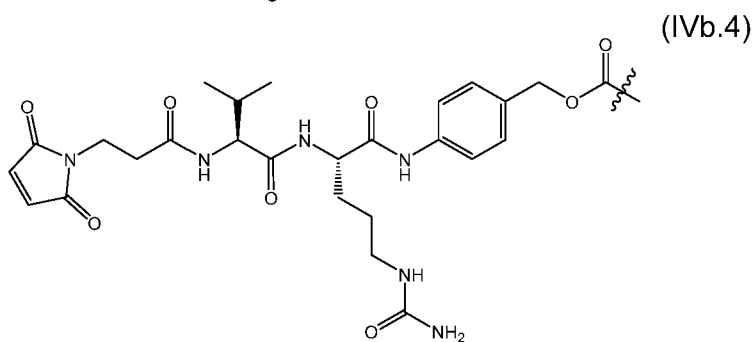
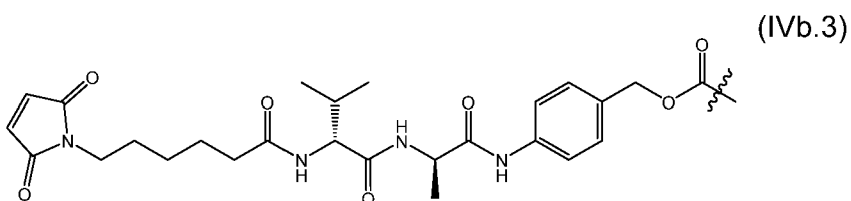
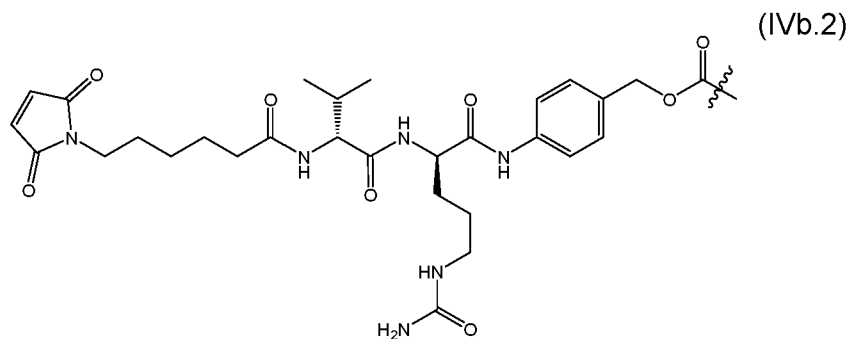
[0212] Specific exemplary embodiments of linkers according to structural formula (IVa) that may be included in the anti-glyco-cMET ADCs of the disclosure include the linkers illustrated below (as illustrated, the linkers include a group suitable for covalently linking the linker to an antibody):

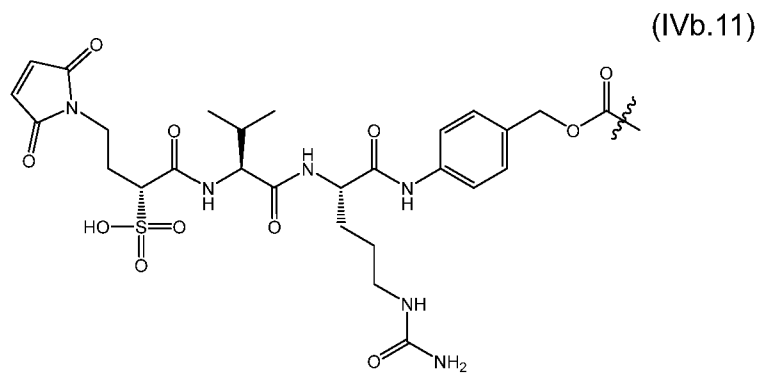
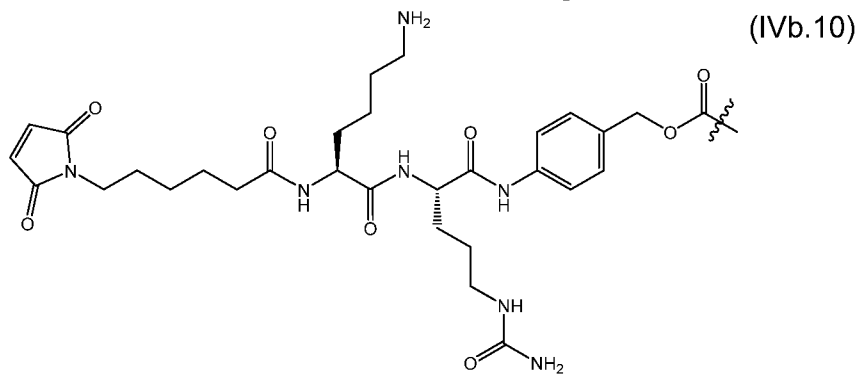
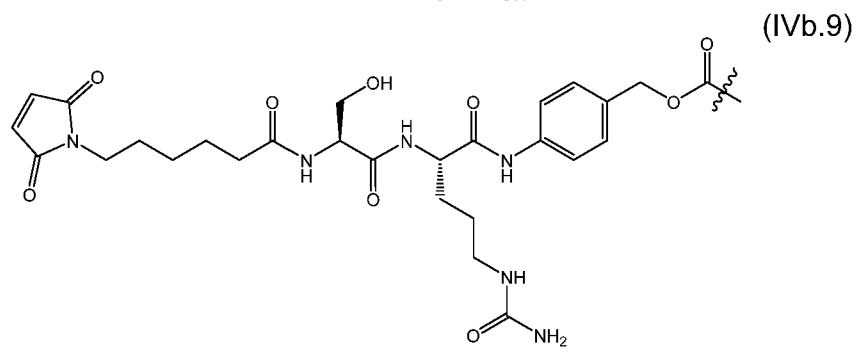
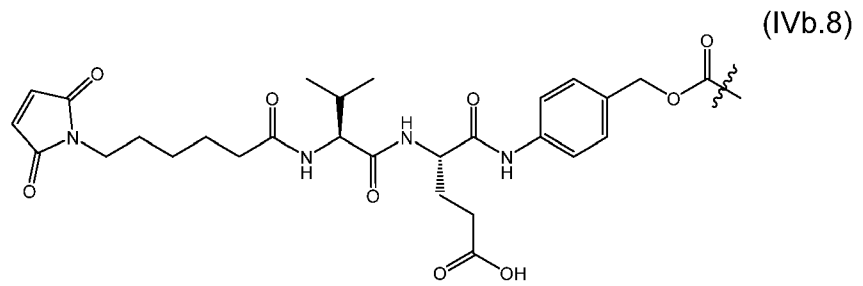
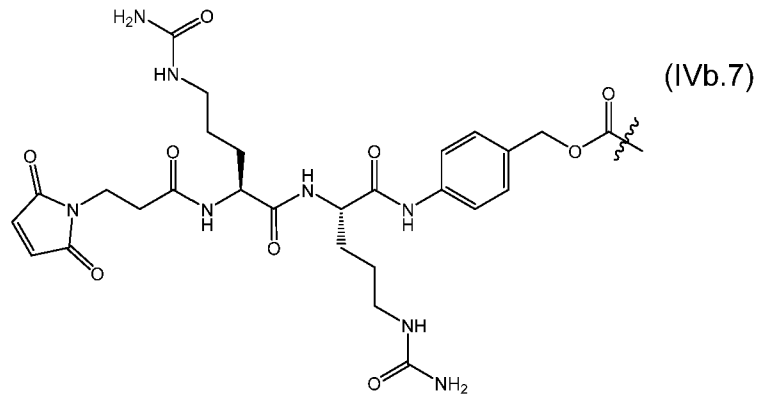




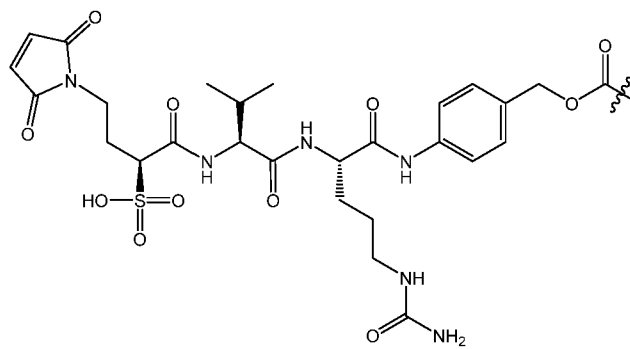
[0213] Specific exemplary embodiments of linkers according to structural formula (IVb) that may be included in the anti-glyco-cMET ADCs of the disclosure include the linkers illustrated below (as illustrated, the linkers include a group suitable for covalently linking the linker to an antibody):



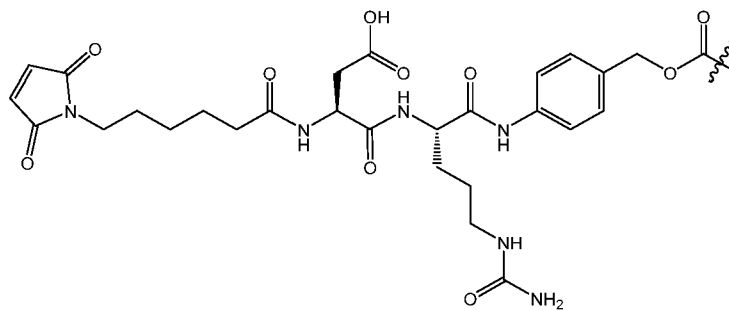




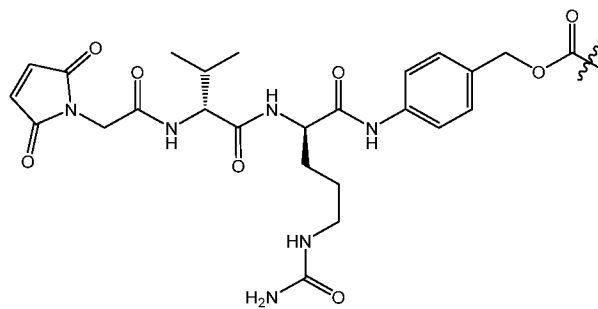
(IVb.12)



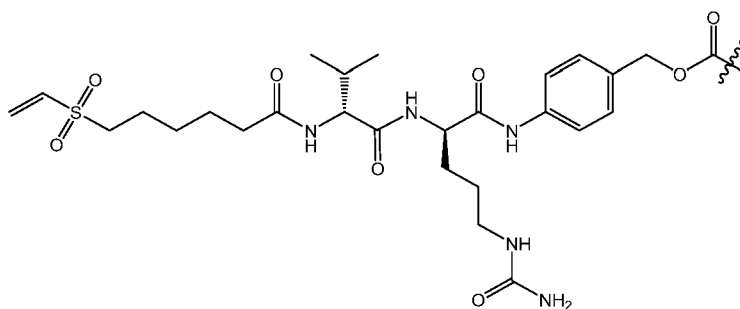
(IVb.13)



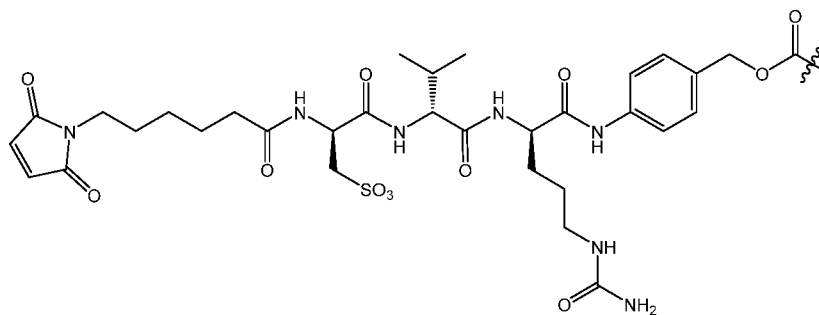
(IVb.14)



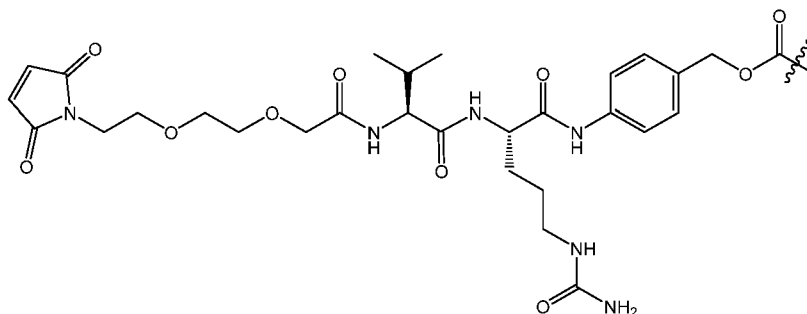
(IVb.15)



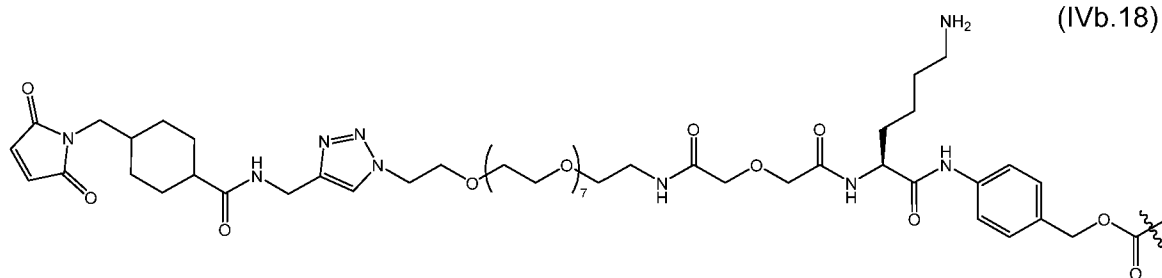
(IVb.16)



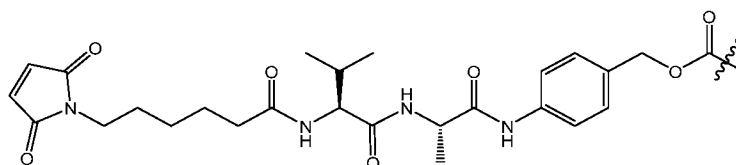
(IVb.17)



(IVb.18)

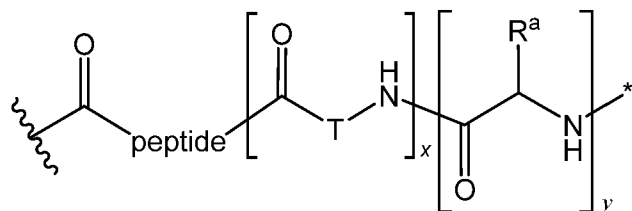


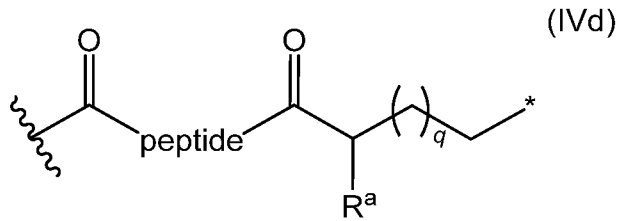
(IVb.19)



[0214] In certain embodiments, the linker comprises an enzymatically cleavable peptide moiety, for example, a linker comprising structural formula (IVc) or (IVd):

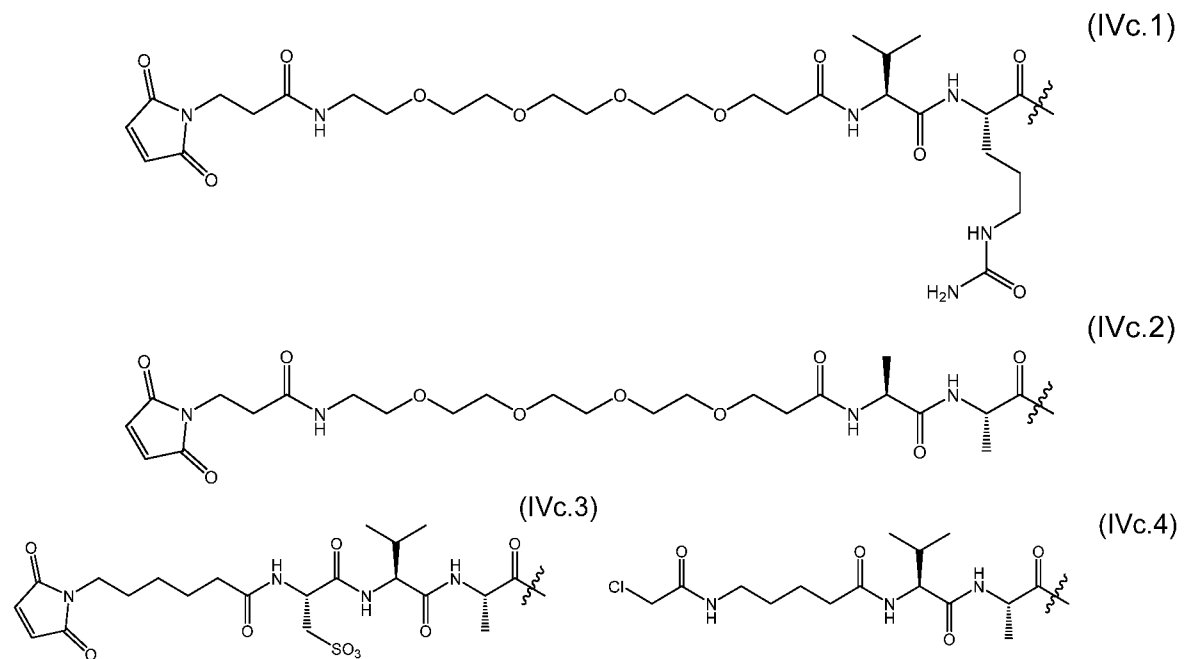
(IVc)

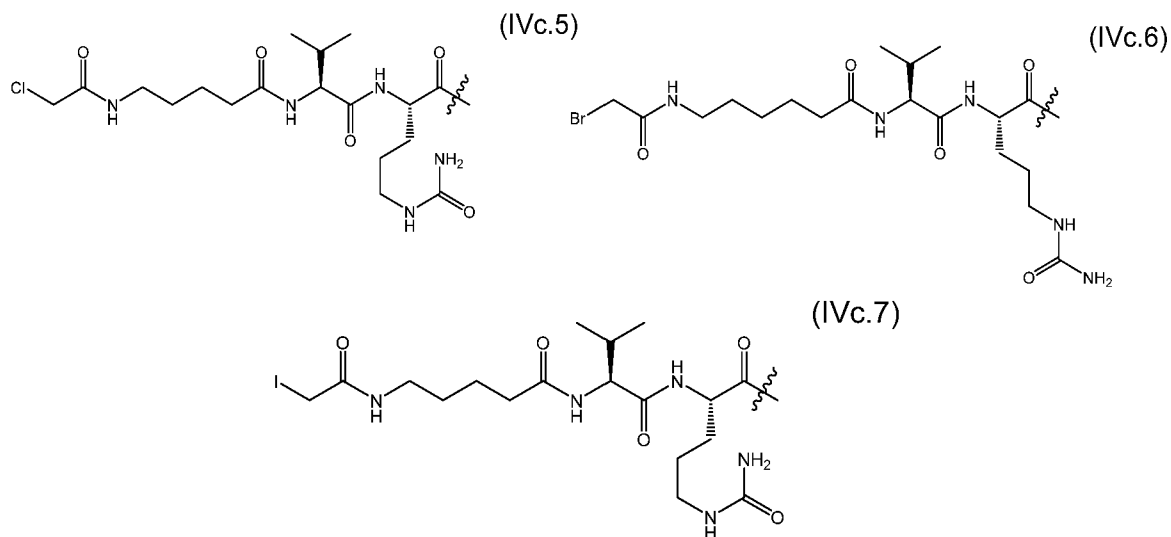




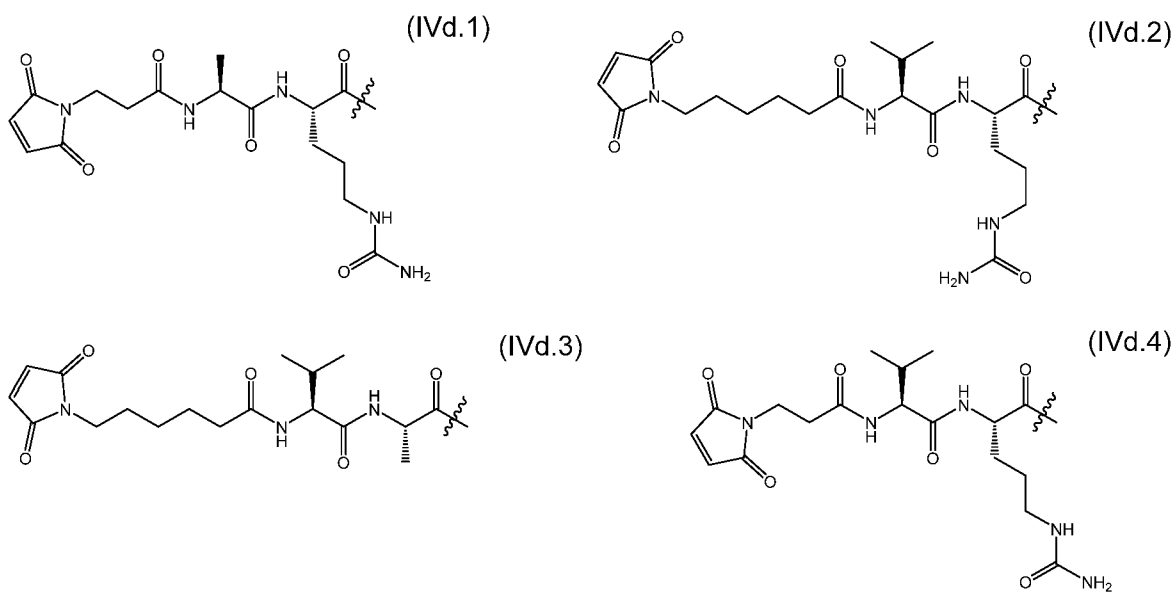
or a salt thereof, wherein: peptide represents a peptide (illustrated C→N and not showing the carboxy and amino “termini”) cleavable by a lysosomal enzyme; T represents a polymer comprising one or more ethylene glycol units or an alkylene chain, or combinations thereof; R^a is selected from hydrogen, alkyl, sulfonate and methyl sulfonate; p is an integer ranging from 0 to 5; q is 0 or 1; x is 0 or 1; y is 0 or 1; x^{linker} represents the point of attachment of the linker to a cytotoxic and/or cytostatic agent; and * represents the point of attachment to the remainder of the linker.

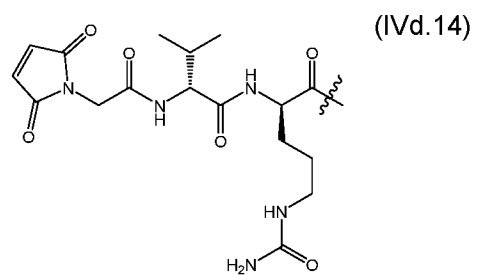
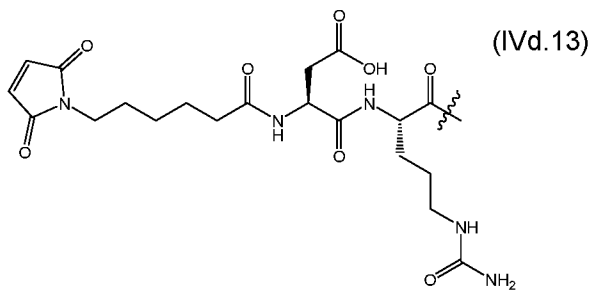
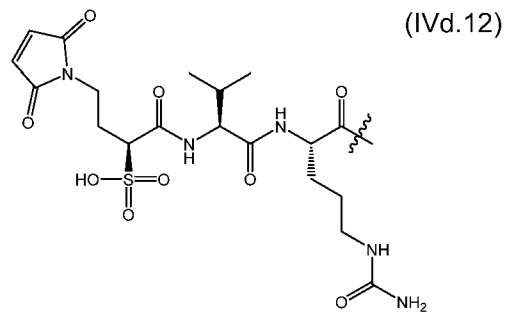
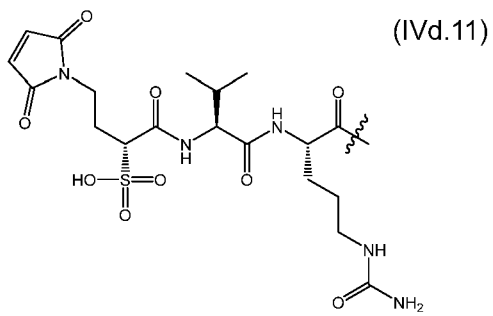
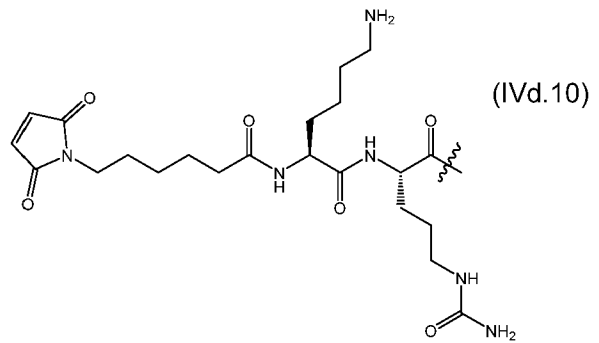
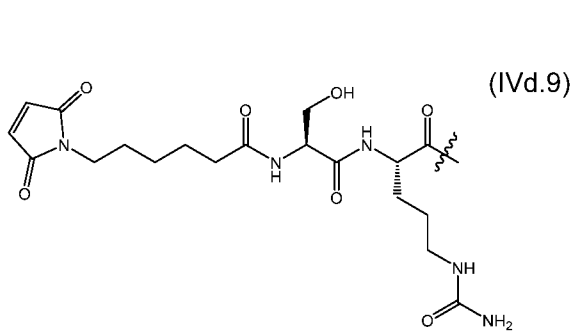
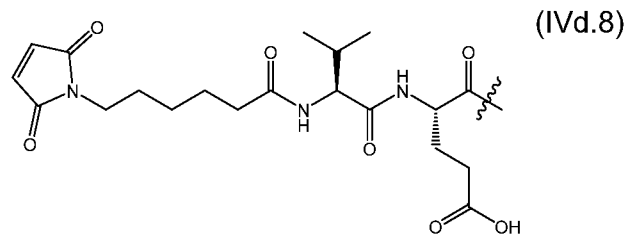
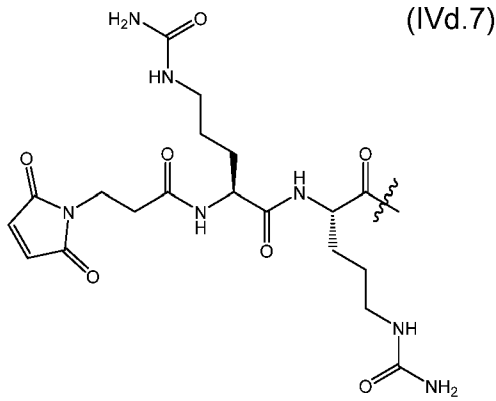
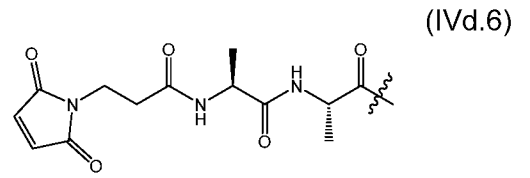
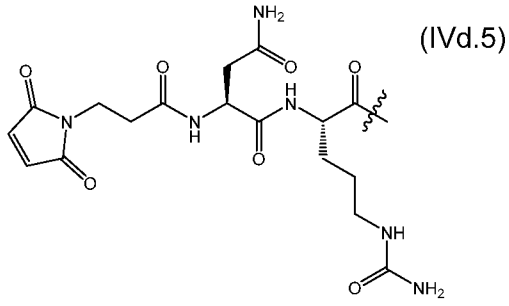
[0215] Specific exemplary embodiments of linkers according to structural formula (IVc) that may be included in the anti-glyco-cMET ADCs of the disclosure include the linkers illustrated below (as illustrated, the linkers include a group suitable for covalently linking the linker to an antibody):

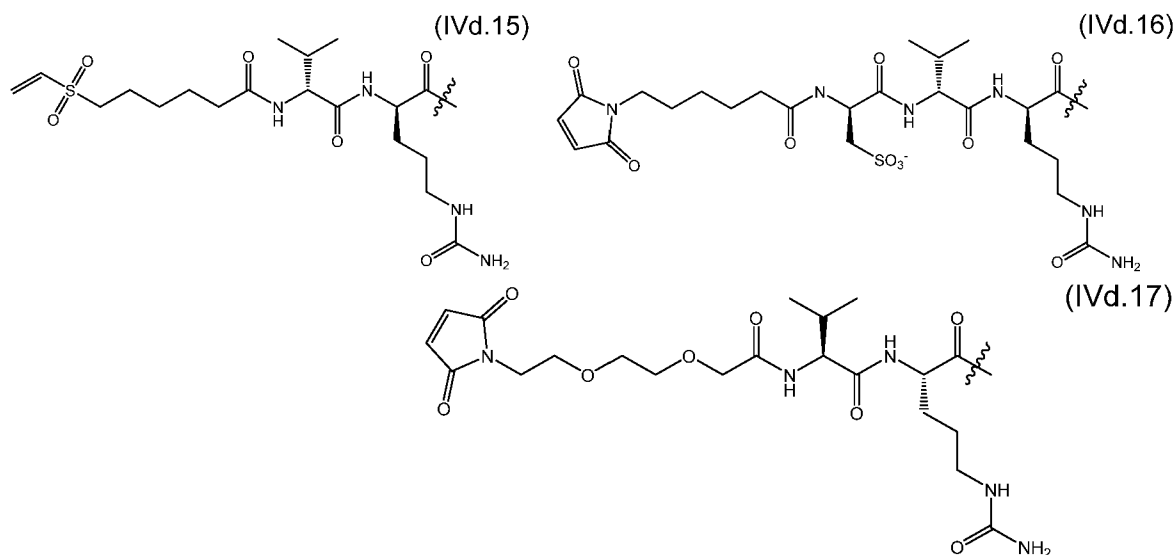




[0216] Specific exemplary embodiments of linkers according to structural formula (IVd) that may be included in the anti-glyco-cMET ADCs of the disclosure include the linkers illustrated below (as illustrated, the linkers include a group suitable for covalently linking the linker to an antibody):







[0217] In certain embodiments, the linker comprising structural formula (IVa), (IVb), (IVc), or (IVd) further comprises a carbonate moiety cleavable by exposure to an acidic medium. In particular embodiments, the linker is attached through an oxygen to a cytotoxic and/or cytostatic agent.

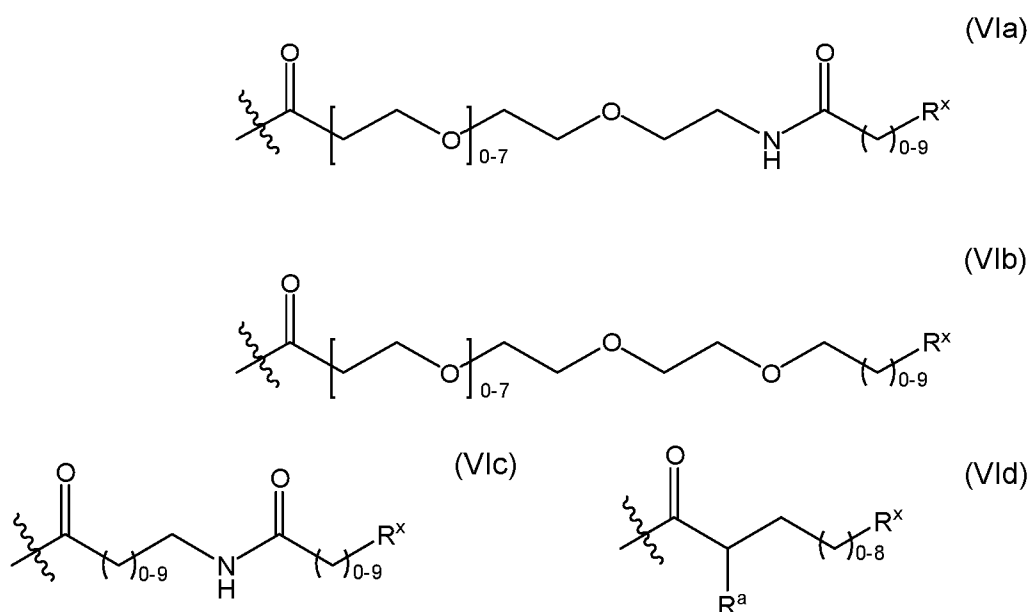
5.2.4. Non-Cleavable Linkers

[0218] Although cleavable linkers may provide certain advantages, the linkers comprising the anti-glyco-cMET ADC of the disclosure need not be cleavable. For noncleavable linkers, the release of drug does not depend on the differential properties between the plasma and some cytoplasmic compartments. The release of the drug is postulated to occur after internalization of the ADC via antigen-mediated endocytosis and delivery to lysosomal compartment, where the antibody is degraded to the level of amino acids through intracellular proteolytic degradation. This process releases a drug derivative, which is formed by the drug, the linker, and the amino acid residue to which the linker was covalently attached. The amino acid drug metabolites from conjugates with noncleavable linkers are more hydrophilic and generally less membrane permeable, which leads to less bystander effects and less nonspecific toxicities compared to conjugates with a cleavable linker. In general, ADCs with noncleavable linkers have greater stability in circulation than ADCs with cleavable linkers. Non-cleavable linkers may be alkylene chains, or maybe polymeric in nature, such as, for example, based upon polyalkylene glycol polymers, amide polymers, or may include segments of alkylene chains, polyalkylene glycols and/or amide polymers.

[0219] A variety of non-cleavable linkers used to link drugs to antibodies have been described. See, Jeffrey *et al.*, 2006, *Bioconjug. Chem.* 17; 831-840; Jeffrey *et al.*, 2007, *Bioorg. Med. Chem. Lett.* 17:2278-2280; and Jiang *et al.*, 2005, *J. Am. Chem. Soc.* 127:11254-11255, each

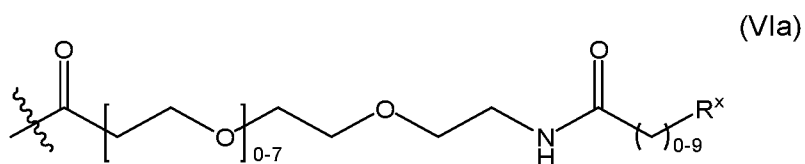
of which is incorporated herein by reference. All of these linkers may be included in the anti-glyco-cMET ADCs of the disclosure.

[0220] In certain embodiments, the linker is non-cleavable *in vivo*, for example a linker according to structural formula (VIa), (VIb), (VIc) or (VIId) (as illustrated, the linkers include a group suitable for covalently linking the linker to an antibody:

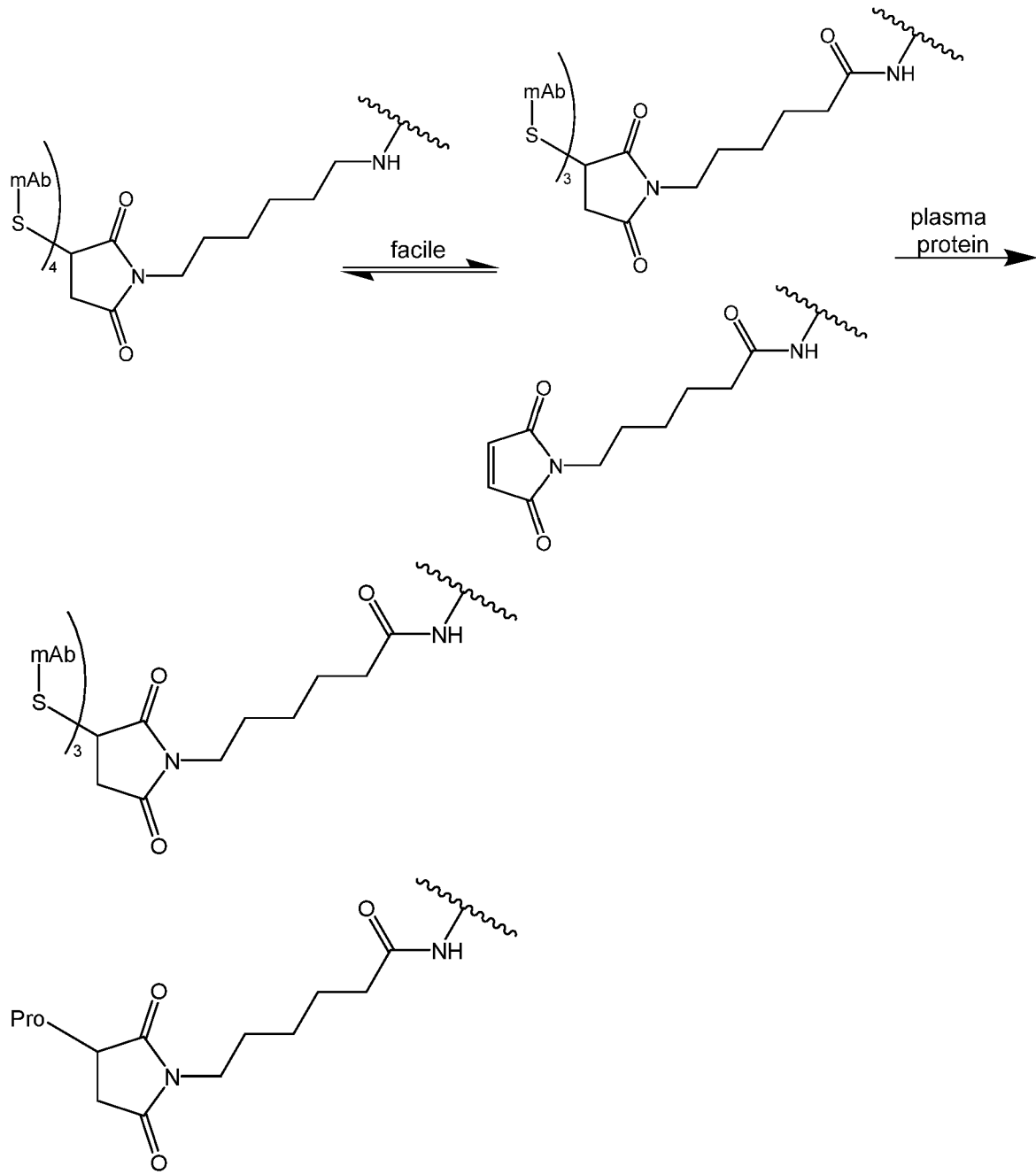


or salts thereof, wherein: R^a is selected from hydrogen, alkyl, sulfonate and methyl sulfonate; R^x is a moiety including a functional group capable of covalently linking the linker to an antibody; and --- represents the point of attachment of the linker to a cytotoxic and/or cytostatic agent.

[0221] Specific exemplary embodiments of linkers according to structural formula (VIa)-(VIId) that may be included in the anti-glyco-cMET ADCs of the disclosure include the linkers illustrated below (as illustrated, the linkers include a group suitable for covalently linking the linker to an antibody, and --- represents the point of attachment to a cytotoxic and/or cytostatic agent):

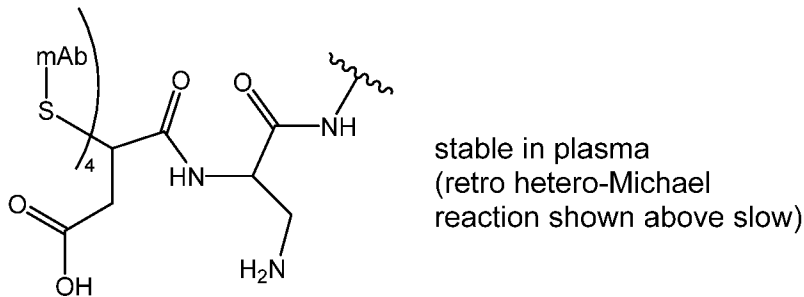
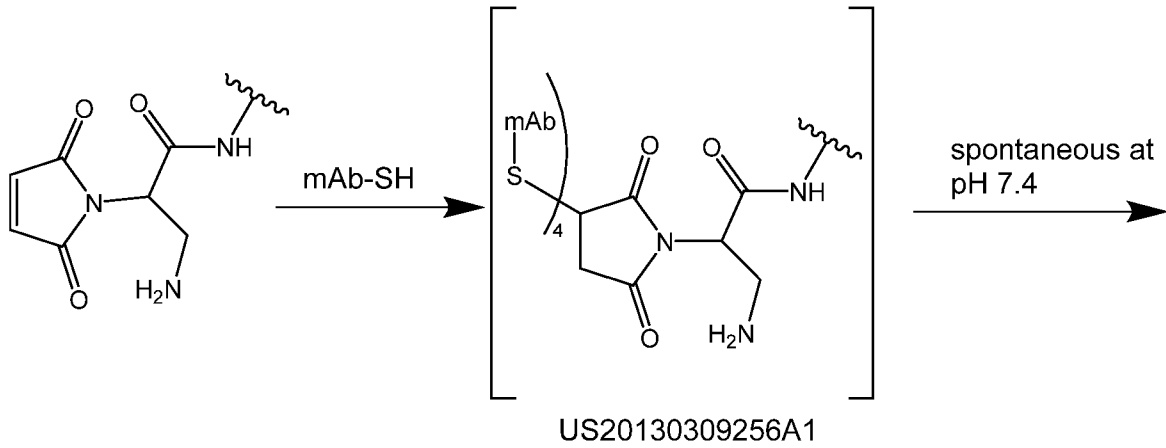


Normal system:

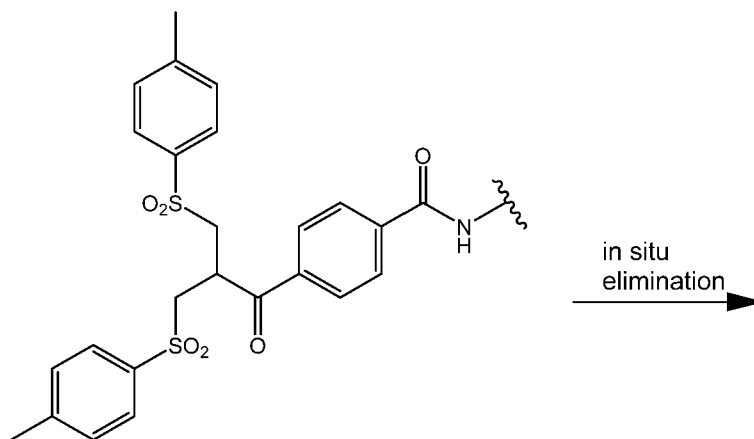


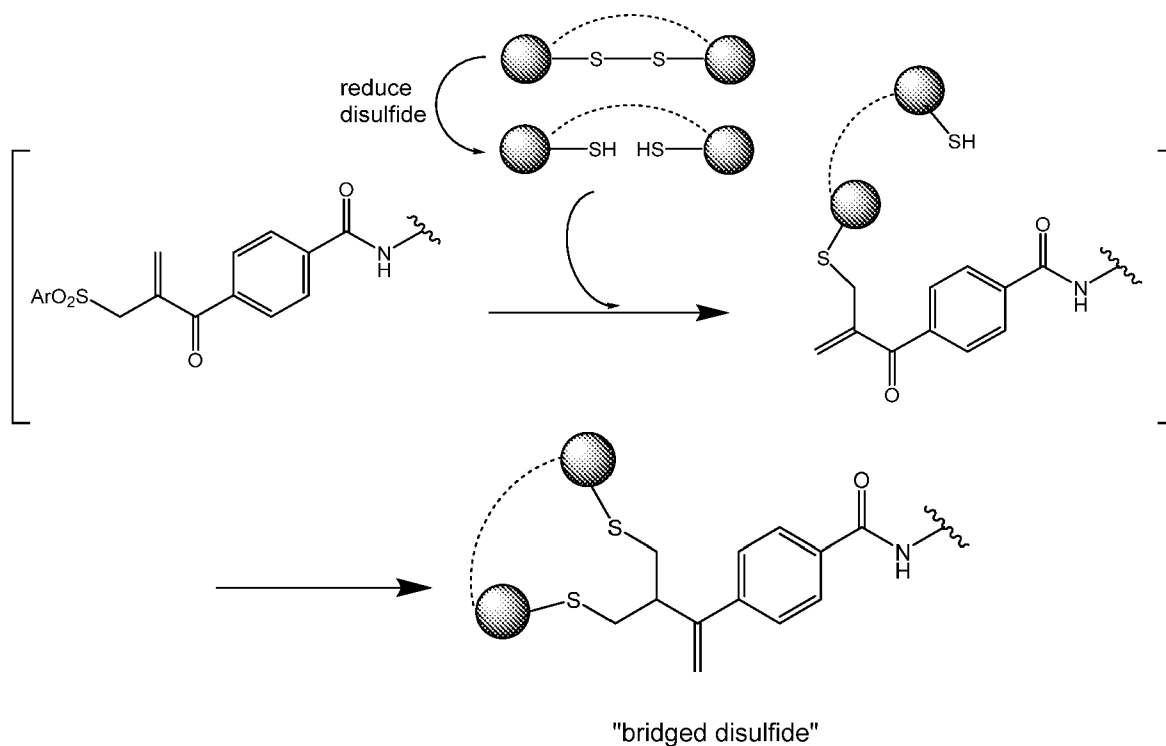
Leads to "DAR loss" over time

SGN MalDPR (maleimido dipropylamino) system:

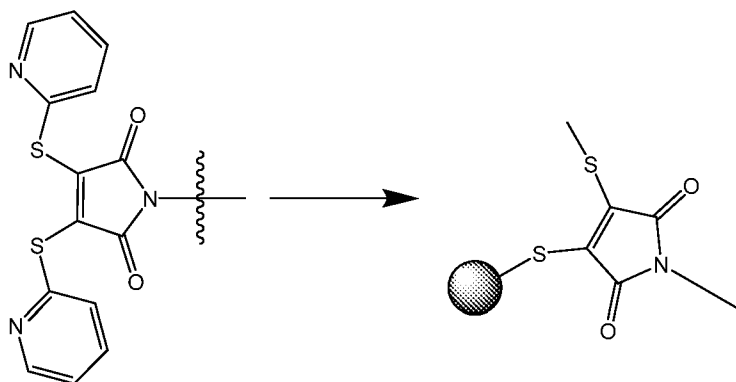


[0224] Polytherics has disclosed a method for bridging a pair of sulfhydryl groups derived from reduction of a native hinge disulfide bond. See, Badescu *et al.*, 2014, *Bioconjugate Chem.* 25:1124-1136. The reaction is depicted in the schematic below. An advantage of this methodology is the ability to synthesize enriched DAR4 ADCs by full reduction of IgGs (to give 4 pairs of sulfhydryls) followed by reaction with 4 equivalents of the alkylating agent. ADCs containing "bridged disulfides" are also claimed to have increased stability.





[0225] Similarly, as depicted below, a maleimide derivative (1, below) that is capable of bridging a pair of sulfhydryl groups has been developed. See WO2013/085925.



5.2.6. Linker Selection Considerations

[0226] As is known by skilled artisans, the linker selected for a particular ADC may be influenced by a variety of factors, including but not limited to, the site of attachment to the antibody (e.g., lys, cys or other amino acid residues), structural constraints of the drug pharmacophore and the lipophilicity of the drug. The specific linker selected for an ADC should seek to balance these different factors for the specific antibody/drug combination. For a review of the factors that are influenced by choice of linkers in ADCs, see Nolting, Chapter 5 "Linker Technology in Antibody-Drug Conjugates," In: *Antibody-Drug Conjugates: Methods in Molecular*

Biology, vol. 1045, pp. 71-100, Laurent Ducry (Ed.), Springer Science & Business Medica, LLC, 2013.

[0227] For example, ADCs have been observed to effect killing of bystander antigen-negative cells present in the vicinity of the antigen-positive tumor cells. The mechanism of bystander cell killing by ADCs has indicated that metabolic products formed during intracellular processing of the ADCs may play a role. Neutral cytotoxic metabolites generated by metabolism of the ADCs in antigen-positive cells appear to play a role in bystander cell killing while charged metabolites may be prevented from diffusing across the membrane into the medium and therefore cannot affect bystander killing. In certain embodiments, the linker is selected to attenuate the bystander killing effect caused by cellular metabolites of the ADC. In certain embodiments, the linker is selected to increase the bystander killing effect.

[0228] The properties of the linker may also impact aggregation of the ADC under conditions of use and/or storage. Typically, ADCs reported in the literature contain no more than 3-4 drug molecules per antibody molecule (see, e.g., Chari, 2008, *Acc Chem Res* 41:98-107). Attempts to obtain higher drug-to-antibody ratios ("DAR") often failed, particularly if both the drug and the linker were hydrophobic, due to aggregation of the ADC (King *et al.*, 2002, *J Med Chem* 45:4336-4343; Hollander *et al.*, 2008, *Bioconjugate Chem* 19:358-361; Burke *et al.*, 2009 *Bioconjugate Chem* 20:1242-1250). In many instances, DARs higher than 3-4 could be beneficial as a means of increasing potency. In instances where the cytotoxic and/or cytostatic agent is hydrophobic in nature, it may be desirable to select linkers that are relatively hydrophilic as a means of reducing ADC aggregation, especially in instances where DARS greater than 3-4 are desired. Thus, in certain embodiments, the linker incorporates chemical moieties that reduce aggregation of the ADCs during storage and/or use. A linker may incorporate polar or hydrophilic groups such as charged groups or groups that become charged under physiological pH to reduce the aggregation of the ADCs. For example, a linker may incorporate charged groups such as salts or groups that deprotonate, e.g., carboxylates, or protonate, e.g., amines, at physiological pH.

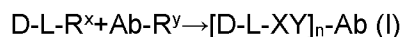
[0229] Exemplary polyvalent linkers that have been reported to yield DARs as high as 20 that may be used to link numerous cytotoxic and/or cytostatic agents to an antibody are described in WO 2009/073445; WO 2010/068795; WO 2010/138719; WO 2011/120053; WO 2011/171020; WO 2013/096901; WO 2014/008375; WO 2014/093379; WO 2014/093394; WO 2014/093640, the content of which are incorporated herein by reference in their entireties.

[0230] In particular embodiments, the aggregation of the ADCs during storage or use is less than about 10% as determined by size-exclusion chromatography (SEC). In particular embodiments, the aggregation of the ADCs during storage or use is less than 10%, such as less than about 5%, less than about 4%, less than about 3%, less than about 2%, less than

about 1%, less than about 0.5%, less than about 0.1%, or even lower, as determined by size-exclusion chromatography (SEC).

5.2.7. Methods of Making Anti-Glyco-cMET ADCs

[0231] The anti-glyco-cMET ADCs of the disclosure may be synthesized using chemistries that are well-known. The chemistries selected will depend upon, among other things, the identity of the cytotoxic and/or cytostatic agent(s), the linker and the groups used to attach linker to the antibody. Generally, ADCs according to formula (I) may be prepared according to the following scheme:



[0232] where D, L, Ab, XY and n are as previously defined, and R^x and R^y represent complementary groups capable of forming a covalent linkages with one another, as discussed above.

[0233] The identities of groups R^x and R^y will depend upon the chemistry used to link synthon D-L- R^x to the antibody. Generally, the chemistry used should not alter the integrity of the antibody, for example its ability to bind its target. Preferably, the binding properties of the conjugated antibody will closely resemble those of the unconjugated antibody. A variety of chemistries and techniques for conjugating molecules to biological molecules such as antibodies are known in the art and in particular to antibodies, are well-known. See, e.g., Amon *et al.*, "Monoclonal Antibodies For Immunotargeting Of Drugs In Cancer Therapy," in: *Monoclonal Antibodies And Cancer Therapy*, Reisfeld *et al.* Eds., Alan R. Liss, Inc., 1985; Hellstrom *et al.*, "Antibodies For Drug Delivery," in: *Controlled Drug Delivery*, Robinson *et al.* Eds., Marcel Dekker, Inc., 2nd Ed. 1987; Thorpe, "Antibody Carriers Of Cytotoxic Agents In Cancer Therapy: A Review," in: *Monoclonal Antibodies '84: Biological And Clinical Applications*, Pinchera *et al.*, Eds., 1985; "Analysis, Results, and Future Prospective of the Therapeutic Use of Radiolabeled Antibody In Cancer Therapy," in: *Monoclonal Antibodies For Cancer Detection And Therapy*, Baldwin *et al.*, Eds., Academic Press, 1985; Thorpe *et al.*, 1982, *Immunol. Rev.* 62:119-58; PCT publication WO 89/12624. Any of these chemistries may be used to link the synthons to an antibody.

[0234] A number of functional groups R^x and chemistries useful for linking synthons to accessible lysine residues are known and include, by way of example and not limitation, NHS-esters and isothiocyanates.

[0235] A number of functional groups R^x and chemistries useful for linking synthons to accessible free sulfhydryl groups of cysteine residues are known and include, by way of example and not limitation, haloacetyls and maleimides.

[0236] However, conjugation chemistries are not limited to available side chain groups. Side chains such as amines may be converted to other useful groups, such as hydroxyls, by linking

an appropriate small molecule to the amine. This strategy can be used to increase the number of available linking sites on the antibody by conjugating multifunctional small molecules to side chains of accessible amino acid residues of the antibody. Functional groups R^x suitable for covalently linking the synthons to these "converted" functional groups are then included in the synthons.

[0237] The antibody may also be engineered to include amino acid residues for conjugation. An approach for engineering antibodies to include non-genetically encoded amino acid residues useful for conjugating drugs in the context of ADCs is described by Axup *et al.*, 2012, Proc Natl Acad Sci USA. 109(40):16101-16106, as are chemistries and functional group useful for linking synthons to the non-encoded amino acids.

[0238] Typically, the synthons are linked to the side chains of amino acid residues of the antibody, including, for example, the primary amino group of accessible lysine residues or the sulfhydryl group of accessible cysteine residues. Free sulfhydryl groups may be obtained by reducing interchain disulfide bonds.

[0239] For linkages where R^y is a sulfhydryl group (for example, when R^x is a maleimide), the antibody is generally first fully or partially reduced to disrupt interchain disulfide bridges between cysteine residues.

[0240] Cysteine residues that do not participate in disulfide bridges may be engineered into an antibody by mutation of one or more codons. Reducing these unpaired cysteines yields a sulfhydryl group suitable for conjugation. Preferred positions for incorporating engineered cysteines include, by way of example and not limitation, positions S112C, S113C, A114C, S115C, A176C, 5180C, S252C, V286C, V292C, S357C, A359C, S398C, S428C (Kabat numbering) on the human IgG₁ heavy chain and positions V110C, S114C, S121C, S127C, S168C, V205C (Kabat numbering) on the human Ig kappa light chain (see, e.g., U.S. Pat. No. 7,521,541, U.S. Pat. No. 7,855,275 and U.S. Pat. No. 8,455,622).

[0241] As will be appreciated by skilled artisans, the number of cytotoxic and/or cytostatic agents linked to an antibody molecule may vary, such that a collection of ADCs may be heterogeneous in nature, where some antibodies contain one linked agent, some two, some three, *etc.* (and some none). The degree of heterogeneity will depend upon, among other things, the chemistries used for linking the cytotoxic and/or cytostatic agents. For example, where the antibodies are reduced to yield sulfhydryl groups for attachment, heterogeneous mixtures of antibodies having zero, 2, 4, 6 or 8 linked agents per molecule are often produced. Furthermore, by limiting the molar ratio of attachment compound, antibodies having zero, 1, 2, 3, 4, 5, 6, 7 or 8 linked agents per molecule are often produced. Thus, it will be understood that depending upon context, stated DARs may be averages for a collection of antibodies. For example, "DAR4" can refer to an ADC preparation that has not been subjected to purification to isolate specific DAR peaks and can comprise a heterogeneous mixture of ADC molecules

having different numbers of cytostatic and/or cytotoxic agents attached per antibody (e.g., 0, 2, 4, 6, 8 agents per antibody), but has an average drug-to-antibody ratio of 4. Similarly, in some embodiments, "DAR2" refers to a heterogeneous ADC preparation in which the average drug-to-antibody ratio is 2.

[0242] When enriched preparations are desired, antibodies having defined numbers of linked cytotoxic and/or cytostatic agents may be obtained via purification of heterogeneous mixtures, for example, via column chromatography, e.g., hydrophobic interaction chromatography.

[0243] Purity may be assessed by a variety of methods, as is known in the art. As a specific example, an ADC preparation may be analyzed via HPLC or other chromatography and the purity assessed by analyzing areas under the curves of the resultant peaks.

5.3 Chimeric Antigen Receptors

[0244] The present disclosure provides chimeric antigen receptors (CARs) comprising the anti-glyco-cMET antibodies or antigen-binding fragments described herein. In some embodiments, the CAR comprises one or more scFvs (e.g., one or two) as described herein. For example, a CAR can comprise two scFvs covalently connected by a linker sequence (e.g., of 4-15 amino acids). Exemplary linkers include GGGGS (SEQ ID NO:293) and (GGGGS)₃ (SEQ ID NO:346).

[0245] The CARs of the disclosure typically comprise an extracellular domain operably linked to a transmembrane domain which is in turn operably linked to an intracellular domain for signaling. The CARs can further comprise a signal peptide at the N-terminus of the extracellular domain (e.g., a human CD8 signal peptide). In some embodiments, a CAR of the disclosure comprises a human CD8 signal peptide comprising the amino acid sequence MALPVTALLLPLALLLHAARP (SEQ ID NO:294).

[0246] The extracellular domains of the CARs of the disclosure comprise the sequence of an anti-glyco-cMET antibody or antigen-binding fragment (e.g., as described in Section 5.1 or numbered embodiments 689 to 724).

[0247] Exemplary transmembrane domain sequence and intracellular domain sequences are described in Section 5.3.1 and 5.3.2, respectively.

[0248] Several fusion proteins described herein (e.g., numbered embodiments 664 to 688) are CARs, and the CAR-related disclosures (e.g., numbered embodiments 689 to 724) apply to such fusion proteins. Other fusion proteins described herein (e.g., in numbered embodiments 735 to 834) are chimeric T cell receptors, and the chimeric TCR-related disclosures apply to such fusion proteins.

5.3.1. Transmembrane Domain

[0249] With respect to the transmembrane domain, the CAR can be designed to comprise a transmembrane domain that is operably linked (*e.g.*, fused) to the extracellular domain of the CAR.

[0250] The transmembrane domain may be derived either from a natural or from a synthetic source. Where the source is natural, the domain may be derived from any membrane-bound or transmembrane protein. Transmembrane regions of particular use in this disclosure may be derived from (*i.e.*, comprise at least the transmembrane region(s) of) the alpha, beta or zeta chain of the T-cell receptor, CD28, CD3 epsilon, CD45, CD4, CD5, CD8, CD9, CD16, CD22, CD33, CD37, CD64, CD80, CD86, CD134, CD137, CD154. In some instances, a variety of human hinges can be employed as well including the human Ig (immunoglobulin) hinge.

[0251] In one embodiment, the transmembrane domain is synthetic (*i.e.*, non-naturally occurring). Examples of synthetic transmembrane domains are peptides comprising predominantly hydrophobic residues such as leucine and valine. Preferably a triplet of phenylalanine, tryptophan and valine will be found at each end of a synthetic transmembrane domain. Optionally, a short oligo- or polypeptide linker, preferably between 2 and 10 amino acids in length may form the linkage between the transmembrane domain and the cytoplasmic signaling domain of the CAR. A glycine-serine doublet provides a particularly suitable linker.

[0252] In one embodiment, the transmembrane domain in the CAR of the disclosure is the CD8 transmembrane domain. In one embodiment, the CD8 transmembrane domain comprises the amino acid sequence YLHLGALGRDLWGPSPVTGYHPLL (SEQ ID NO:295).

[0253] In one embodiment, the transmembrane domain in the CAR of the disclosure is the CD28 transmembrane domain. In one embodiment, the CD28 transmembrane domain comprises the amino acid sequence FWVLVWVGGVLACYSLLVTVAFIIFWV (SEQ ID NO:296).

[0254] In some instances, the transmembrane domain of the CAR of the disclosure is linked to the extracellular domain by a CD8a hinge domain. In one embodiment, the CD8a hinge domain comprises the amino acid sequence

TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFAC (SEQ ID NO:297). In another embodiment, the CD8a hinge domain comprises the amino acid sequence

TTTPAPRPPTPAPTIASPLSLRPEACRPAAGGAVHTRGLDFACD (SEQ ID NO:298). In another embodiment, the CD8a hinge domain comprises the amino acid sequence

TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFACD (SEQ ID NO:349).

[0255] In some instances, the transmembrane domain of the CAR of the disclosure is linked to the extracellular domain by a human IgG4-short hinge. In one embodiment, the human IgG4-short hinge comprises the amino acid sequence ESKYGPPCPSCP (SEQ ID NO:299).

[0256] In some instances, the transmembrane domain of the CAR of the disclosure is linked to the extracellular domain by a human IgG4-long hinge. In one embodiment, the human IgG4-long hinge comprises the amino acid sequence

ESKYGPPCPPCPAPPVAGPSVFLFPPKPKDTLMISRTPEVTCVVDVVSQEDPEVQFNWYVDG
VEVHNAKTKPREEQFQSTYRVVSVLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPR
EPQVYTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTTPVLDSDGSFFLY
SRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLSLGLGKM (SEQ ID NO:300).

5.3.2. Intracellular Domain

[0257] The intracellular signaling domain of the CAR of the disclosure is responsible for activation of at least one of the normal effector functions of the immune cell in which the CAR is expressed. The term “effector function” refers to a specialized function of a cell. Effector function of a T cell, for example, may be cytolytic activity or helper activity including the secretion of cytokines. Thus the term “intracellular signaling domain” refers to the portion of a protein which transduces the effector function signal and directs the cell to perform a specialized function. While usually the entire intracellular signaling domain can be employed, in many cases it is not necessary to use the entire chain. To the extent that a truncated portion of the intracellular signaling domain is used, such truncated portion may be used in place of the intact chain as long as it transduces the effector function signal. The term intracellular signaling domain is thus meant to include any truncated portion of the intracellular signaling domain sufficient to transduce the effector function signal.

[0258] Preferred examples of intracellular signaling domains for use in the CAR of the disclosure include the cytoplasmic sequences of the T cell receptor (TCR) and co-receptors that act in concert to initiate signal transduction following antigen receptor engagement, as well as any derivative or variant of these sequences and any synthetic sequence that has the same functional capability.

[0259] Signals generated through the TCR alone may be insufficient for full activation of the T cell and a secondary or co-stimulatory signal is also required. Thus, T cell activation can be said to be mediated by two distinct classes of cytoplasmic signaling sequence: those that initiate antigen-dependent primary activation through the TCR (primary cytoplasmic signaling sequences) and those that act in an antigen-independent manner to provide a secondary or co-stimulatory signal (secondary cytoplasmic signaling sequences).

[0260] Primary cytoplasmic signaling sequences regulate primary activation of the TCR complex either in a stimulatory way, or in an inhibitory way. Primary cytoplasmic signaling sequences that act in a stimulatory manner may contain signaling motifs which are known as immunoreceptor tyrosine-based activation motifs or ITAMs.

[0261] Examples of ITAM containing primary cytoplasmic signaling sequences that are of particular use in the CARs of the disclosure include those derived from TCR zeta, FcR gamma, FcR beta, CD3 gamma, CD3 delta, CD3 epsilon, CD5, CD22, CD79a, CD79b, and CD66d. It is particularly preferred that cytoplasmic signaling molecule in the CAR of the disclosure comprises a cytoplasmic signaling sequence from CD3-zeta.

[0262] In a preferred embodiment, the cytoplasmic domain of the CAR is designed to include an ITAM containing primary cytoplasmic signaling sequences domain (e.g., that of CD3-zeta) by itself or combined with any other desired cytoplasmic domain(s) useful in the context of the CAR of the disclosure. For example, the cytoplasmic domain of the CAR can include a CD3 zeta chain portion and a costimulatory signaling region.

[0263] The costimulatory signaling region refers to a portion of the CAR comprising the intracellular domain of a costimulatory molecule. A costimulatory molecule is a cell surface molecule other than an antigen receptor or its ligands that is required for an efficient response of lymphocytes to an antigen. Examples of such molecules include CD27, CD28, 4-1BB (CD137), OX40, CD30, CD40, PD-1, ICOS, lymphocyte function-associated antigen-1 (LFA-1), CD2, CD7, LIGHT, NKG2C, B7-H3, and a ligand that specifically binds with CD83, DAP10, GITR, and the like.

[0264] The cytoplasmic signaling sequences within the cytoplasmic signaling portion of the CAR of the disclosure may be linked to each other in a random or specified order. Optionally, a short oligo- or polypeptide linker, preferably between 2 and 10 amino acids in length may form the linkage. A glycine-serine doublet provides a particularly suitable linker.

[0265] In one embodiment, the cytoplasmic domain comprises the signaling domain of CD3-zeta and the signaling domain of CD28. In some embodiments, the signaling domain of CD3-zeta comprises the amino acid sequence
RVKFSRSADAPAYQQGQNQLYNELNLGRREEYDVLDKRRGRDPEMGGKPRRKNPQEGLYN
ELQKDKMAEAYSEIGMKGERRRGKGGHDGLYQGLSTATKDTYDALHMQALPPR (SEQ ID
NO:301). In some embodiments, the signaling domain of CD28 comprises the amino acid
sequence RSKRSRLLHSDYMNMTPRRPGPTRKHYQPYAPPRDFAAYRS (SEQ ID NO:302).

[0266] In another embodiment, the cytoplasmic domain comprises the signaling domain of CD3-zeta and the signaling domain of 4-1BB.

[0267] In another embodiment, the cytoplasmic domain comprises the signaling domain of CD3-zeta and the signaling domain of CD2. In some embodiments, the signaling domain of CD2 comprises the amino acid sequence
TKRKKQRSRRNDEELETRAHRVATEERGRKPHQIPASTPQN PATSQHPPPPGHRSQAPSHR
PPPPGHRVQHQPQKRPPAPSGTQVHQKGPPLPRPRVQPKPPHGA AENSLSPSSN (SEQ ID
NO:303).

[0268] In another embodiment, the cytoplasmic domain comprises the signaling domain of CD3-zeta, the signaling domain of CD28, and the signaling domain of CD2.

[0269] In another embodiment, the cytoplasmic domain comprises the signaling domain of CD3-zeta, the signaling domain of 4-1BB, and the signaling domain of CD2.

[0270] Inclusion of the CD2 signaling domain in the cytoplasmic domain allows for the tuning of CAR T cell cytokine production (see US Pat. No. 9,783,591, the contents of which are incorporated herein by reference in their entireties). As disclosed in US Pat. No. 9,783,591, inclusion of the CD2 signaling domain in the CAR cytoplasmic domain significantly alters CAR T cell cytokine production in both positive and negative directions, with the effect being dependent on the presence and identity of other costimulatory molecules in the costimulatory signaling region of the cytoplasmic domain. For example, in some embodiments, inclusion of the CD2 signaling domain and the CD28 signaling domain in the costimulatory signaling region of the cytoplasmic domain results in the release of significantly less IL2 relative to T cells expressing a CAR with CD28 but not CD2. A CAR T cell releasing less IL2 can result in reduced proliferation of immunosuppressive Treg cells. In some embodiments, inclusion of the CD2 signaling domain in the costimulatory signaling region of the cytoplasmic domain significantly reduces calcium influx in the CAR T cell. This has been shown to reduce activation-induced CAR T cell death.

5.4 Chimeric T Cell Receptors

[0271] The present disclosure provides chimeric T cell receptors (TCRs) comprising the anti-glyco-cMET antibodies or antigen-binding fragments described herein. The chimeric TCRs provide an anti-glyco-cMET specific antibody and TCR chimera that specifically binds to anti-glyco-cMET, and are capable of recruiting at least one TCR-associated signaling molecule (e.g., CD3 $\gamma\epsilon$, CD3 $\delta\epsilon$, and ζ). In some embodiments, the chimeric TCR comprises one or more antigen-binding fragments capable of binding glyco-cMET. Examples of antigen-binding fragments include by way of example and not limitation, Fab, Fab', F (ab')₂, Fv fragments, single chain Fv fragments (scFV) and single domain fragments. In some embodiments, an antigen-binding fragment of a chimeric T cell receptor comprises at least one anti-glyco-cMET variable heavy chain and at least one anti-glyco-cMET variable light chain as described herein.

[0272] TCRs occur as either an $\alpha\beta$ heterodimer or as a $\gamma\delta$ heterodimer, with T cells expressing either the $\alpha\beta$ form or the $\gamma\delta$ form TCR on the cell surface. The four chains (α , β , γ , δ) each have a characteristic extracellular structure consisting of a highly polymorphic "immunoglobulin variable region"-like N-terminal domain and an "immunoglobulin constant region"-like second domain. Each of these domains has a characteristic intra-domain disulfide bridge. The constant region is proximal to the cell membrane, followed by a connecting peptide, a transmembrane region and a short cytoplasmic tail. The covalent linkage between the 2 chains of the heterodimeric TCR is formed by the cysteine residue located within the short connecting

peptide sequence bridging the extracellular constant domain and the transmembrane region which forms a disulfide bond with the paired TCR chain cysteine residue at the corresponding position (Lefranc and Lefranc, "The T Cell Receptor FactsBook," Academic Press, 2001).

[0273] Several examples of chimeric TCRs are known in the art. See, e.g., Kuwana *et al.*, Biochem Biophys Res Commun. 149(3):960-968; Gross *et al.*, 1989, Proc Natl Acad Sci USA. 86:10024-10028; Gross & Eshhar, 1992, FASEB J. 6(15):3370-3378; Liu *et al.*, 2021, Sci Transl Med, 13:eabb5191, WO 2016/187349, WO 2017/070608, WO 2020/029774, and US Patent No. 7,741,465, the contents of each of which are incorporated herein by reference in their entireties.

[0274] A chimeric TCR generally comprises a first polypeptide chain comprising a first TCR domain, a second polypeptide chain comprising a second TCR domain, and an anti-glyco-cMET antigen binding fragment described herein. In some embodiments, the chimeric TCR comprises a single anti-glyco-cMET antigen binding fragment. In other embodiments, the chimeric TCR comprises a two or more anti-glyco-cMET antigen binding fragments. In certain embodiments, the chimeric TCR comprises two anti-glyco-cMET antigen binding fragments.

[0275] In some embodiments, the anti-glyco-cMET antigen binding fragment is an scFv described herein. In embodiments in which the chimeric TCR includes a single anti-glyco-cMET antigen binding fragment, a single anti-glyco-cMET scFv can be included in either the first polypeptide chain or the second polypeptide chain of the chimeric TCR. In embodiments in which the chimeric TCR includes, e.g., two anti-glyco-cMET antigen binding fragments, two anti-glyco-cMET scFVs can be included in either the first polypeptide chain or the second polypeptide chain of the chimeric TCR, or a first scFv can be included in the first polypeptide chain and a second scFv can be included in the second polypeptide chain. In embodiments in which two scFvs are included in one of either the first polypeptide chain or the second polypeptide chain of the chimeric TCR, the two scFvs can be linked via a peptide linker. In some embodiments, the chimeric TCR comprises two or more anti-glyco-cMET scFvs having the same amino acid sequence. In other embodiments, the chimeric TCR comprises two or more anti-glyco-cMET scFvs having different amino acid sequences.

[0276] In other embodiments, the anti-glyco-cMET antigen binding fragment is an Fv fragment. In some embodiments, an anti-glyco-cMET variable heavy chain (VH) described herein is included in one of the two polypeptide chains that associate to form the chimeric TCR. An anti-glyco-cMET variable light chain (VL) described herein can be included in the polypeptide chain that does not include the anti-glyco-cMET VH. When the first and second polypeptide chains dimerize, the anti-glyco-cMET VH and VL are brought together to form an anti-glyco-cMET Fv fragment. In some embodiments, the VH is included in the first polypeptide chain and the VL is included in the second polypeptide chain. In other embodiments, the VH is included in the second polypeptide chain and the VL is included in the first polypeptide chain.

[0277] In other embodiments, the anti-glyco-cMET antigen fragment is a Fab- domain, comprising VH, VL, CH1, and CL domains. In some embodiments, an anti-glyco-cMET variable heavy chain (VH) described herein and a CH1 domain is included in the first or second polypeptide chain. In some embodiments, an anti-glyco-cMET variable light chain (VL) described herein and a CL domain are included in the first or second polypeptide chain that does not include the anti-glyco-cMET VH and CH1. In other embodiments, an anti-glyco-cMET variable heavy chain (VH) and a CL domain is included in the first or second polypeptide chain. In some embodiments, an anti-glyco-cMET variable light chain (VL) and a CH1 domain are included in the polypeptide chain that does not include the anti-glyco-cMET VH and CL. When the first and second polypeptide chains dimerize, the anti-glyco-cMET VH and VL, and the CH1 and CL, are brought together to form an anti-glyco-cMET Fab domain. In some embodiments, the VH and the CH1 or CL is included in the first polypeptide chain, and the VL and the CL or CH1 is included in the second polypeptide chain. In other embodiments, the VH and the CH1 or CL is included in the second polypeptide chain, and the VL and the CH1 or CL is included in the first polypeptide chain.

[0278] In other embodiments, the anti-glyco-cMET VH and CH1 or CL are included in the first polypeptide chain of the second polypeptide chain, and the chimeric TCR further comprises a third polypeptide comprising the VL and either a CL domain or a CH1 domain. The third polypeptide is capable of associating with the VH and CH1 or CL of the first or second polypeptide chain, thus forming a Fab domain. In some embodiments, both the first and second polypeptide chains include a VH and a CH1 domain or a CL domain. Where both the first and second polypeptide chains include a VH and a CH1 or CL, a third polypeptide comprising a VL and a CL or CH1 associates with the first polypeptide chain to form a first Fab domain, and a fourth polypeptide comprising a VL and a CL or CH1 associates with the second polypeptide chain to form a second Fab domain.

[0279] First and second TCR domains are included in the first and second polypeptide chains, respectively, with the first TCR domain comprising a first TCR transmembrane domain from a first TCR subunit and the second TCR domain comprising a second TCR transmembrane domain from a second TCR subunit. In some embodiments, the first TCR subunit is a TCR α chain and the second TCR subunit is a TCR β chain. In other embodiments, the first TCR subunit is a TCR β chain and the second TCR subunit is a TCR α chain. In In some embodiments, the first TCR subunit is a TCR γ chain and the second TCR subunit is a TCR δ chain. In other embodiments, the first TCR subunit is a TCR δ chain and the second TCR subunit is a TCR γ chain. A TCR transmembrane domain from a TCR subunit can be a native TCR transmembrane domain, a natural or engineered variant thereof, or a fragment of the native or variant TCR transmembrane domain. In some embodiments, the first and/or second TCR transmembrane domains comprise, individually, an amino acid sequence of a TCR

transmembrane domain contained in one of SEQ ID NOS:77-80 of WO 2017/070608, which is incorporated by reference in its entirety. In other embodiments, the first and/or second TCR transmembrane domains comprise, individually, an amino acid sequence of SEQ ID NOS:1-4 of WO 2017/070608.

[0280] In some embodiments, in addition to the first and second TCR transmembrane domains, the first and second TCR domains also include first and second connecting peptides, respectively. The first and second connecting peptides are positioned at the N-terminus of the first and second TCR transmembrane domains, respectively. In some embodiments, the first connecting peptide comprises all or a portion of the connecting peptide of the first TCR subunit and/or the second connecting peptide comprises all or a portion of the connecting peptide of the second TCR subunit. In some embodiments, the first transmembrane domain and the first connecting peptide are derived from different TCR subunits and/or the second transmembrane domain and the second connecting peptide are derived from different TCR subunits. A connecting peptide from a TCR subunit can be a native TCR connecting peptide, a natural or engineered variant thereof, or a fragment of the native or variant TCR connecting peptide. In some embodiments, the first and/or second connecting peptides comprise, individually, an amino acid sequence of a connecting peptide contained in one of SEQ ID NOS:77-80 of WO 2017/070608. In other embodiments, the first and/or second connecting peptides comprise, individually, an amino acid sequence of SEQ ID NOS:5-12 of WO 2017/070608.

[0281] In some embodiments, the first and second TCR domains comprise a first and second TCR constant domain, respectively. The first and second TCR constant domains are positioned at the C-terminus of the first and second TCR transmembrane domains, respectively. If the first and/or second TCR domains include a TCR connecting peptide, the TCR constant domain can be positioned at the C-terminus of the TCR connecting peptide. In some embodiments, the first TCR constant domain comprises all or a portion of the constant domain of the first TCR subunit and/or the second TCR constant domain comprises all or a portion of the constant domain of the second TCR subunit. For example, in some embodiments, the first and/or second TCR constant domains are derived from TCR α and β subunit constant domains, or TCR γ and δ subunit constant domains. A TCR constant domain from a TCR subunit can be a native TCR intra constant cellular domain, a natural or engineered variant thereof, or a fragment of the native or variant TCR constant domain. In some embodiments, the first and/or second TCR constant domain comprise, individually an amino acid sequence of SEQ ID NOS:172, 174, 176, 178, 180, or 182, or the wildtype equivalent thereof.

[0282] In some embodiments, the first and second TCR domains comprise first and second TCR intracellular domains, respectively. The first and second TCR intracellular domains are positioned at the C-terminus of the first and second TCR transmembrane domains, respectively. In some embodiments, the first TCR intracellular domain comprises all or a portion

of the intracellular domain of the first TCR subunit and/or the second TCR intracellular domain comprises all or a portion of the intracellular domain of the second TCR subunit. A TCR intracellular domain from a TCR subunit can be a native TCR intracellular domain, a natural or engineered variant thereof, or a fragment of the native or variant TCR intracellular domain. In some embodiments, the first and/or second TCR intracellular domains comprise, individually, an amino acid sequence of a TCR intracellular domain contained in one of SEQ ID NOS:77-80 of WO 2017/070608. In other embodiments, the first and/or second TCR intracellular domain comprise, individually, an amino acid sequence of SEQ ID NOS:13-14 of WO 2017/070608.

[0283] In some embodiments, the first polypeptide chain of the chimeric TCR further comprises a first accessory intracellular domain C-terminal to the first TCR transmembrane domain and/or the second polypeptide chain of the chimeric TCR further comprises a second accessory intracellular domain C-terminal to the second transmembrane domain. In some embodiments, the first and/or second accessory intracellular domains comprise a TCR costimulatory domain. In some embodiments, the TCR costimulatory domain comprises all or a portion of the amino acid sequence of SEQ ID NO:70 or 71 of WO 2017/070608.

[0284] In some embodiments the first TCR domain is a fragment of the first TCR subunit and/or the second TCR subunit is a fragment of the second TCR subunit.

[0285] The first and second polypeptide chains that form the chimeric TCR are linked. In some embodiments, the first and second polypeptide chains that form the chimeric TCR are linked by a disulfide bond. In some embodiments, first and second polypeptide chains that form the chimeric TCR are linked by a disulfide bond between a residue in the first connecting peptide and a residue in the second connecting peptide.

[0286] In some embodiments, the first and second polypeptide chains are linked or otherwise associate. In some embodiments, the associated first and second polypeptide chains are capable of recruiting at least one TCR-associated signaling modules, such as, *e.g.*, CD3 $\delta\epsilon$, CD3 $\gamma\epsilon$, and $\zeta\zeta$. In certain embodiments, the associated first and second polypeptide chains are capable of recruiting each of CD3 $\delta\epsilon$, CD3 $\gamma\epsilon$, and $\zeta\zeta$, forming a TCR-CD3 complex.

[0287] In some embodiments, the first polypeptide chain comprises a first linker between the first TCR domain and an anti-glyco-cMET VH or VL of the scFv, Fv, or Fab fragment included in the first polypeptide chain. In some embodiments, the second polypeptide chain comprises a second linker between the second TCR domain and an anti-glyco-cMET VH or VL of the scFv, Fv, or Fab fragment included in the second polypeptide chain. In some embodiments, the first peptide linker and/or the second peptide linker comprises between about 5 to about 70 amino acids. In some embodiment, the first and/or second linker comprises a constant domain or fragment thereof from an immunoglobulin or T cell receptor subunit. In some embodiments, the first and/or second linker comprises an immunoglobulin constant domain or fragment thereof. For example in those embodiments described above comprising a CH1 or CL domain, the CH1

or CL domain functions as a linker between the TCR domain and the anti-glyco-cMET binding fragment, or a subpart (e.g., VH or VL) thereof. The immunoglobulin constant domain can also be, in addition to CH1 or CL, a CH2, CH3, or CH4 domain or fragment thereof. The immunoglobulin constant domains can be derived from an IgG (e.g., IgG1, IgG2, IgG3, or IgG4), IgA (e.g., IgA1 or IgA2), IgD, IgM, or IgE heavy chain. In some embodiments the constant domains can be derived from a human (e.g., IgG1, IgG2, IgG3, or IgG4), IgA (e.g., IgA1 or IgA2), IgD, IgM, or IgE heavy chain. In other embodiments, a TCR constant domain or fragment thereof described above functions as a linker between the TCR domain and the anti-glyco-cMET binding fragment, or a subpart (e.g., VH or VL) thereof. In some embodiments, the first and second linkers are capable of binding to one another.

[0288] In some embodiments, the first and second polypeptide chains are connected, at least temporarily, by a cleavable peptide linker. In some embodiments, the cleavable peptide linker is a furin-p2A cleavable peptide. The cleavable peptide linker can facilitate expression of the two polypeptide chains. The cleavable peptide linker can be configured to temporarily associate the first polypeptide chain with the second polypeptide chain during and/or shortly after protein translation.

[0289] In some embodiments, the chimeric TCR is a synthetic T cell receptor and antigen receptor (STAR), as described in Liu et al., 2021, Sci Transl Med, and WO 2020/029774, the contents of each of which are incorporated herein by reference in their entireties.

[0290] In some aspects, the STAR comprises, from N- to C-terminus, a first polypeptide chain comprising an anti-glyco- cMET variable heavy chain and a TCR α chain constant region domain; a cleavable peptide linker; and a second polypeptide chain comprising an anti-glyco- cMET variable light chain and a TCR β constant region domain (configuration STAR 1).

[0291] In other aspects, the STAR comprises, from N- to C-terminus, a first polypeptide chain comprising an anti-glyco- cMET variable heavy chain and a TCR β chain constant region domain; a cleavable peptide linker; and a second polypeptide chain comprising an anti-glyco- cMET variable light chain and a TCR α constant region domain (configuration STAR 2).

[0292] In other aspects, the STAR comprises, from N- to C-terminus, a first polypeptide chain comprising an anti-glyco- cMET variable light chain and a TCR α chain constant region domain; a cleavable peptide linker; and a second polypeptide chain comprising an anti-glyco- cMET variable heavy chain and a TCR β constant region domain (configuration STAR 3).

[0293] In other aspects, the STAR comprises, from N- to C-terminus, a first polypeptide chain comprising an anti-glyco- cMET variable light chain and a TCR β chain constant region domain; a cleavable peptide linker; and a second polypeptide chain comprising an anti-glyco- cMET variable heavy chain and a TCR α constant region domain (configuration STAR 4).

[0294] In certain embodiments, the TCR α chain constant region domain and the TCR β chain constant region domain of any one of configurations STAR 1 through STAR 4 can be replaced by TCR γ and TCR δ constant region domains, respectively.

[0295] The chimeric TCRs of the present disclosure can form complexes with TCR-associated signaling molecules (e.g., CD3 $\gamma\epsilon$, CD3 $\delta\epsilon$, and $\zeta\zeta$) endogenously expressed in T cells. These complexes provide for TCR signaling controlled by binding of the anti-glyco-cMET heavy and light variable chains by its target.

[0296] Chimeric TCRs of the disclosure are further described in numbered embodiments 735 to 834.

5.4.1. TCR Constant Domains

[0297] With respect to the TCR constant domains, the chimeric TCR can be designed to comprise constant regions that are derived from, e.g., human peripheral blood T cells. Nucleotide and corresponding amino acid sequences for TCR constant regions for use in chimeric TCRs according to the disclosure are provided in Table 5.

Table 5 Nucleotide and Amino Acid Sequences for TCR Constant Regions		
Description	Sequence	SEQ ID NO:
TCR α Constant Region – Nucleic Acid (human)	gatatccagaaccctgaccctgctgtctatcaactccgggactctaaatcca gtgacaagtctgtctgcctattcaccgatttgattctcaaacaaatgtgtcac aaagtaaggattctgtgtgtatatcacagacaaatgtgtctagacatgag gtctatggactcaagagcaacagtgctgtggcctggagcaacaaatcga ctttgcatgtgcaaacgccttcaacaacagcattattccagaagacaccttct tcccagcccagaaagtctctgtgatgcaagctggcgagaaaagcttg aaacagatacgaacctaaacttcaaacctgtcagtgattgggtccgaat cctcctcctgaaagtgccgggttaatctgctcatgacgctgaggctgtggt ccagc	304
TCR α Constant Region – Amino Acid (human)	XIQNPDPVYQLRDSKSSDKSVCLFTDFDSQTNVSQS KDSVYITDKCVLDMRSMDFKSN SAVAWSNK SDFAC ANAFNNSIIPEDTFFPSPESSCDVKLVEKSFETDTNLN FQNLSVIGFRILLKLVAGFNLLMTLRLWSS X=Asp, Asn, His, Tyr	305
TCR α Constant Region – Amino Acid (murine); Cysteine mutant	Aatatccagaaccagaaacctgctgtgtaccagttaaaagatcctcggctc caggacagaccctctgcctgttcaccgacttgactcccaaatcaatgtgc cgaaaaccttggaatctggaacgttcatcactgacaaaactgtgctggac atgaaagctatggattccaagagcaatggggccattgcctggagcaacca gacaagcttcacctgccaagatatctcaagagaccaacgccacctacc ccagttcagacgttccctgtgatgccacgttgactgagaaaagcttgaac agatatgaacctaaacttcaaacctgtcagttatgggactccgaatcctcc tgctgaaagtagccggatttaacctgctcatgacgctgaggctgtggtccag ttga	306
TCR α Constant Region – Amino Acid (murine); Cysteine mutant	XIQNPEPAVYQLKDPQRSQDSTLCLFTDFDSQINVPKT MESGTFITDKTVLDMKAMDSKSN GAIAWSNQT SFTC QDIFKETNATYPSSDVPCDATL TEKSFETDMNLNFQN LSVMGLRILLKLVAGFNLLMTLRLWSS X at 1, x=Asp, Asn, His, Tyr	307
TCR β Constant Region – Nucleic Acid (human)	gaggacctgaaaaacgtgttcccacccgaagtggccgtcttccaaccatc agaagcagagatctcccacacccaaaaggccacactgggtgtcctggcc acaggcttctcccaccacgtggagctgagctgggtggatgggaa ggagggtcacagtggggtctgcacagaccccgagcccctcaaggagca	308

Table 5 Nucleotide and Amino Acid Sequences for TCR Constant Regions		
Description	Sequence	SEQ ID NO:
	gcccgcctcaatgactccagatactgctgagcagccgcctgagggctc ggccacctctggcagaacccccgcaaccactccgctgtcaagtcagtt ctacgggctctcggagaatgacgagtggaaccaggataggccaaacc gtcaccagatcgtcagcgcgaggcctgggtagagcagactgtggcttt acctcgggtgctaccagcaaggggtcctgtctgccaccatcctatgaga tctgctaggggaaggccaccctgatgctgtgctggcagcgccttggtg atggccatggcaagagaaaggattc	
TCRβ Constant Region – Amino Acid (human)	EDLKNVFPPEVAVFEPSEAEISHTQKATLVCLATGFFP DHVELSWWWNGKEVHSGVCTDPQPLKEQPALNDSR YCLSSRLRVSATFWQNP RNHFRCQVQFYGLSENDE WTQDRAKPVTQIVSAEAWGRADCGFTSVSYQQGVL SATILYEILLGKATLYAVLV SALVLMAMVKRKD	309
TCRβ Constant Region – Amino Acid (murine); Cysteine mutant	gaggatcagagaaatgactccaccaaggctcctgttgagccatcaa aagcagagattgcaaacaacaagaaggctaccctcgtgctggccag gggcttctccctgaccacgtggagctgagctgggggtaatggcaagga ggccaagtgagggtcagcacggaccctcaggcctacaaggagagcaat tatagctactgctgagcagccgctgagggctctgctacctctggcaca atcctcgcaaccactccgctgccaagtcagttccatgggcttcagagga ggacaagtgccagagggctcacccaaacctgtcacacagaacatcagt gcagaggcctgggcccagcagactgtgggattacctcagcatcctatca acaaggggtctgtctgccaccatcctatgagatcctgctagggaaagcc accctgatgctgtgctgtcagtaactgggtgatggctatggtcaaaag aagaattca	310
TCRβ Constant Region – Amino Acid (murine); Cysteine mutant	EDLRNVTPPKVSLFEP SKAEIANKQKATLVCLARGFFP DHVELSWWWNGKEVHSGVSTDPQAYKESNYSYCLS SRLRVSATFWHNP RNHFRCQVQFHGLSEEDKWPEG SPKPVTQNISAEAWGRADCGITSASYQQGVL SATILY EILLGKATLYAVLVSTLVMMAMVKRKNS	311
TCRγ Constant Region – Amino Acid (human)	DKQLDADVSPKPTIFLPSIAETKLQKAGTYLCLLEKFFP DVIKIHWQEKSNTILGSQEGN TMKTNDTYMKFSWLT VPEKSLDKEHRCIVRHENNKNGVDQEIIFFPIKTDVITM DPKDNC SKDANDTLLLQLTNTSAYMYLLLLLKSVMY FAITCCLLRRTAFCCNGEKS	312
TCRγ Constant Region – Amino Acid (murine)	XKRLDADISPKPTIFLPSVAETNLHKTGTYLCLLEKFFP DVIRVYWKEKDGNTILDSQEGDTLKTNDTYMKFSWLT VPERAMGKEHRCIVKHENNKGGADQEIFFPSIKKVAV STKPTTCWQDKNDVLQLQFTITSAYTYLLLLLKSVIYL AIISFLLRRTSVCGNEKKS X = any naturally occurring amino acid	313
TCRδ Constant Region – Amino Acid (human)	SQPHTKPSVFMKNGTNVACL VKEFYPKDIRINLVSS KKITEFDPAIVISPSGKYNAVKLGKYEDSNSVTCSVQH DNKTVHSTDFEVKTDSTDHV KPKETENTKQPSKSCH KPKAIVHTEKVNMMSLTVLGLRMLFAKTAVNFLLTAK LFFL	314
TCRδ Constant Region – Amino Acid (murine)	XSQPPAKPSVFIMKNGTNVACL VKDFYPKEVTISLRSS KKIVEFDPAIVISPSGKYSAVKLGQYGDSNSVTCSVQH NSETVHSTDFEPYANSFNNEKLPEPENDTQISEPCYG PRVTVHTEKVNMMSLTVLGLRLLFAKTI AINFLLVKLF F X = any naturally occurring amino acid	315

[0298] In certain embodiments the TCR constant domain of the chimeric TCR can be modified to provide for additional bonds between two TCR constant domains of the chimeric TCR. In some embodiments, the residue corresponding to position 48 of the wildtype human TCRα

constant domain is mutated to cysteine and the residue corresponding to position 57 of the wildtype human TCR β constant domain is mutated to cysteine, as shown in Table 5. This results in the formation of a disulfide linkage between TCR α and TCR β constant domains, resulting in a disulfide bond between the first and second polypeptide chains of the chimeric TCR. In some embodiments, the residue corresponding to position 85 of the wildtype human TCR α constant domain is mutated to alanine and the residue corresponding to position 88 of the wildtype human TCR β constant domain is mutated to glycine, as shown in Table 5. Again, this results in the formation of a disulfide linkage between TCR α and TCR β constant regions.

5.4.2. Cleavable Linkers

[0299] In some embodiments, the two polypeptide chains of the chimeric TCRs of the disclosure are linked via a cleavable peptide linker. In some embodiments, the two polypeptide chains of the chimeric TCR are linked via a furin-P2A peptide linker, which provides a protease cleavage site between the two polypeptide chains. The two polypeptide chains can thus be transcribed and translated into a fusion protein, which is subsequently cleaved by a protease into two distinct protein subunits. In some embodiments, the two resulting protein subunits are covalently bound through disulfide bonds, and subsequently form a complex with the endogenous CD3 subunits of T cells.

[0300] In some embodiments, the furin-P2A peptide linker comprises the sequence RAKRSGSGATNFSLKQAGDVEENPGP (SEQ ID NO:316).

[0301] In some embodiments, the furin-P2A peptide linker comprises the sequence ATNFSLKQAGDVEENPGP (SEQ ID NO:317).

5.5 Neuraminidase

[0302] Sialic acids are terminal sugars of glycans on either glycoproteins or glycolipids on the cell surface, and have been shown to be aberrantly expressed during tumor transformation and malignant progression. Hypersialylation frequently occurs in tumor tissues due to aberrant expression of sialyltransferases/sialidases. This can result in accelerated cancer progression. Sialylation facilitates immune escape, enhances tumor proliferation and metastasis, helps tumor angiogenesis, and assists in resisting apoptosis and cancer therapy.

[0303] Host cells (e.g., T cells, NK cells) expressing a CAR of the disclosure can be engineered to coexpress a cell surface or secreted neuraminidase (sialidase) along with the CAR. The cell surface neuraminidase, anchored to the cell surface via a heterologous transmembrane, gives the host cell glycoediting activity. This enhances cytotoxic effects and anti-tumor efficacy of the CAR-T cell and immune cells such as innate NK cells and monocytes. Host cells coexpressing a CAR and an engineered neuraminidase are described in PCT Publication No WO2020/236964, which is incorporated herein by reference in its entirety.

[0304] A neuraminidase can be coexpressed in a host cell along with a CAR described herein. Exemplary host cells coexpressing a neuraminidase and a CAR are described in the specific embodiments.

[0305] The neuraminidase can be included as a domain of a fusion protein described herein.

[0306] In certain embodiments, the neuraminidase is EC 3.2.1.18 or EC 3.2.1.129.

[0307] In some embodiments, the neuraminidase is derived from *Micromonospora viridifaciens*.

[0308] In some aspects, the neuraminidase comprises an amino acid sequence having at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to:

GGSPVPPGGEPLYTEQDLAVNGREGFPNYRIPALTVTPDGDLLASYDGRPTGIDAPGPNSILQ
 RRSTDGGRTWGEQQVVSAGQTTAPIKGFSDPSYLVRETGTIFNFHVYSQRQGFAGSRPGTD
 PADPNVLHANVATSTDGGLTWSHRTITADITPDGWRSRFAASGEGIQLRYGPHAGRLIQQYTI
 INAAGAFQAVSVYSDDHGRTWRAGEAVGVGMDENKTVELSDGRVLLNSRDSARSQGYRQVAV
 STDGGHSYGPVTIDRDLDPPTNNASIIRAFDAPAGSARAKVLLFSNAASQTSRSQGTIRMSCD
 DGQTPVSKVFQPGSMSYSTLTALPDGTYGLLYEPGTGIRYANFNLAWLGG (SEQ ID
 NO:318).

[0309] The neuraminidase can be retained at a surface of a host cell engineered to express the neuraminidase, or can be secreted by a host cell engineered to express the neuraminidase. The host cell engineered to express the neuraminidase can include, for example, a vector encoding the neuraminidase.

5.6 MicAbodies

[0310] The present disclosure provides MicAbodies comprising the anti-glyco-cMET antibodies and antigen-binding fragments of the disclosure. MicAbodies are fusion proteins comprising an antibody or antigen-binding fragment and an engineered MHC-class I-chain-related (MIC) protein domain. MIC proteins are the natural ligands of human NKG2D receptors expressed on the surface of NK cells, and the $\alpha 1$ - $\alpha 2$ domain of MIC proteins provides the binding site for the NKG2D receptor. By fusing an engineered MIC protein domain (e.g. an engineered $\alpha 1$ - $\alpha 2$ domain) to a cancer-targeting antibody or antigen-binding fragment, T-cells expressing an engineered NKG2D receptor capable of binding the engineered MIC protein domain can be targeted to cancer cells. Engineered MIC protein domains that can be included in MicAbodies of the disclosure, and NKG2D receptors capable of binding the engineered MIC protein domains, CARs and CAR T cells comprising the NKG2D receptors are described in U.S. publication nos. US 2011/0183893, US2011/0311561, US 2015/0165065, and US 2016/0304578 and PCT publication nos. WO 2016/090278, WO 2017/024131, WO 2017/222556, and WO 2019/191243, the contents of which are incorporated herein by reference in their entireties.

[0311] In some embodiments, the MicAbodies of the disclosure comprise $\alpha 1$ - $\alpha 2$ domains which are at least 80% identical or homologous to the $\alpha 1$ - $\alpha 2$ domain of an NKG2D ligand (e.g., MICA, MICB, ULBP1, ULBP2, ULBP3, ULBP4, ULBP5, ULBP6, or OMCP). Exemplary amino acid sequences of MICA, MICB, ULBP1, ULBP2, ULBP3, ULBP4, ULBP5, ULBP6, and OMCP are set forth as SEQ ID NOS: 1-9 of WO 2019/191243, respectively, the sequences of which are incorporated herein by reference. In other embodiments, the $\alpha 1$ - $\alpha 2$ domain is 85% identical to a native or natural $\alpha 1$ - $\alpha 2$ domain of an NKG2D ligand. In yet other embodiments, the $\alpha 1$ - $\alpha 2$ domain is 90% identical to a native or natural $\alpha 1$ - $\alpha 2$ domain of a natural NKG2D ligand protein and binds non-natural NKG2D.

[0312] In some embodiments, the MicAbodies of the disclosure comprise $\alpha 1$ - $\alpha 2$ domains which are at least 80% identical or homologous to a native or natural $\alpha 1$ - $\alpha 2$ domain of a human MICA or MICB protein and bind NKG2D. In some embodiments, the $\alpha 1$ - $\alpha 2$ domain is 85% identical to a native or natural $\alpha 1$ - $\alpha 2$ domain of a human MICA or MICB protein and binds NKG2D. In other embodiments, the $\alpha 1$ - $\alpha 2$ domain is 90%, 95%, 96%, 97%, 98%, or 99% identical to a native or natural $\alpha 1$ - $\alpha 2$ platform domain of a human MICA or MICB protein and binds NKG2D.

[0313] In some embodiments, specific mutations in $\alpha 1$ - $\alpha 2$ domains of NKG2D ligands can be made to create non-natural $\alpha 1$ - $\alpha 2$ domains that bind non-natural NKG2D receptors, themselves engineered so as to have reduced affinity for natural NKG2D ligands. This can be done, for example, through genetic engineering. A non-natural NKG2D receptor so modified can be used to create on the surface of NK- or T-cells of the immune system an NKG2D-based CAR that can preferentially bind to and be activated by molecules comprised of the non-natural $\alpha 1$ - $\alpha 2$ domains. These pairs of non-natural NKG2D receptors and their cognate non-natural NKG2D ligands can provide important safety, efficacy, and manufacturing advantages for treating cancer and viral infections as compared to traditional CAR-T cells and CAR-NK cells. Activation of CAR-T cells and CAR-NK cells having a NKG2D-based CAR can be controlled by administration of a MicAbody. In the event that an adverse event develops, the dosing regimen of the MicAbody can be modified rather than having to deploy an induced suicide mechanism to destroy the infused CAR cells.

[0314] MicAbodies can be generated by attaching an antibody or antigen-binding fragment to an engineered $\alpha 1$ - $\alpha 2$ domain via a linker, e.g., APTSSSGGGGS (SEQ ID NO:319), GGGGS (SEQ ID NO:320), or GGGGS (SEQ ID NO:293). For example, an $\alpha 1$ - $\alpha 2$ domain can be fused to the C-terminus of an IgG heavy chain or light chain, for example, as described in WO 2019/191243.

[0315] In some embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

EPHSLRYNLTVLSWDGVSQSGFLTEVHLDGQPFLRCRQKCRAPQGQWAEDVLGNKTWD
RETRDLTGWGTLLMTLAHIKDQKEGLHSLQEIRVCEIHEDNSTRSSQHFYYDGELFLSQNLET

LEWTMPQSSRAQTLAMNVRNFLKEDAMETDIGYRLMRADCLSELRRYLKSGVLRRTV (SEQ ID NO:321) (MICA25.17).

[0316] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

EPHSLRYNLTVLSWDGVSQSGFLTEVHLDGQPFLRCRQKCRAPQGGWAEDVLGNKTWD
RETRDLTGWGTFLRMTLAHIKDQKEGLHSLQEIRVCEIHEDNSTRSSQHFYYDGELFLSQNLET
LEWTMPQSSRAQTLAMNVRNFLKEDAMETDRSGLLMRADCLSELRRYLKSGVLRRTV (SEQ
ID NO:322) (MICA25.18).

[0317] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

AAEPHLSYDITVIPKFRPGPRWCAVQGQVDEKTFLLHYDCGNKTVTPVSPLGKKLNVTTAWKA
QNPVLRVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFD
SEKRMWTTVHPGARKMKEKWENDKVVATTLYTWSMGDCIGWLEDFLMGMDSTLEPSAGAP
(SEQ ID NO:323) (ULBP2.S1).

[0318] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

AAEPHLSYDITVIPKFRPGPRWCAVQGQVDEKTFLLHYDCGNKTVTPVSPLGKKLNVTTAWKA
QNPVLRVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFD
SEKRMWTTVHPGARKMKEKWENDKVVATLMRIWSMGDCIGWLEDFLMGMDSTLEPSAGAP
(SEQ ID NO:324) (ULBP2.S2).

[0319] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

AAEPHLSYDITVIPKFRPGPRWCAVQGQVDEKTFLLHYDCGNKTVTPVSPLGKKLNVTTAWKA
QNPVLRVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFD
SEKRMWTTVHPGARKMKEKWENDKVVATKLYLWSMGDCIGWLEDFLMGMDSTLEPSAGAP
(SEQ ID NO:325) (ULBP2.S3).

[0320] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

AAEPHSLWYNFTIIHLPRHGQQWCEVQSQVDQKNFLSYDCGSDKVLSMGHLEEQLYATDAW
GKQLEMLREVGQRLRLELADTELEDFTPSGPLTLQVRMSCESEADGYIRGSWQFSFDGRKFL
LFDSNNRKWTVHAGARRMKEKWEKDSGLTTDLIRSMGDCKSWLRDFLMHRKKRLEPTAP
(SEQ ID NO:326) (ULBP3.S1).

[0321] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

AAEPHSLWYNFTIIHLPRHGQQWCEVQSQVDQKNFLSYDCGSDKVLSMGHLEEQLYATDAW
GKQLEMLREVGQRLRLELADTELEDFTPSGPLTLQVRMSCESEADGYIRGSWQFSFDGRKFL

LFDSNNRKWTVVHAGARRMKEKWEKDSGLTTYFYLRSMGDCKSWLRDFLMHRKKRLEPTAP (SEQ ID NO:327) (ULBP3.S2).

[0322] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

EPHLSYDITVIPKFRPGPRWCAVQQQVDEKTFLHYDCGNKTVTPVSPLGKKLNVTTAWKAQN
PVLREVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFDSE
KRMWTTVHPGARKMKEKWENDKVVATILWQTSMGDCIGWLEDFLMGMDSTLEPS (SEQ ID
NO:328) (ULBP2.C).

[0323] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

EPHLSYDITVIPKFRPGPRWCAVQQQVDEKTFLHYDCGNKTVTPVSPLGKKLNVTTAWKAQN
PVLREVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFDSE
KRMWTTVHPGARKMKEKWENDKVVATLLWGWSMGDCIGWLEDFLMGMDSTLEPS (SEQ ID
NO:329) (ULBP2.R).

[0324] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

EPHLSYDITVIPKFRPGPRWCAVQQQVDEKTFLHYDCGNKTVTPVSPLGKKLNVTTAWKAQN
PVLREVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFDSE
KRMWTTVHPGARKMKEKWENDKVVATMFWSWSMGDCIGWLEDFLMGMDSTLEPS (SEQ ID
NO:330) (ULBP2.AA).

[0325] In other embodiments, the MicAbodies of the disclosure comprise an engineered $\alpha 1$ - $\alpha 2$ domain comprising the amino acid sequence

EPHLSYDITVIPKFRPGPRWCAVQQQVDEKTFLHYDCGNKTVTPVSPLGKKLNVTTAWKAQN
PVLREVVDILTEQLWDIQLENYTPKEPLTLQARMSCEQKAEGHSSGSWQFSFDGQIFLLFDSE
KRMWTTVHPGARKMKEKWENDKVVATLMWQWSMGDCIGWLEDFLMGMDSTLEPS (SEQ ID
NO:331) (ULBP2.AB).

[0326] An exemplary engineered NKG2D receptor comprises the amino acid sequence

NSLNFNQEVIPLTESYCGPCPKNWICYKNNCYQFFDESKNWYESQASCMSQNASLLKVYSKE
DQDLLKLVKSYHWMGLVHIPTNGSWQWEDGSILSPNLLTIEMQKGDALYASSFKGYIENCST
PNTYICMQRTV (SEQ ID NO:332) in which the tyrosine at position 73 has been replaced with
another amino acid, for example alanine.

[0327] Another exemplary engineered NKG2D receptor comprises the amino acid sequence

FLNSLNFNQEVIPLTESYCGPCPKNWICYKNNCYQFFDESKNWYESQASCMSQNASLLKVYS
KEDQDLLKLVKSYHWMGLVHIPTNGSWQWEDGSILSPNLLTIEMQKGDALYASSFKGYIENC
STPNTYICMQRTV (SEQ ID NO:333) in which the tyrosines at positions 75 and 122 have

been replaced with another amino acid, for example alanine at position 75 and phenylalanine at position 122.

5.7 Nucleic Acids, Recombinant Vectors and Host Cells

[0328] The present disclosure encompasses nucleic acid molecules encoding immunoglobulin light and heavy chain genes for anti-glyco-cMET antibodies, vectors comprising such nucleic acids, and host cells capable of producing the anti-glyco-cMET antibodies of the disclosure. In certain aspects, the nucleic acid molecules encode, and the host cells are capable of expressing, the anti-glyco-cMET antibodies and antibody-binding fragments of the disclosure (*e.g.*, as described in Section 5.1 and numbered embodiments 1 to 657) as well as fusion proteins (*e.g.*, as described in numbered embodiments 664 to 688), and chimeric antigen receptors (*e.g.*, as described in Section 5.3 and numbered embodiments 689 to 724) and chimeric TCRs (*e.g.*, as described in Section 5.4 and numbered embodiments 735 to 834) containing them. Exemplary vectors of the disclosure are described in numbered embodiments 837 to 839 and exemplary host cells are described in numbered embodiments 840 to 846.

[0329] An anti-glyco-cMET antibody of the disclosure can be prepared by recombinant expression of immunoglobulin light and heavy chain genes in a host cell. To express an antibody recombinantly, a host cell is transfected with one or more recombinant expression vectors carrying DNA fragments encoding the immunoglobulin light and heavy chains of the antibody such that the light and heavy chains are expressed in the host cell and, optionally, secreted into the medium in which the host cells are cultured, from which medium the antibodies can be recovered. Standard recombinant DNA methodologies are used to obtain antibody heavy and light chain genes, incorporate these genes into recombinant expression vectors and introduce the vectors into host cells, such as those described in *Molecular Cloning: A Laboratory Manual, Second Edition* (Sambrook, Fritsch and Maniatis (eds), Cold Spring Harbor, N. Y., 1989), *Current Protocols in Molecular Biology* (Ausubel, F. M. *et al.*, eds., Greene Publishing Associates, 1989) and in U.S. Pat. No. 4,816,397.

[0330] To generate nucleic acids encoding such anti-glyco-cMET antibodies, DNA fragments encoding the light and heavy chain variable regions are first obtained. These DNAs can be obtained by amplification and modification of germline DNA or cDNA encoding light and heavy chain variable sequences, for example using the polymerase chain reaction (PCR). Germline DNA sequences for human heavy and light chain variable region genes are known in the art (see, *e.g.*, the "VBASE" human germline sequence database; see also Kabat *et al.*, 1991, *Sequences of Proteins of Immunological Interest, Fifth Edition*, U.S. Department of Health and Human Services, NIH Publication No. 91-3242; Tomlinson *et al.*, 1992, *J. Mol. Biol.* 22T:116-198; and Cox *et al.*, 1994, *Eur. J. Immunol.* 24:827-836; the contents of each of which are incorporated herein by reference).

[0331] Once DNA fragments encoding anti-glyco-cMET antibody-related V_H and V_L segments are obtained, these DNA fragments can be further manipulated by standard recombinant DNA techniques, for example to convert the variable region genes to full-length antibody chain genes, to Fab fragment genes or to a scFv gene. In these manipulations, a V_H - or V_L -encoding DNA fragment is operatively linked to another DNA fragment encoding another protein, such as an antibody constant region or a flexible linker. The term "operatively linked," as used in this context, is intended to mean that the two DNA fragments are joined such that the amino acid sequences encoded by the two DNA fragments remain in-frame.

[0332] The isolated DNA encoding the V_H region can be converted to a full-length heavy chain gene by operatively linking the V_H -encoding DNA to another DNA molecule encoding heavy chain constant regions (CH_1 , CH_2 , CH_3 and, optionally, CH_4). The sequences of human heavy chain constant region genes are known in the art (see, e.g., Kabat *et al.*, 1991, Sequences of Proteins of Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, NIH Publication No. 91-3242) and DNA fragments encompassing these regions can be obtained by standard PCR amplification. The heavy chain constant region can be an IgG₁, IgG₂, IgG₃, IgG₄, IgA, IgE, IgM or IgD constant region, but in certain embodiments is an IgG₁ or IgG₄ constant region. For a Fab fragment heavy chain gene, the V_H -encoding DNA can be operatively linked to another DNA molecule encoding only the heavy chain CH1 constant region.

[0333] The isolated DNA encoding the V_L region can be converted to a full-length light chain gene (as well as a Fab light chain gene) by operatively linking the V_L -encoding DNA to another DNA molecule encoding the light chain constant region, CL. The sequences of human light chain constant region genes are known in the art (see, e.g., Kabat *et al.*, 1991, Sequences of Proteins of Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, NIH Publication No. 91-3242) and DNA fragments encompassing these regions can be obtained by standard PCR amplification. The light chain constant region can be a kappa or lambda constant region, but in certain embodiments is a kappa constant region.

[0334] To create a scFv gene, the V_H - and V_L -encoding DNA fragments can be operatively linked to another fragment encoding a flexible linker, e.g., encoding the amino acid sequence $(Gly_4\sim Ser)_3$, such that the V_H and V_L sequences can be expressed as a contiguous single-chain protein, with the V_H and V_L regions joined by the flexible linker (see, e.g., Bird *et al.*, 1988, Science 242:423-426; Huston *et al.*, 1988, Proc. Natl. Acad. Sci. USA 85:5879-5883; McCafferty *et al.*, 1990, Nature 348:552-554).

[0335] To express the anti-glyco-cMET antibodies of the disclosure, DNAs encoding partial or full-length light and heavy chains, obtained as described above, are inserted into expression vectors such that the genes are operatively linked to transcriptional and translational control sequences. In this context, the term "operatively linked" is intended to mean that an antibody

gene is ligated into a vector such that transcriptional and translational control sequences within the vector serve their intended function of regulating the transcription and translation of the antibody gene. The expression vector and expression control sequences are chosen to be compatible with the expression host cell used. The antibody light chain gene and the antibody heavy chain gene can be inserted into separate vectors or, more typically, both genes are inserted into the same expression vector.

[0336] The antibody genes are inserted into the expression vector by standard methods (*e.g.*, ligation of complementary restriction sites on the antibody gene fragment and vector, or blunt end ligation if no restriction sites are present). Prior to insertion of the anti-glyco-cMET antibody-related light or heavy chain sequences, the expression vector can already carry antibody constant region sequences. For example, one approach to converting the anti-glyco-cMET monoclonal antibody-related V_H and V_L sequences to full-length antibody genes is to insert them into expression vectors already encoding heavy chain constant and light chain constant regions, respectively, such that the V_H segment is operatively linked to the CH segment(s) within the vector and the V_L segment is operatively linked to the CL segment within the vector. Additionally or alternatively, the recombinant expression vector can encode a signal peptide that facilitates secretion of the antibody chain from a host cell. The antibody chain gene can be cloned into the vector such that the signal peptide is linked in-frame to the amino terminus of the antibody chain gene. The signal peptide can be an immunoglobulin signal peptide or a heterologous signal peptide (*i.e.*, a signal peptide from a non-immunoglobulin protein).

[0337] In addition to the antibody chain genes, the recombinant expression vectors of the disclosure carry regulatory sequences that control the expression of the antibody chain genes in a host cell. The term "regulatory sequence" is intended to include promoters, enhancers and other expression control elements (*e.g.*, polyadenylation signals) that control the transcription or translation of the antibody chain genes. Such regulatory sequences are described, for example, in Goeddel, *Gene Expression Technology: Methods in Enzymology* 185, Academic Press, San Diego, Calif., 1990. It will be appreciated by those skilled in the art that the design of the expression vector, including the selection of regulatory sequences may depend on such factors as the choice of the host cell to be transformed, the level of expression of protein desired, *etc.* Suitable regulatory sequences for mammalian host cell expression include viral elements that direct high levels of protein expression in mammalian cells, such as promoters and/or enhancers derived from cytomegalovirus (CMV) (such as the CMV promoter/enhancer), Simian Virus 40 (SV40) (such as the SV40 promoter/enhancer), adenovirus, (*e.g.*, the adenovirus major late promoter (AdMLP)) and polyoma. For further description of viral regulatory elements, and sequences thereof, see, *e.g.*, U.S. Pat. No. 5,168,062 by Stinski, U.S. Pat. No. 4,510,245 by Bell *et al.*, and U.S. Pat. No. 4,968,615 by Schaffner *et al.*

[0338] In addition to the antibody chain genes and regulatory sequences, the recombinant expression vectors of the disclosure can carry additional sequences, such as sequences that regulate replication of the vector in host cells (e.g., origins of replication) and selectable marker genes. The selectable marker gene facilitates selection of host cells into which the vector has been introduced (see, e.g., U.S. Pat. Nos. 4,399,216, 4,634,665 and 5,179,017, all by Axel *et al.*). For example, typically the selectable marker gene confers resistance to drugs, such as G418, hygromycin or methotrexate, on a host cell into which the vector has been introduced. Suitable selectable marker genes include the dihydrofolate reductase (DHFR) gene (for use in DHFR⁻ host cells with methotrexate selection/amplification) and the neo gene (for G418 selection). For expression of the light and heavy chains, the expression vector(s) encoding the heavy and light chains is transfected into a host cell by standard techniques. The various forms of the term “transfection” are intended to encompass a wide variety of techniques commonly used for the introduction of exogenous DNA into a prokaryotic or eukaryotic host cell, e.g., electroporation, lipofection, calcium-phosphate precipitation, DEAE--dextran transfection and the like.

[0339] It is possible to express the antibodies of the disclosure in either prokaryotic or eukaryotic host cells. In certain embodiments, expression of antibodies is performed in eukaryotic cells, e.g., mammalian host cells, of optimal secretion of a properly folded and immunologically active antibody. Exemplary mammalian host cells for expressing the recombinant antibodies of the disclosure include Chinese Hamster Ovary (CHO cells) (including DHFR⁻ CHO cells, described in Urlaub and Chasin, 1980, Proc. Natl. Acad. Sci. USA 77:4216-4220, used with a DHFR selectable marker, e.g., as described in Kaufman and Sharp, 1982, Mol. Biol. 159:601-621), NSO myeloma cells, COS cells and SP2 cells. When recombinant expression vectors encoding antibody genes are introduced into mammalian host cells, the antibodies are produced by culturing the host cells for a period of time sufficient to allow for expression of the antibody in the host cells or secretion of the antibody into the culture medium in which the host cells are grown. Antibodies can be recovered from the culture medium using standard protein purification methods. Host cells can also be used to produce portions of intact antibodies, such as Fab fragments or scFv molecules. It is understood that variations on the above procedure are within the scope of the present disclosure. For example, it can be desirable to transfect a host cell with DNA encoding either the light chain or the heavy chain (but not both) of an anti-glyco-cMET antibody of this disclosure.

[0340] For expression of a CAR of the disclosure, for example as described in Section 5.3 and in numbered embodiments 689 to 724, it is preferable that the host cell is a T cell, preferably a human T cell. In some embodiments, the host cell exhibits an anti-tumor immunity when the cell is cross-linked with cMET on a tumor cell. Detailed methods for producing the T cells of the disclosure are described in Section 5.7.1.

[0341] For expression of a chimeric TCR of the disclosure, for example as described in Section 5.4 and in numbered embodiments 735 to 834, it is preferable that the host cell is a T cell, preferably a human T cell. In some embodiments, the host cell exhibits an anti-tumor immunity when the cell is cross-linked with glyco-cMET on a tumor cell. Detailed methods for producing the T cells of the disclosure are described in Section 5.7.1.

[0342] Recombinant DNA technology can also be used to remove some or all of the DNA encoding either or both of the light and heavy chains that is not necessary for binding to glyco-cMET. The molecules expressed from such truncated DNA molecules are also encompassed by the antibodies of the disclosure.

[0343] For recombinant expression of an anti-glyco-cMET antibody of the disclosure, the host cell can be co-transfected with two expression vectors of the disclosure, the first vector encoding a heavy chain derived polypeptide and the second vector encoding a light chain derived polypeptide. The two vectors can contain identical selectable markers, or they can each contain a separate selectable marker. Alternatively, a single vector can be used which encodes both heavy and light chain polypeptides.

[0344] Once a nucleic acid encoding one or more portions of an anti-glyco-cMET antibody, further alterations or mutations can be introduced into the coding sequence, for example to generate nucleic acids encoding antibodies with different CDR sequences, antibodies with reduced affinity to the Fc receptor, or antibodies of different subclasses.

[0345] The anti-glyco-cMET antibodies of the disclosure can also be produced by chemical synthesis (*e.g.*, by the methods described in Solid Phase Peptide Synthesis, 2nd ed., 1984 The Pierce Chemical Co., Rockford, Ill.). Variant antibodies can also be generated using a cell-free platform (see, *e.g.*, Chu *et al.*, Biochemia No. 2, 2001 (Roche Molecular Biologicals) and Murray *et al.*, 2013, Current Opinion in Chemical Biology, 17:420-426).

[0346] Once an anti-glyco-cMET antibody of the disclosure has been produced by recombinant expression, it can be purified by any method known in the art for purification of an immunoglobulin molecule, for example, by chromatography (*e.g.*, ion exchange, affinity, and sizing column chromatography), centrifugation, differential solubility, or by any other standard technique for the purification of proteins. Further, the anti-glyco-cMET antibodies of the present disclosure and/or binding fragments can be fused to heterologous polypeptide sequences described herein or otherwise known in the art to facilitate purification.

[0347] Once isolated, the anti-glyco-cMET antibody can, if desired, be further purified, *e.g.*, by high performance liquid chromatography (see, *e.g.*, Fisher, Laboratory Techniques In Biochemistry And Molecular Biology, Work and Burdon, eds., Elsevier, 1980), or by gel filtration chromatography on a Superdex™ 75 column (Pharmacia Biotech AB, Uppsala, Sweden).

5.7.1. Recombinant Production of CARs and Chimeric TCRs in T Cells

[0348] In some embodiments, nucleic acids encoding the anti-glyco-cMET CARs or chimeric TCRs of the disclosure are delivered into cells using a retroviral or lentiviral vector. CAR- or chimeric TCR-expressing retroviral and lentiviral vectors can be delivered into different types of eukaryotic cells as well as into tissues and whole organisms using transduced cells as carriers or cell-free local or systemic delivery of encapsulated, bound or naked vectors. The method used can be for any purpose where stable expression is required or sufficient.

[0349] In other embodiments, the CAR or chimeric TCR sequences are delivered into cells using *in vitro* transcribed mRNA. *In vitro* transcribed mRNA CAR or chimeric TCR can be delivered into different types of eukaryotic cells as well as into tissues and whole organisms using transfected cells as carriers or cell-free local or systemic delivery of encapsulated, bound or naked mRNA. The method used can be for any purpose where transient expression is required or sufficient.

[0350] In another embodiment, the desired CAR or chimeric TCR can be expressed in the cells by way of transposons.

[0351] One advantage of RNA transfection methods of the disclosure is that RNA transfection is essentially transient and a vector-free: an RNA transgene can be delivered to a lymphocyte and expressed therein following a brief *in vitro* cell activation, as a minimal expressing cassette without the need for any additional viral sequences. Under these conditions, integration of the transgene into the host cell genome is unlikely. Cloning of cells is not necessary because of the efficiency of transfection of the RNA and its ability to uniformly modify the entire lymphocyte population.

[0352] Genetic modification of T cells with *in vitro*-transcribed RNA (IVT-RNA) makes use of two different strategies both of which have been successively tested in various animal models. Cells are transfected with *in vitro*-transcribed RNA by means of lipofection or electroporation. Preferably, it is desirable to stabilize IVT-RNA using various modifications in order to achieve prolonged expression of transferred IVT-RNA.

[0353] Some IVT vectors are known in the literature which are utilized in a standardized manner as template for *in vitro* transcription and which have been genetically modified in such a way that stabilized RNA transcripts are produced. Currently protocols used in the art are based on a plasmid vector with the following structure: a 5' RNA polymerase promoter enabling RNA transcription, followed by a gene of interest which is flanked either 3' and/or 5' by untranslated regions (UTR), and a 3' polyadenyl cassette containing 50-70 A nucleotides. Prior to *in vitro* transcription, the circular plasmid is linearized downstream of the polyadenyl cassette by type II restriction enzymes (recognition sequence corresponds to cleavage site). The polyadenyl cassette thus corresponds to the later poly(A) sequence in the transcript. As a result of this procedure, some nucleotides remain as part of the enzyme cleavage site after linearization and

extend or mask the poly (A) sequence at the 3' end. It is not clear, whether this nonphysiological overhang affects the amount of protein produced intracellularly from such a construct.

[0354] RNA has several advantages over more traditional plasmid or viral approaches. Gene expression from an RNA source does not require transcription and the protein product is produced rapidly after the transfection. Further, since the RNA has to only gain access to the cytoplasm, rather than the nucleus, and therefore typical transfection methods result in an extremely high rate of transfection. In addition, plasmid-based approaches require that the promoter driving the expression of the gene of interest be active in the cells under study.

[0355] In another aspect, the RNA construct can be delivered into the cells by electroporation. See, e.g., the formulations and methodology of electroporation of nucleic acid constructs into mammalian cells as taught in US 2004/0014645, US 2005/0052630A1, US 2005/0070841A1, US 2004/0059285A1, US 2004/0092907A1. The various parameters including electric field strength required for electroporation of any known cell type are generally known in the relevant research literature as well as numerous patents and applications in the field. See e.g., U.S. Pat. No. 6,678,556, U.S. Pat. No. 7,171,264, and U.S. Pat. No. 7,173,116. Apparatus for therapeutic application of electroporation are available commercially, e.g., the MedPulser™ DNA Electroporation Therapy System (Inovio/Genetronics, San Diego, Calif.), and are described in patents such as U.S. Pat. No. 6,567,694; U.S. Pat. No. 6,516,223, U.S. Pat. No. 5,993,434, U.S. Pat. No. 6,181,964, U.S. Pat. No. 6,241,701, and U.S. Pat. No. 6,233,482; electroporation may also be used for transfection of cells *in vitro* as described e.g. in US20070128708A1. Electroporation may also be utilized to deliver nucleic acids into cells *in vitro*. Accordingly, electroporation-mediated administration into cells of nucleic acids including expression constructs utilizing any of the many available devices and electroporation systems known to those of skill in the art presents an exciting new means for delivering an RNA of interest to a target cell.

5.7.1.1 Sources of T Cells

[0356] Prior to expansion and genetic modification, a source of T cells is obtained from a subject. The term “subject” is intended to include living organisms in which an immune response can be elicited (e.g., mammals). Examples of subjects include humans, dogs, cats, mice, rats, and transgenic species thereof. Preferably, subjects are human.

[0357] T cells can be obtained from a number of sources, including peripheral blood mononuclear cells, bone marrow, lymph node tissue, cord blood, thymus tissue, tissue from a site of infection, ascites, pleural effusion, spleen tissue, and tumors. In certain embodiments of the present disclosure, any number of T cell lines available in the art, may be used. In certain embodiments of the present disclosure, T cells can be obtained from a unit of blood collected from a subject using any number of techniques known to the skilled artisan, such as Ficoll™

separation. In one preferred embodiment, cells from the circulating blood of an individual are obtained by apheresis. The apheresis product typically contains lymphocytes, including T cells, monocytes, granulocytes, B cells, other nucleated white blood cells, red blood cells, and platelets. In one embodiment, the cells collected by apheresis may be washed to remove the plasma fraction and to place the cells in an appropriate buffer or media for subsequent processing steps. In one embodiment of the disclosure, the cells are washed with phosphate buffered saline (PBS). In an alternative embodiment, the wash solution lacks calcium and may lack magnesium or may lack many if not all divalent cations. Again, surprisingly, initial activation steps in the absence of calcium lead to magnified activation. As those of ordinary skill in the art would readily appreciate a washing step may be accomplished by methods known to those in the art, such as by using a semi-automated "flow-through" centrifuge (for example, the Cobe 2991 cell processor, the Baxter CytoMate, or the Haemonetics Cell Saver 5) according to the manufacturer's instructions. After washing, the cells may be resuspended in a variety of biocompatible buffers, such as, for example, Ca-free, Mg-free PBS, PlasmaLyte A, or other saline solution with or without buffer. Alternatively, the undesirable components of the apheresis sample may be removed and the cells directly resuspended in culture media.

[0358] In another embodiment, T cells are isolated from peripheral blood lymphocytes by lysing the red blood cells and depleting the monocytes, for example, by centrifugation through a PERCOLL™ gradient or by counterflow centrifugal elutriation. A specific subpopulation of T cells, such as CD3⁺, CD28⁺, CD4⁺, CD8⁺, CD45RA⁺ and CD45RO⁺ T cells, can be further isolated by positive or negative selection techniques. For example, in one embodiment, T cells are isolated by incubation with anti-CD3/anti-CD28 (*i.e.*, 3 x 28)-conjugated beads, such as DYNABEADS® M-450 CD3/CD28 T, for a time period sufficient for positive selection of the desired T cells. In one embodiment, the time period is about 30 minutes. In a further embodiment, the time period ranges from 30 minutes to 36 hours or longer and all integer values there between. In a further embodiment, the time period is at least 1, 2, 3, 4, 5, or 6 hours. In yet another preferred embodiment, the time period is 10 to 24 hours. In one preferred embodiment, the incubation time period is 24 hours. For isolation of T cells from patients with leukemia, use of longer incubation times, such as 24 hours, can increase cell yield. Longer incubation times may be used to isolate T cells in any situation where there are few T cells as compared to other cell types, such in isolating tumor infiltrating lymphocytes (TIL) from tumor tissue or from immunocompromised individuals. Further, use of longer incubation times can increase the efficiency of capture of CD8⁺ T cells. Thus, by simply shortening or lengthening the time T cells are allowed to bind to the CD3/CD28 beads and/or by increasing or decreasing the ratio of beads to T cells (as described further herein), subpopulations of T cells can be preferentially selected for or against at culture initiation or at other time points during the process. Additionally, by increasing or decreasing the ratio of anti-CD3 and/or anti-CD28 antibodies on the beads or other surface, subpopulations of T cells can be preferentially

selected for or against at culture initiation or at other desired time points. The skilled artisan would recognize that multiple rounds of selection can also be used in the context of this disclosure. In certain embodiments, it may be desirable to perform the selection procedure and use the “unselected” cells in the activation and expansion process. “Unselected” cells can also be subjected to further rounds of selection.

[0359] Enrichment of a T cell population by negative selection can be accomplished with a combination of antibodies directed to surface markers unique to the negatively selected cells. One method is cell sorting and/or selection via negative magnetic immunoadherence or flow cytometry that uses a cocktail of monoclonal antibodies directed to cell surface markers present on the cells negatively selected. For example, to enrich for CD4⁺ cells by negative selection, a monoclonal antibody cocktail typically includes antibodies to CD14, CD20, CD11b, CD16, HLA-DR, and CD8. In certain embodiments, it may be desirable to enrich for or positively select for regulatory T cells which typically express CD4⁺, CD25⁺, CD62L^{hi}, GITR⁺, and FoxP3⁺. Alternatively, in certain embodiments, T regulatory cells are depleted by anti-CD25 conjugated beads or other similar method of selection.

[0360] For isolation of a desired population of cells by positive or negative selection, the concentration of cells and surface (e.g., particles such as beads) can be varied. In certain embodiments, it may be desirable to significantly decrease the volume in which beads and cells are mixed together (i.e., increase the concentration of cells), to ensure maximum contact of cells and beads. For example, in one embodiment, a concentration of 2 billion cells/ml is used. In one embodiment, a concentration of 1 billion cells/ml is used. In a further embodiment, greater than 100 million cells/ml is used. In a further embodiment, a concentration of cells of 10, 15, 20, 25, 30, 35, 40, 45, or 50 million cells/ml is used. In yet another embodiment, a concentration of cells from 75, 80, 85, 90, 95, or 100 million cells/ml is used. In further embodiments, concentrations of 125 or 150 million cells/ml can be used. Using high concentrations can result in increased cell yield, cell activation, and cell expansion. Further, use of high cell concentrations allows more efficient capture of cells that may weakly express target antigens of interest, such as CD28-negative T cells, or from samples where there are many tumor cells present (i.e., leukemic blood, tumor tissue, etc.). Such populations of cells may have therapeutic value and would be desirable to obtain. For example, using high concentration of cells allows more efficient selection of CD8⁺ T cells that normally have weaker CD28 expression.

[0361] In a related embodiment, it may be desirable to use lower concentrations of cells. By significantly diluting the mixture of T cells and surface (e.g., particles such as beads), interactions between the particles and cells are minimized. This selects for cells that express high amounts of desired antigens to be bound to the particles. For example, CD4⁺ T cells express higher levels of CD28 and are more efficiently captured than CD8⁺ T cells in dilute

concentrations. In one embodiment, the concentration of cells used is 5×10^6 /ml. In other embodiments, the concentration used can be from about 1×10^5 /ml to 1×10^6 /ml, and any integer value in between.

[0362] In other embodiments, the cells may be incubated on a rotator for varying lengths of time at varying speeds at either 2-10° C. or at room temperature.

[0363] T cells for stimulation can also be frozen after a washing step. Washing not to be bound by theory, the freeze and subsequent thaw step provides a more uniform product by removing granulocytes and to some extent monocytes in the cell population. After the washing step that removes plasma and platelets, the cells may be suspended in a freezing solution. While many freezing solutions and parameters are known in the art and will be useful in this context, one method involves using PBS containing 20% DMSO and 8% human serum albumin, or culture media containing 10% Dextran 40 and 5% Dextrose, 20% Human Serum Albumin and 7.5% DMSO, or 31.25% Plasmalyte-A, 31.25% Dextrose 5%, 0.45% NaCl, 10% Dextran 40 and 5% Dextrose, 20% Human Serum Albumin, and 7.5% DMSO or other suitable cell freezing media containing for example, Hespan and PlasmaLyte A, the cells then are frozen to -80° C. at a rate of 1° per minute and stored in the vapor phase of a liquid nitrogen storage tank. Other methods of controlled freezing may be used as well as uncontrolled freezing immediately at -20° C. or in liquid nitrogen.

[0364] In certain embodiments, cryopreserved cells are thawed and washed as described herein and allowed to rest for one hour at room temperature prior to activation using the methods of the present disclosure.

[0365] Also contemplated in the context of the disclosure is the collection of blood samples or apheresis product from a subject at a time period prior to when the expanded cells as described herein might be needed. As such, the source of the cells to be expanded can be collected at any time point necessary, and desired cells, such as T cells, isolated and frozen for later use in T cell therapy for any number of diseases or conditions that would benefit from T cell therapy, such as those described herein. In one embodiment a blood sample or an apheresis is taken from a generally healthy subject. In certain embodiments, a blood sample or an apheresis is taken from a generally healthy subject who is at risk of developing a disease, but who has not yet developed a disease, and the cells of interest are isolated and frozen for later use. In certain embodiments, the T cells may be expanded, frozen, and used at a later time. In certain embodiments, samples are collected from a patient shortly after diagnosis of a particular disease as described herein but prior to any treatments. In a further embodiment, the cells are isolated from a blood sample or an apheresis from a subject prior to any number of relevant treatment modalities, including but not limited to treatment with agents such as natalizumab, efalizumab, antiviral agents, chemotherapy, radiation, immunosuppressive agents, such as cyclosporin, azathioprine, methotrexate, mycophenolate, and FK506, antibodies, or other

immunoablative agents such as CAMPATH, anti-CD3 antibodies, cytoxan, fludarabine, cyclosporin, FK506, rapamycin, mycophenolic acid, steroids, FR901228, and irradiation. These drugs inhibit either the calcium dependent phosphatase calcineurin (cyclosporine and FK506) or inhibit the p70S6 kinase that is important for growth factor induced signaling (rapamycin). (Liu *et al.*, Cell 66:807-815, 1991; Henderson *et al.*, Immun. 73:316-321, 1991; Bierer *et al.*, Curr. Opin. Immun. 5:763-773, 1993). In a further embodiment, the cells are isolated for a patient and frozen for later use in conjunction with (*e.g.*, before, simultaneously or following) bone marrow or stem cell transplantation or T cell ablative therapy using either chemotherapy agents such as, fludarabine, external-beam radiation therapy (XRT), cyclophosphamide.

[0366] In a further embodiment of the present disclosure, T cells are obtained from a patient directly following treatment. In this regard, it has been observed that following certain cancer treatments, in particular treatments with drugs that damage the immune system, shortly after treatment during the period when patients would normally be recovering from the treatment, the quality of T cells obtained may be optimal or improved for their ability to expand *ex vivo*. Likewise, following *ex vivo* manipulation using the methods described herein, these cells may be in a preferred state for enhanced engraftment and *in vivo* expansion. Thus, it is contemplated within the context of the present disclosure to collect blood cells, including T cells, dendritic cells, or other cells of the hematopoietic lineage, during this recovery phase. Further, in certain embodiments, mobilization (for example, mobilization with GM-CSF) and conditioning regimens can be used to create a condition in a subject wherein repopulation, recirculation, regeneration, and/or expansion of particular cell types is favored, especially during a defined window of time following therapy. Illustrative cell types include T cells, B cells, dendritic cells, and other cells of the immune system.

5.7.1.2 Activation and Expansion of T Cells

[0367] T cells are activated and expanded generally using methods as described, for example, in U.S. Pat. Nos. 6,352,694; 6,534,055; 6,905,680; 6,692,964; 5,858,358; 6,887,466; 6,905,681; 7,144,575; 7,067,318; 7,172,869; 7,232,566; 7,175,843; 5,883,223; 6,905,874; 6,797,514; 6,867,041; and U.S. Patent Application Publication No. 20060121005.

[0368] Generally, the T cells of the disclosure are expanded by contact with a surface having attached thereto an agent that stimulates a CD3/TCR complex associated signal and a ligand that stimulates a co-stimulatory molecule on the surface of the T cells. In particular, T cell populations may be stimulated as described herein, such as by contact with an anti-CD3 antibody, or antigen-binding fragment thereof, or an anti-CD2 antibody immobilized on a surface, or by contact with a protein kinase C activator (*e.g.*, bryostatin) in conjunction with a calcium ionophore. For co-stimulation of an accessory molecule on the surface of the T cells, a ligand that binds the accessory molecule is used. For example, a population of T cells can be contacted with an anti-CD3 antibody and an anti-CD28 antibody, under conditions appropriate

for stimulating proliferation of the T cells. To stimulate proliferation of either CD4⁺ T cells or CD8⁺ T cells, an anti-CD3 antibody and an anti-CD28 antibody. Examples of an anti-CD28 antibody include 9.3, B-T3, XR-CD28 (Diaclone, Besancon, France) can be used as can other methods commonly known in the art (Berg *et al.*, Transplant Proc. 30(8):3975-3977, 1998; Haanen *et al.*, J. Exp. Med. 190(9):13191328, 1999; Garland *et al.*, J. Immunol Meth. 227(1-2):53-63, 1999).

[0369] In certain embodiments, the primary stimulatory signal and the co-stimulatory signal for the T cell may be provided by different protocols. For example, the agents providing each signal may be in solution or coupled to a surface. When coupled to a surface, the agents may be coupled to the same surface (*i.e.*, in "cis" formation) or to separate surfaces (*i.e.*, in "trans" formation). Alternatively, one agent may be coupled to a surface and the other agent in solution. In one embodiment, the agent providing the co-stimulatory signal is bound to a cell surface and the agent providing the primary activation signal is in solution or coupled to a surface. In certain embodiments, both agents can be in solution. In another embodiment, the agents may be in soluble form, and then cross-linked to a surface, such as a cell expressing Fc receptors or an antibody or other binding agent which will bind to the agents. In this regard, see for example, U.S. Patent Application Publication Nos. 20040101519 and 20060034810 for artificial antigen presenting cells (APCs) that are contemplated for use in activating and expanding T cells in the present disclosure.

[0370] In one embodiment, the two agents are immobilized on beads, either on the same bead, *i.e.*, "cis," or to separate beads, *i.e.*, "trans." By way of example, the agent providing the primary activation signal is an anti-CD3 antibody or an antigen-binding fragment thereof and the agent providing the co-stimulatory signal is an anti-CD28 antibody or antigen-binding fragment thereof; and both agents are co-immobilized to the same bead in equivalent molecular amounts. In one embodiment, a 1:1 ratio of each antibody bound to the beads for CD4⁺ T cell expansion and T cell growth is used. In certain aspects of the present disclosure, a ratio of anti CD3:CD28 antibodies bound to the beads is used such that an increase in T cell expansion is observed as compared to the expansion observed using a ratio of 1:1. In one particular embodiment an increase of from about 1 to about 3 fold is observed as compared to the expansion observed using a ratio of 1:1. In one embodiment, the ratio of CD3:CD28 antibody bound to the beads ranges from 100:1 to 1:100 and all integer values there between. In one aspect of the present disclosure, more anti-CD28 antibody is bound to the particles than anti-CD3 antibody, *i.e.*, the ratio of CD3:CD28 is less than one. In certain embodiments of the disclosure, the ratio of anti CD28 antibody to anti CD3 antibody bound to the beads is greater than 2:1. In one particular embodiment, a 1:100 CD3:CD28 ratio of antibody bound to beads is used. In another embodiment, a 1:75 CD3:CD28 ratio of antibody bound to beads is used. In a further embodiment, a 1:50 CD3:CD28 ratio of antibody bound to beads is used. In another

embodiment, a 1:30 CD3:CD28 ratio of antibody bound to beads is used. In one preferred embodiment, a 1:10 CD3:CD28 ratio of antibody bound to beads is used. In another embodiment, a 1:3 CD3:CD28 ratio of antibody bound to the beads is used. In yet another embodiment, a 3:1 CD3:CD28 ratio of antibody bound to the beads is used.

[0371] Ratios of particles to cells from 1:500 to 500:1 and any integer values in between may be used to stimulate T cells or other target cells. As those of ordinary skill in the art can readily appreciate, the ratio of particles to cells may depend on particle size relative to the target cell. For example, small sized beads could only bind a few cells, while larger beads could bind many. In certain embodiments the ratio of cells to particles ranges from 1:100 to 100:1 and any integer values in-between and in further embodiments the ratio comprises 1:9 to 9:1 and any integer values in between, can also be used to stimulate T cells. The ratio of anti-CD3- and anti-CD28-coupled particles to T cells that result in T cell stimulation can vary as noted above, however certain preferred values include 1:100, 1:50, 1:40, 1:30, 1:20, 1:10, 1:9, 1:8, 1:7, 1:6, 1:5, 1:4, 1:3, 1:2, 1:1, 2:1, 3:1, 4:1, 5:1, 6:1, 7:1, 8:1, 9:1, 10:1, and 15:1 with one preferred ratio being at least 1:1 particles per T cell. In one embodiment, a ratio of particles to cells of 1:1 or less is used. In one particular embodiment, a preferred particle: cell ratio is 1:5. In further embodiments, the ratio of particles to cells can be varied depending on the day of stimulation. For example, in one embodiment, the ratio of particles to cells is from 1:1 to 10:1 on the first day and additional particles are added to the cells every day or every other day thereafter for up to 10 days, at final ratios of from 1:1 to 1:10 (based on cell counts on the day of addition). In one particular embodiment, the ratio of particles to cells is 1:1 on the first day of stimulation and adjusted to 1:5 on the third and fifth days of stimulation. In another embodiment, particles are added on a daily or every other day basis to a final ratio of 1:1 on the first day, and 1:5 on the third and fifth days of stimulation. In another embodiment, the ratio of particles to cells is 2:1 on the first day of stimulation and adjusted to 1:10 on the third and fifth days of stimulation. In another embodiment, particles are added on a daily or every other day basis to a final ratio of 1:1 on the first day, and 1:10 on the third and fifth days of stimulation. One of skill in the art will appreciate that a variety of other ratios may be suitable for use in the present disclosure. In particular, ratios will vary depending on particle size and on cell size and type.

[0372] In further embodiments of the present disclosure, the cells, such as T cells, are combined with agent-coated beads, the beads and the cells are subsequently separated, and then the cells are cultured. In an alternative embodiment, prior to culture, the agent-coated beads and cells are not separated but are cultured together. In a further embodiment, the beads and cells are first concentrated by application of a force, such as a magnetic force, resulting in increased ligation of cell surface markers, thereby inducing cell stimulation.

[0373] By way of example, cell surface proteins may be ligated by allowing paramagnetic beads to which anti-CD3 and anti-CD28 are attached (3 x 28 beads) to contact the T cells. In

one embodiment the cells (for example, 10^4 to 10^9 T cells) and beads (for example, DYNABEADS® M-450 CD3/CD28 T paramagnetic beads at a ratio of 1:1) are combined in a buffer, preferably PBS (without divalent cations such as, calcium and magnesium). Again, those of ordinary skill in the art can readily appreciate any cell concentration may be used. For example, the target cell may be very rare in the sample and comprise only 0.01% of the sample or the entire sample (*i.e.*, 100%) may comprise the target cell of interest. Accordingly, any cell number is within the context of the present disclosure. In certain embodiments, it may be desirable to significantly decrease the volume in which particles and cells are mixed together (*i.e.*, increase the concentration of cells), to ensure maximum contact of cells and particles. For example, in one embodiment, a concentration of about 2 billion cells/ml is used. In another embodiment, greater than 100 million cells/ml is used. In a further embodiment, a concentration of cells of 10, 15, 20, 25, 30, 35, 40, 45, or 50 million cells/ml is used. In yet another embodiment, a concentration of cells from 75, 80, 85, 90, 95, or 100 million cells/ml is used. In further embodiments, concentrations of 125 or 150 million cells/ml can be used. Using high concentrations can result in increased cell yield, cell activation, and cell expansion. Further, use of high cell concentrations allows more efficient capture of cells that may weakly express target antigens of interest, such as CD28-negative T cells. Such populations of cells may have therapeutic value and would be desirable to obtain in certain embodiments. For example, using high concentration of cells allows more efficient selection of CD8+ T cells that normally have weaker CD28 expression.

[0374] In one embodiment of the present disclosure, the mixture may be cultured for several hours (about 3 hours) to about 14 days or any hourly integer value in between. In another embodiment, the mixture may be cultured for 21 days. In one embodiment of the disclosure the beads and the T cells are cultured together for about eight days. In another embodiment, the beads and T cells are cultured together for 2-3 days. Several cycles of stimulation may also be desired such that culture time of T cells can be 60 days or more. Conditions appropriate for T cell culture include an appropriate media (*e.g.*, Minimal Essential Media or RPMI Media 1640 or, X-vivo 15, (Lonza)) that may contain factors necessary for proliferation and viability, including serum (*e.g.*, fetal bovine or human serum), interleukin-2 (IL-2), insulin, IFN- γ , IL-4, IL-7, GM-CSF, IL-10, IL-12, IL-15, TGF β , and TNF- α or any other additives for the growth of cells known to the skilled artisan. Other additives for the growth of cells include, but are not limited to, surfactant, plasmanate, and reducing agents such as N-acetyl-cysteine and 2-mercaptoethanol. Media can include RPMI 1640, AIM-V, DMEM, MEM, α -MEM, F-12, X-Vivo 15, and X-Vivo 20, Optimizer, with added amino acids, sodium pyruvate, and vitamins, either serum-free or supplemented with an appropriate amount of serum (or plasma) or a defined set of hormones, and/or an amount of cytokine(s) sufficient for the growth and expansion of T cells. Antibiotics, *e.g.*, penicillin and streptomycin, are included only in experimental cultures, not in cultures of cells that are to be infused into a subject. The target cells are maintained under

conditions necessary to support growth, for example, an appropriate temperature (e.g., 37° C.) and atmosphere (e.g., air plus 5% CO₂).

[0375] T cells that have been exposed to varied stimulation times may exhibit different characteristics. For example, typical blood or apheresed peripheral blood mononuclear cell products have a helper T cell population (T_H, CD4⁺) that is greater than the cytotoxic or suppressor T cell population (T_C, CD8⁺). Ex vivo expansion of T cells by stimulating CD3 and CD28 receptors produces a population of T cells that prior to about days 8-9 consists predominately of T_H cells, while after about days 8-9, the population of T cells comprises an increasingly greater population of T_C cells. Accordingly, depending on the purpose of treatment, infusing a subject with a T cell population comprising predominately of T_H cells may be advantageous. Similarly, if an antigen-specific subset of T_C cells has been isolated it may be beneficial to expand this subset to a greater degree.

[0376] Further, in addition to CD4 and CD8 markers, other phenotypic markers vary significantly, but in large part, reproducibly during the course of the cell expansion process. Thus, such reproducibility enables the ability to tailor an activated T cell product for specific purposes.

5.8 Compositions

[0377] The anti-glyco-cMET antibodies, fusion proteins, and/or anti-glyco-cMET ADCs of the disclosure may be in the form of compositions comprising the anti-glyco-cMET antibody, fusion protein and/or ADC and one or more carriers, excipients and/or diluents. The compositions may be formulated for specific uses, such as for veterinary uses or pharmaceutical uses in humans. The form of the composition (e.g., dry powder, liquid formulation, etc.) and the excipients, diluents and/or carriers used will depend upon the intended uses of the antibody, fusion protein and/or ADC and, for therapeutic uses, the mode of administration.

[0378] For therapeutic uses, the compositions may be supplied as part of a sterile, pharmaceutical composition that includes a pharmaceutically acceptable carrier. This composition can be in any suitable form (depending upon the desired method of administering it to a patient). The pharmaceutical composition can be administered to a patient by a variety of routes such as orally, transdermally, subcutaneously, intranasally, intravenously, intramuscularly, intratumorally, intrathecally, topically or locally. The most suitable route for administration in any given case will depend on the particular antibody and/or ADC, the subject, and the nature and severity of the disease and the physical condition of the subject. Typically, the pharmaceutical composition will be administered intravenously or subcutaneously.

[0379] Pharmaceutical compositions can be conveniently presented in unit dosage forms containing a predetermined amount of an anti-glyco-cMET antibody and/or anti-glyco-cMET ADC of the disclosure per dose. The quantity of antibody and/or ADC included in a unit dose

will depend on the disease being treated, as well as other factors as are well known in the art. Such unit dosages may be in the form of a lyophilized dry powder containing an amount of antibody and/or ADC suitable for a single administration, or in the form of a liquid. Dry powder unit dosage forms may be packaged in a kit with a syringe, a suitable quantity of diluent and/or other components useful for administration. Unit dosages in liquid form may be conveniently supplied in the form of a syringe pre-filled with a quantity of antibody and/or ADC suitable for a single administration.

[0380] The pharmaceutical compositions may also be supplied in bulk from containing quantities of ADC suitable for multiple administrations.

[0381] Pharmaceutical compositions may be prepared for storage as lyophilized formulations or aqueous solutions by mixing an antibody, fusion protein, and/or ADC having the desired degree of purity with optional pharmaceutically acceptable carriers, excipients or stabilizers typically employed in the art (all of which are referred to herein as "carriers"), *i.e.*, buffering agents, stabilizing agents, preservatives, isotoniifiers, non-ionic detergents, antioxidants, and other miscellaneous additives. See, Remington's Pharmaceutical Sciences, 16th edition (Osol, ed. 1980). Such additives should be nontoxic to the recipients at the dosages and concentrations employed.

[0382] Buffering agents help to maintain the pH in the range which approximates physiological conditions. They may be present at a wide variety of concentrations, but will typically be present in concentrations ranging from about 2 mM to about 50 mM. Suitable buffering agents for use with the present disclosure include both organic and inorganic acids and salts thereof such as citrate buffers (*e.g.*, monosodium citrate-disodium citrate mixture, citric acid-trisodium citrate mixture, citric acid-monosodium citrate mixture, *etc.*), succinate buffers (*e.g.*, succinic acid-monosodium succinate mixture, succinic acid-sodium hydroxide mixture, succinic acid-disodium succinate mixture, *etc.*), tartrate buffers (*e.g.*, tartaric acid-sodium tartrate mixture, tartaric acid-potassium tartrate mixture, tartaric acid-sodium hydroxide mixture, *etc.*), fumarate buffers (*e.g.*, fumaric acid-monosodium fumarate mixture, fumaric acid-disodium fumarate mixture, monosodium fumarate-disodium fumarate mixture, *etc.*), gluconate buffers (*e.g.*, gluconic acid-sodium glyconate mixture, gluconic acid-sodium hydroxide mixture, gluconic acid-potassium glyconate mixture, *etc.*), oxalate buffer (*e.g.*, oxalic acid-sodium oxalate mixture, oxalic acid-sodium hydroxide mixture, oxalic acid-potassium oxalate mixture, *etc.*), lactate buffers (*e.g.*, lactic acid-sodium lactate mixture, lactic acid-sodium hydroxide mixture, lactic acid-potassium lactate mixture, *etc.*) and acetate buffers (*e.g.*, acetic acid-sodium acetate mixture, acetic acid-sodium hydroxide mixture, *etc.*). Additionally, phosphate buffers, histidine buffers and trimethylamine salts such as Tris can be used.

[0383] Preservatives may be added to retard microbial growth, and can be added in amounts ranging from about 0.2%-1% (w/v). Suitable preservatives for use with the present disclosure

include phenol, benzyl alcohol, meta-cresol, methyl paraben, propyl paraben, octadecyldimethylbenzyl ammonium chloride, benzalconium halides (*e.g.*, chloride, bromide, and iodide), hexamethonium chloride, and alkyl parabens such as methyl or propyl paraben, catechol, resorcinol, cyclohexanol, and 3-pentanol. Isotonicifiers sometimes known as "stabilizers" can be added to ensure isotonicity of liquid compositions of the present disclosure and include polyhydric sugar alcohols, for example trihydric or higher sugar alcohols, such as glycerin, erythritol, arabitol, xylitol, sorbitol and mannitol. Stabilizers refer to a broad category of excipients which can range in function from a bulking agent to an additive which solubilizes the therapeutic agent or helps to prevent denaturation or adherence to the container wall. Typical stabilizers can be polyhydric sugar alcohols (enumerated above); amino acids such as arginine, lysine, glycine, glutamine, asparagine, histidine, alanine, ornithine, L-leucine, 2-phenylalanine, glutamic acid, threonine, *etc.*, organic sugars or sugar alcohols, such as lactose, trehalose, stachyose, mannitol, sorbitol, xylitol, ribitol, myoinositol, galactitol, glycerol and the like, including cyclitols such as inositol; polyethylene glycol; amino acid polymers; sulfur containing reducing agents, such as urea, glutathione, thiocetic acid, sodium thioglycolate, thioglycerol, α -monothioglycerol and sodium thio sulfate; low molecular weight polypeptides (*e.g.*, peptides of 10 residues or fewer); proteins such as human serum albumin, bovine serum albumin, gelatin or immunoglobulins; hydrophylic polymers, such as polyvinylpyrrolidone monosaccharides, such as xylose, mannose, fructose, glucose; disaccharides such as lactose, maltose, sucrose and trehalose; and trisaccharides such as raffinose; and polysaccharides such as dextran. Stabilizers may be present in amounts ranging from 0.5 to 10 wt % per wt of ADC.

[0384] Non-ionic surfactants or detergents (also known as "wetting agents") may be added to help solubilize the glycoprotein as well as to protect the glycoprotein against agitation-induced aggregation, which also permits the formulation to be exposed to shear surface stressed without causing denaturation of the protein. Suitable non-ionic surfactants include polysorbates (20, 80, *etc.*), poloxamers (184, 188 *etc.*), and pluronic polyols. Non-ionic surfactants may be present in a range of about 0.05 mg/mL to about 1.0 mg/mL, for example about 0.07 mg/mL to about 0.2 mg/mL.

[0385] Additional miscellaneous excipients include bulking agents (*e.g.*, starch), chelating agents (*e.g.*, EDTA), antioxidants (*e.g.*, ascorbic acid, methionine, vitamin E), and cosolvents.

5.9 Methods of Use

[0386] The anti-glyco-cMET antibody or binding fragments described herein can be used in various diagnostic and therapeutic methods. In some embodiments, a patient can be diagnosed with a cancer using any method as described herein (*e.g.*, as described in Section 5.9.1) and subsequently treated using any method as described herein (*e.g.*, as described in Section 5.9.2). The diagnostic methods described herein (*e.g.*, as described in Section 5.9.1)

can be utilized to monitor the patient's cancer status during or following cancer therapy (including but not limited to cancer therapy as described in Section 5.9.2).

5.9.1. Diagnostic Methods

[0387] The anti-glyco-cMET antibody or binding fragments (including immunoconjugates and labeled antibodies and binding fragments) can be used in diagnostic assays. For example, the antibodies and binding fragments can be employed in immunoassays, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays, including immunohistochemistry, enzyme-linked immunosorbent assay (ELISA), fluorescence-activated cell sorting (FACS), and Western blots.

[0388] An anti-glyco-cMET antibody or antigen-binding fragment of the disclosure can be used in a method of detecting a biomarker in a sample comprising one or more EVs (e.g., a liquid biopsy). In such embodiments, an EV surface biomarker is recognized by the anti-glyco-cMET antibody or antigen-binding fragment of the disclosure. Exemplary methods of detecting the biomarker include, but are not limited to, capture assays, immunoassays, such as immunoprecipitation; Western blot; ELISA; immunohistochemistry; immunocytochemistry; flow cytometry; and immuno-PCR. In some embodiments, an immunoassay can be a chemiluminescent immunoassay. In some embodiments, an immunoassay can be a high-throughput and/or automated immunoassay platform.

[0389] The anti-glyco-cMET antibody or binding fragments described herein also are useful for radiographic *in vivo* imaging, wherein an antibody labeled with a detectable moiety such as a radio-opaque agent or radioisotope is administered to a subject, preferably into the bloodstream, and the presence and location of the labeled antibody in the host is assayed. This imaging technique is useful in the staging and treatment of malignancies.

5.9.2. Therapeutic Methods

[0390] The anti-glyco-cMET antibody or binding fragments, fusion proteins, ADCs and CARs, and chimeric TCRs described herein are useful for treatment of glyco-cMET expressing cancers, including lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.

[0391] Thus, the disclosure provides anti-glyco-cMET antibodies, binding fragments, fusion proteins, ADCs, CARs, and chimeric TCRs as described herein for use as a medicament, for example for use in the treatment of cancer, e.g., any of the cancers identified in the previous paragraph, for use in a diagnostic assay, and for use in radiographic *in vivo* imaging. The disclosure further provides for the use of the anti-glyco-cMET antibodies, binding fragments, fusion proteins, ADCs, CARs and chimeric TCRs as described herein in the manufacture of a medicament, for example for the treatment of cancer, e.g., any of the cancers identified in the previous paragraph.

[0392] When using the CARs or chimeric TCRs of the disclosure for therapy, the therapeutic methods of the disclosure comprise administering to a subject with a glyco-cMET-expressing tumor an effective amount of a genetically modified cell engineered to express a CAR or chimeric TCR of the disclosure, for example a CAR as described in Section 5.3 or in numbered embodiments 689 to 724, a chimeric TCR as described in Section 5.4 or in numbered embodiments 735 to 834 or a MicAbody as described in Section 5.6. Methods of modifying cells, particularly T cells, to express a CAR or chimeric TCR, are described in Section 5.7.1.

[0393] When using the MicAbodies of the disclosure for therapy, the therapeutic methods of the disclosure comprise administering to a subject with a glyco-cMET-expressing tumor therapeutically effective amounts of a MicAbody of the disclosure, for example a MicAbody described in Section 5.6, and a genetically modified T-cell engineered to express a CAR comprising a NKG2D receptor capable of specifically binding the MicAbody.

5.10 cMET Peptides

[0394] Also provided are isolated cMET glycopeptides, or glyco-cMET peptides, comprising the amino acid PTKSFISGGSTITGVGKLN (SEQ ID NO:286), or a fragment thereof. In some embodiments, the cMET glycopeptide is glycosylated with O-linked GalNAc on the serine residue at amino acid position 10 and the threonine residue at amino acid position 11 of PTKSFISGGSTITGVGKLN (SEQ ID NO:286). In some embodiments the cMET glycopeptide comprises the amino acid PTKSFISGG**ST**ITGVGKLN (SEQ ID NO:285) or a fragment thereof, with O-linked GalNAc on the serine and threonine residues shown with bold and underlined text. Exemplary isolated cMET glycopeptides are described in numbered embodiments 894 to 920.

[0395] The present disclosure encompasses synthetic synthesis of the isolated cMET glycoproteins and recombinant methods for producing the isolated cMET glycoproteins.

[0396] In certain embodiments, the isolated cMET peptides are synthesized using a solid-phase peptide synthesis (SPPS) strategy. SPPS methods are known in the art. SPPS provides for the rapid assembly of a polypeptide through successive reactions of amino acid derivatives on a solid support. Through repeated cycles of alternating N-terminal deprotection and coupling reactions, successive amino acid derivatives are added to the polypeptide. In other embodiments, isolated cMET peptides are synthesized using a solution-phase peptide synthesis strategy. Solution-phase peptide synthesis methods are known in the art.

[0397] To ensure proper O-linked glycosylation with GalNAc on the serine at amino acid position 10 of SEQ ID NO:285 and the threonine at amino acid position 11 of SEQ ID NO:285, pre-synthesized glycosylated amino acids can be used in the elongation reactions.

[0398] Nucleic acid molecules encoding the isolated cMET glycopeptides, vectors comprising such nucleic acids, and host cells capable of producing the isolated cMET glycopeptides of the disclosure are provided. In certain aspects, the nucleic acid molecules encode, and the host cells are capable of expressing, the cMET glycopeptide as well as fusion proteins that include the cMET glycoproteins.

[0399] An isolated cMET glycopeptide of the disclosure can be prepared by recombinant expression in a host cell. To express a cMET glycopeptide recombinantly, a host cell is transfected with a recombinant expression vector carrying DNA encoding the glycopeptide such that the glycopeptide is expressed in the host cell and, optionally, secreted into the medium in which the host cells are cultured, from which medium the glycoproteins can be recovered (*i.e.*, isolated). Standard recombinant DNA methodologies are used to obtain a cMET glycoprotein gene, incorporate the gene into recombinant expression vectors and introduce the vectors into host cells, such as those described in *Molecular Cloning; A Laboratory Manual, Second Edition* (Sambrook, Fritsch and Maniatis (eds), Cold Spring Harbor, N. Y., 1989), 122 *Current Protocols in Molecular Biology* (Ausubel, F. M. *et al.*, eds., Greene Publishing Associates, 1989) and in U.S. Pat. No. 4,816,397.

[0400] It is possible to express the cMET glycoproteins of the disclosure in either prokaryotic or eukaryotic host cells. In certain embodiments, expression of cMET glycoprotein is performed in eukaryotic cells, e.g., mammalian host cells. To produce the isolated cMET glycoproteins of the disclosure, a host cell is selected based on its ability to glycosylate the serine at amino acid position 10 of SEQ ID NO:285 and the threonines at amino acid positions 10 and 11 of SEQ ID NO:285. An exemplary host cell is the COSMC HEK293 cell.

5.10.1. cMET Peptide Compositions

[0401] The cMET glycopeptides of the disclosure may be in the form of compositions comprising the cMET glycopeptide and one or more carriers, excipients, diluents and/or adjuvants. The compositions may be formulated for specific uses, such as for veterinary uses or pharmaceutical uses in humans. The form of the composition (e.g., dry powder, liquid formulation, etc.) and the excipients, diluents and/or carriers used will depend upon the intended uses of the cMET glycopeptide and, for therapeutic uses, the mode of administration.

[0402] For therapeutic uses, the compositions may be supplied as part of a sterile, pharmaceutical composition that includes a pharmaceutically acceptable carrier and/or a pharmaceutically acceptable adjuvant. This composition can be in any suitable form (depending upon the desired method of administering it to a patient). The pharmaceutical composition can be administered to a patient by a variety of routes such as orally, transdermally, subcutaneously, intranasally, intravenously, intramuscularly, intratumorally, intrathecally, topically or locally. The most suitable route for administration in any given case will depend on the particular cMET glycopeptide to be administered, the subject, and the nature and severity of

the disease and the physical condition of the subject. Typically, the pharmaceutical composition will be administered intravenously or subcutaneously.

[0403] Pharmaceutical compositions can be conveniently presented in unit dosage forms containing a predetermined amount of an cMET glycopeptide of the disclosure per dose. The quantity of cMET glycopeptide included in a unit dose will depend on the disease being treated, as well as other factors as are well known in the art. Such unit dosages may be in the form of a lyophilized dry powder containing an amount of cMET glycopeptide suitable for a single administration, or in the form of a liquid. Dry powder unit dosage forms may be packaged in a kit with a syringe, a suitable quantity of diluent and/or other components useful for administration. Unit dosages in liquid form may be conveniently supplied in the form of a syringe pre-filled with a quantity of cMET glycopeptide suitable for a single administration.

[0404] The pharmaceutical compositions may also be supplied in bulk form containing quantities of cMET glycopeptide suitable for multiple administrations.

[0405] Pharmaceutical compositions may be prepared for storage as lyophilized formulations or aqueous solutions by mixing a cMET glycopeptide having the desired degree of purity with optional pharmaceutically acceptable carriers, excipients, adjuvants or stabilizers typically employed in the art (all of which are referred to herein as "carriers"), *i.e.*, buffering agents, stabilizing agents, preservatives, isotonicifiers, non-ionic detergents, antioxidants, and other miscellaneous additives. See, Remington's Pharmaceutical Sciences, 16th edition (Osol, ed. 1980). Such additives should be nontoxic to the recipients at the dosages and concentrations employed.

[0406] In some embodiments, the composition includes one or more pharmaceutically acceptable adjuvants. Adjuvants include, for example, aluminum salts (e.g., amorphous aluminum hydroxyphosphate sulfate (AAHS), aluminum hydroxide, aluminum phosphate, potassium aluminum sulfate (Alum)), dsRNA analogues, lipid A analogues, flagellin, imidazoquinolines, CpG ODN, saponins (e.g., QS21), C-type lectin ligands (e.g., TDB), CD1d ligands (α-galactosylceramide), M F59, AS01, AS02, AS03, AS04, AS15, AF03, GLA-SE, IC31, CAF01, and virosomes. Other adjuvants known in the art, including chemical adjuvants, genetic adjuvants, protein adjuvants, and lipid adjuvants, can also be included in the compositions.

[0407] Buffering agents help to maintain the pH in the range which approximates physiological conditions. They may be present at a wide variety of concentrations, but will typically be present in concentrations ranging from about 2 mM to about 50 mM. Suitable buffering agents for use with the present disclosure include both organic and inorganic acids and salts thereof such as citrate buffers (e.g., monosodium citrate-disodium citrate mixture, citric acid-trisodium citrate mixture, citric acid-monosodium citrate mixture, etc.), succinate buffers (e.g., succinic acid-monosodium succinate mixture, succinic acid-sodium hydroxide mixture, succinic acid-disodium succinate mixture, etc.), tartrate buffers (e.g., tartaric acid-sodium tartrate mixture, tartaric acid-

potassium tartrate mixture, tartaric acid-sodium hydroxide mixture, etc.), fumarate buffers (e.g., fumaric acid-monosodium fumarate mixture, fumaric acid-disodium fumarate mixture, monosodium fumarate-disodium fumarate mixture, etc.), gluconate buffers (e.g., gluconic acid-sodium glyconate mixture, gluconic acid-sodium hydroxide mixture, gluconic acid-potassium glyconate mixture, etc.), oxalate buffer (e.g., oxalic acid-sodium oxalate mixture, oxalic acid-sodium hydroxide mixture, oxalic acid-potassium oxalate mixture, etc.), lactate buffers (e.g., lactic acid-sodium lactate mixture, lactic acid-sodium hydroxide mixture, lactic acid-potassium lactate mixture, etc.) and acetate buffers (e.g., acetic acid-sodium acetate mixture, acetic acid-sodium hydroxide mixture, etc.). Additionally, phosphate buffers, histidine buffers and trimethylamine salts such as Tris can be used.

[0408] Preservatives may be added to retard microbial growth, and can be added in amounts ranging from about 0.2%-1% (w/v). Suitable preservatives for use with the present disclosure include phenol, benzyl alcohol, meta-cresol, methyl paraben, propyl paraben, octadecyldimethylbenzyl ammonium chloride, benzalconium halides (e.g., chloride, bromide, and iodide), hexamethonium chloride, and alkyl parabens such as methyl or propyl paraben, catechol, resorcinol, cyclohexanol, and 3-pentanol. Isotonicifiers sometimes known as "stabilizers" can be added to ensure isotonicity of liquid compositions of the present disclosure and include polyhydric sugar alcohols, for example trihydric or higher sugar alcohols, such as glycerin, erythritol, arabitol, xylitol, sorbitol and mannitol. Stabilizers refer to a broad category of excipients which can range in function from a bulking agent to an additive which solubilizes the therapeutic agent or helps to prevent denaturation or adherence to the container wall. Typical stabilizers can be polyhydric sugar alcohols (enumerated above); amino acids such as arginine, lysine, glycine, glutamine, asparagine, histidine, alanine, ornithine, L-leucine, 2-phenylalanine, glutamic acid, threonine, etc., organic sugars or sugar alcohols, such as lactose, trehalose, stachyose, mannitol, sorbitol, xylitol, ribitol, myoinositol, galactitol, glycerol and the like, including cyclitols such as inositol; polyethylene glycol; amino acid polymers; sulfur containing reducing agents, such as urea, glutathione, thiocetic acid, sodium thioglycolate, thioglycerol, α -monothioglycerol and sodium thio sulfate; low molecular weight polypeptides (e.g., peptides of 10 residues or fewer); proteins such as human serum albumin, bovine serum albumin, gelatin or immunoglobulins; hydrophylic polymers, such as polyvinylpyrrolidone monosaccharides, such as xylose, mannose, fructose, glucose; disaccharides such as lactose, maltose, sucrose and trehalose; and trisaccharides such as raffinose; and polysaccharides such as dextran. Stabilizers may be present in amounts ranging from 0.5 to 10 wt % per wt of cMET peptide.

[0409] Non-ionic surfactants or detergents (also known as "wetting agents") may be added to help solubilize the glycoprotein as well as to protect the glycoprotein against agitation-induced aggregation, which also permits the formulation to be exposed to shear surface stressed without causing denaturation of the protein. Suitable non-ionic surfactants include polysorbates

(20, 80, etc.), poloxamers (184, 188 etc.), and pluronic polyols. Non-ionic surfactants may be present in a range of about 0.05 mg/mL to about 1.0 mg/mL, for example about 0.07 mg/mL to about 0.2 mg/mL.

[0410] Additional miscellaneous excipients include bulking agents (e.g., starch), chelating agents (e.g., EDTA), antioxidants (e.g., ascorbic acid, methionine, vitamin E), and cosolvents.

[0411] Exemplary cMET peptide compositions of the disclosure are described in numbered embodiments 921 and 922.

5.10.2. Methods of Using cMET Peptides

[0412] The cMET peptides described herein can be used in the production of antibodies against a tumor-associated form of cMET. The cMET peptide can be administered to an animal. The amount of peptide administered can be effective to cause the animal to produce antibodies against the peptide. As used herein, "animal" refers to multicellular eukaryotic organism from the biological kingdom Animalia. In some embodiments, the animal is a mammal. In some embodiments, the animal is a mouse or a rabbit. Resulting antibodies can then be collected from the animal. The cMET peptide can be administered as purified peptide or as part of a composition provided herein.

[0413] The cMET peptides described herein can be used to elicit an immune response against a tumor-associated form of cMET. The cMET peptide can be administered to an animal in an amount effective to cause the animal to mount an immune response (e.g., produce antibodies) against the peptide.

[0414] Exemplary methods for using the cMET peptides of the disclosure are described in numbered embodiments 923 to 926.

6. EXAMPLES

6.1 Example 1: Identification and Characterization of Anti-Glyco-cMET Antibodies

6.1.1. Overview

[0415] Glycans are essential membrane components and neoplastic transformation of human cells is virtually always associated with aberrant glycosylation of proteins and lipids. There are several types of protein glycosylation, including N-glycosylation and many types of O-glycosylation, but one of the most diverse types is the mucin type GalNAc type O-glycosylation (hereafter called O-glycosylation). Cancer associated changes in O-glycans are particularly interesting and the most frequently observed aberrant glycophenotype is expression of the most immature truncated O-glycan structures designated Tn (GalNAc α 1-O-Ser/Thr), STn (NeuAc α 2-6GalNAc α 1-O-Ser/Thr), and T (Gal β 1-3GalNAc α 1-O-Ser/Thr) antigens. Truncated O-glycans are observed on almost all epithelial cancer cells and strongly correlated with poor prognosis. In addition, it is becoming increasingly clear that glycans also have pivotal roles in

cancer development, with truncated O-glycans affecting differentiation, cell-cell and cell-matrix interactions, directly inducing oncogenic features in predisposed cells.

[0416] The inventors have identified cMET glycopeptide epitopes in human cancer cells and used the defined glyco-peptides to develop cancer specific anti-glyco-cMET monoclonal antibodies.

6.1.2. Materials and Methods

6.1.2.1 Synthesis of Tn cMET glycopeptide

[0417] The cMET glycopeptide, PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285) , with O-linked GalNAc on the serine and threonine residues shown with bold and underlined text was synthesized using a standard Fmoc peptide synthesis strategy. Pre-synthesized glycosylated amino acids were coupled to the elongating peptide at specific locations using solid or solution phase peptide chemistry in a stepwise fashion. After completing the full sequence and removing all protecting groups, the resulting glycopeptide was purified by high-performance liquid chromatography (HPLC) and characterized by mass spectrometry (electrospray ionization in positive mode).

6.1.2.2 Synthesis of recombinant Tn-glycosylated cMET

[0418] 1×10^8 COSMC KO HEK293 cells in 30 mL Opti-MEM were transfected using 30 μ g of a plasmid encoding his-tagged human cMET and 60 μ L 293fectin™ Transfection Reagent (Gibco). Following 48 hours of culture, the cells were spun down and the his-tagged recombinant cMET protein was purified from the supernatant using a 50% Ni-NTA agarose slurry column (Invitrogen), eluting with 250mM imidazole. To increase purity, this purification step was repeated. The recombinant SC-cMET protein was concentrated in PBS using Amicon Ultra centrifugal filters.

6.1.2.3 Mouse Immunization Protocol

[0419] Female Balb/c mice were immunized subcutaneously with the Tn-glycosylated cMET glycopeptide conjugated to KLH (keyhole limpet hemocyanin) through a maleimide linker. The mice were immunized on days 0, 14, and 35 with 50 μ g, 45 μ g, and 45 μ g of KLH-glycopeptide, respectively. The first immunization used Freund's complete adjuvant. All subsequent immunizations used Freund's incomplete adjuvant. On Day 45, tail bleeds were evaluated for polyclonal response. On day 56 or after, mice to be fused were boosted with 15 μ g of KLH-glycopeptide in Freund's incomplete adjuvant 3 to 5 days before hybridoma fusion. Splenocytes from mice were fused with SP2/0-Ag14 (ATCC, cat# CRL-1581) myeloma cells using the Electro Cell Manipulator (ECM2001) from BTX Harvard Apparatus. Hybridomas were seeded in 96-well plates, cultured, scaled, and evaluated and selected for specificity towards cMET-Tn using ELISA, FLOW cytometry, and immunofluorescence to obtain monoclonal antibodies having specificity for cMET-Tn.

6.1.2.4 Rabbit Immunization Protocol

[0420] New Zealand White Rabbits were immunized with Tn-glycosylated cMET glycopeptide conjugated to KLH through a maleimide linker. The rabbits were immunized on days 0, 28, and 47 with 50-200 ug of KLH-glycopeptide. On day 58, bleeds were evaluated for polyclonal response. On day 66 or later, rabbits of interest were boosted with 50-200 µg of KLH-glycopeptide 12 days before terminal bleed for B cell harvest. B cells were enriched, seeded in 96-well plates, cultured, and evaluated for specificity towards cMET-Tn using ELISA and FLOW cytometry. B cells of interests were cloned, expressed, and screened on ELISA, FLOW cytometry, and Immunofluorescences to obtain monoclonal antibodies having specificity to cMET-Tn.

6.1.2.5 ELISA

[0421] 96-well Corning high bind microplates (Fisher) were coated overnight at 4 °C with various concentrations of protein, peptide, or glycopeptide in 0.2 M bicarbonate-carbonate buffer (pH 9.4). The plates were then blocked for 1 hour at room temperature with Phosphate-buffered saline (PBS) (pH 7.4) containing 2.5% BSA. Contents of the plate were discarded and purified antibody, or hybridoma supernatants, or blood serum for polyclonal responses, were added at various concentrations and incubated for two hours at room temperature. Plates were washed with tris-buffered saline with 0.05% Tween-20 and then incubated for 1 hour at room temperature with a 1:3000 dilution of HRP conjugated goat anti-mouse IgG Fc γ (Sigma). The plates were washed again and developed with TMB chromogen substrate. After proper development (approximately 2-3 min), the reaction was stopped with 0.2 N H₂SO₄ and the absorbance was read at 450 nm. Data was analysed in GraphPad Prism Software.

6.1.2.6 Flow Cytometry

[0422] Adherent cells were dissociated with TrypLE select (Gibco) and washed from flask surface with cell culture media (RPMI w/ L-glutamine, 1% PenStrep, 1x Glutamax & 10% FBS). Cells were washed several times by centrifugation at 300*g for 5 min at 4 °C followed by resuspension in PBS with 1% BSA (PBS/1%BSA). Cells were resuspended between 5x10⁵ cells/ml to 2x10⁶ cell/ml and then distributed into a 96 well U-bottom plate. Diluted commercial antibody (0.25-2 µg /ml), or hybridoma supernatants, or blood serum for polyclonal responses, were added to cells and incubated for 1 hr on ice. Following several washes with PBS/1% BSA, cells were incubated for 30 min on ice with a 1:1600 dilution of AlexaFluor647 conjugated F(ab)₂ goat anti-mouse IgG Fc γ (JacksonImmunoResearch). Cells were washed again with PBS/1% BSA and then fixed in 1% formaldehyde in PBS/1% BSA. Cells were analysed on either a 2 or 4 laser Attune NXT flow cytometer. Data was processed in FlowJo Software.

6.1.2.7 Immunofluorescence

[0423] Cells were seeded to 50% confluency in glass bottom 96 well plate (Greiner Bio) and incubated 12-18 hours at 37 °C 5% CO₂. Following overnight growth, media from slides was

removed and cells were fixed with 4% formaldehyde in PBS (pH 7.4) for 10 min at room temperature. Slides were washed in PBS. Diluted commercial antibody (1-4 µg /ml), or hybridoma supernatants, or blood serum for polyclonal responses, were added to the slides and the slides were incubated overnight at 4 °C. The slides were washed in PBS and stained with a 1:800 dilution of AlexaFluor488 conjugated F(ab)₂ rabbit anti-mouse IgG (H+L) (Invitrogen) for 45 min at room temperature. The slides were washed in PBS and incubated with 4 µg /ml DAPI. DAPI was removed and PBS was added prior to imaging on a Nikon Ti LTTL microscope.

6.1.3. Results

6.1.3.1 Glycopeptide Specific antibodies to Tn-cMET

[0424] Glycopeptide reactive antibodies were generated using the Tn-glycosylated cMET glycopeptide. Mouse antibodies 15C4, 8H3, and 16E12, and rabbit antibodies 14E9, 19H2, and 39A3 showed superior selectivity. These 6 antibodies were moved forward with further characterization.

5.1.3.2 Characterization of mAbs 14E9, 19H2, and 39A3 binding specificity

[0425] To characterize the binding specificity of 14E9, 19H2, and 39A3, ELISA against Tn-glycosylated cMET and Tn-glycosylated Syndecan2 peptides was performed. It was found that in the context of ELISA, all 3 rabbit cMET mAbs only reacted with the Tn-glycosylated cMET peptide (FIG. 1).

6.2 Example 2: Functional characterization of 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 antibodies by Octet and Biacore

6.2.1. Overview

[0426] 15C4, 8H3, and 16E12 were characterized by Biacore to test the reactivity of anti-cMET mAbs to titrated cMET peptides. 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 were also characterized by Octet to test the reactivity of the anti-cMET mAbs to peptides with different glycosylated residues (including a non-glycosylated peptide) as shown in Table 6.

Table 6	
Peptide	Sequence (Bold and Underlined= GaINAc Site)
cMET-Tn	PTKSFISGG <u>S</u> TITGVGKNLN (SEQ ID NO:285)
cMET-Tn (S)	PTKSFISGG <u>S</u> TITGVGKNLN (SEQ ID NO:334)
cMET-Tn (T)	PTKSFISGG <u>T</u> TITGVGKNLN (SEQ ID NO:335)
cMET	PTKSFISGGSTITGVGKNLN (SEQ ID NO:286)

6.2.2. Materials and Methods

6.2.2.1 Surface Plasmon Resonance

[0427] Antibody affinity assays can be carried out using surface plasmon resonance (e.g., using a Biacore system (Cytiva)). In a surface plasmon resonance assay, one or more antibodies can be immobilized onto a biosensor and presented with an analyte (e.g., the cMET-Tn peptide Biotin- PTKSFISGGSTITGVGKNLN (the amino acid portion of which is SEQ ID NO:285; bold and underlined residues indicate GalNAc glycosylation sites) or a negative control analyte such as an unglycosylated cMET peptide (Biotin- PTKSFISGGSTITGVGKNLN (the amino acid portion of which is SEQ ID NO:286)). The antibodies are contacted with different concentrations of the analyte, for example concentrations of 2.5 nM, 7.4 nM, 22 nM, 66 nM and 200 nM. Affinity is measured using multi-cycle kinetics in triplicate for each analyte concentration, with 1 min association and 5 min dissociation. When comparing the binding affinities of two antibodies, the same concentration of both antibodies was used (e.g., measured using a 1 μ M concentration of each antibody). The affinity is determined by fitting the binding curve to a specific model: kinetic fit (1:1 model) or if applicable heterogenous ligand binding model. The error (>95% confidence) was calculated by how close the generated curve matches the model.

6.2.2.2 Bio-Layer Interferometry (Octet)

[0428] Antibody affinity and epitope binning of monoclonal antibodies can be assessed against specific antigens using bio-layer interferometry (BLI). In a BLI assay, the antigen can be immobilized onto a biosensor (e.g., the cMET-Tn peptide Biotin- PTKSFISGGSTITGVGKNLN (the amino acid portion of which is SEQ ID NO:285) or a negative control analyte such as an unglycosylated cMET peptide (Biotin- PTKSFISGGSTITGVGKNLN (the amino acid portion of which is SEQ ID NO:286)) and presented to one antibody for affinity measurements or two competing antibodies in tandem (or consecutive steps) for epitope binning. The binding to non-overlapping epitopes occurs if saturation with the first antibody does not block the binding of the second antibody. The affinity is determined by fitting the binding curve to a specific model: a 1:1 monovalent model or a 2:1 bivalent model. The error (>95% confidence) is calculated by how close the generated curve matches the model.

6.2.2.1 Flow Cytometry

[0429] Adherent cells were dissociated with TrypLE select (Gibco) and washed from the flask surface with cell culture media (RPMI w/ L-glutamine, 1% PenStrep, & 10% FBS). Cells were washed several times by centrifugation at 300*g for 5 min at 4 °C followed by resuspension in PBS with 1% BSA (PBS/1% BSA). Cells were resuspended between 5x10⁵ cells/ml to 2x10⁶ cell/ml and then distributed into a 96 well U-bottom plate. Diluted commercial antibody (0.25-2 μ g/ml), or hybridoma supernatants, or blood serum for polyclonal responses, were added to cells and incubated for 1 hr on ice. Following several washes with PBS/1% BSA, cells were incubated for 30 min on ice with a 1:1600 dilution of AlexaFluor647 conjugated F(ab)₂ goat anti-

mouse IgG Fc γ (JacksonImmunoResearch). Cells were washed again with PBS/1% BSA and then fixed in 1% formaldehyde in PBS/1% BSA. Cells were analysed on either a 2 or 4 laser Attune NXT flow cytometer. Data was processed in FlowJo Software.

6.2.2.2 Immunofluorescence

[0430] Cells were seeded to 50% confluency in glass chamber slides (nunc) and incubated 12-18 hours at 37 °C, 5% CO₂. Following overnight growth, media from slides was removed and cells were fixed with 4% formaldehyde in PBS (pH 7.4) for 10 min at room temperature. Slides were washed in PBS and blocked with PBS/2% BSA for 1 hour. Diluted commercial antibody (1-4 μ g/ml), or hybridoma supernatants, or blood serum for polyclonal responses, were added to the slides and the slides were incubated overnight at 4 °C. The slides were washed in PBS and stained with a 1:800 dilution of AlexaFluor488 conjugated F(ab)₂ rabbit anti-mouse IgG (H+L) (Invitrogen) for 45 min at room temperature. The slides were washed in PBS and mounted using Prolong Gold Antifade Mountant with DAPI (Thermofisher) and examined using an Olympus FV3000 confocal microscope.

6.2.3. Results

6.2.3.1 Glycopeptide specific antibodies to Tn-cMET

[0431] Glycopeptide reactive antibodies were generated using the Tn-glycosylated cMET glycopeptide. Antibodies generated using the cMET glycopeptide, including 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3, proved superior in selectivity.

6.2.3.2 Binding specificities of mAb 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3

[0432] The affinities of 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 against the various cMET glycopeptides were determined by Biacore and Octet. Table 7 summarizes dissociation constants (K_d) for 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 against different glycoforms of cMET peptide, as well as unglycosylated cMET and MUC1-Tn.

Table 7: Dissociation Constants for monoclonal cMET antibodies								
Antibody	Affinity (Biacore)			Apparent Affinity (Octet)				
	cMET-Tn	cMET	MUC1-Tn	cMET-Tn	cMET-Tn (S)	cMET-Tn (T)	cMET	MUC1-Tn
15C4	7.00 nM	>400 nM	>400 nM	2.83 nM	>10 μ M	>10 μ M	>10 μ M	>10 μ M
8H3	6.26 nM	>400 nM	>400 nM	7 nM	>10 μ M	>10 μ M	>10 μ M	>10 μ M
16E12	1.43 nM	>400 nM	>400 nM	19.2 nM	>10 μ M	>1.6 μ M	>10 μ M	2 μ M

14E9	NA	NA	NA	1.5 nM	NA	NA	>10 uM	>10 uM
19H2	NA	NA	NA	>10 uM	NA	NA	>10 uM	>10 uM
39A3	NA	NA	NA	52 nM	NA	NA	>10 uM	>10 uM
NA indicates affinity was not measured using the given technology, ND (not determined) indicates the affinity was below the detection limits.								

[0433] To further assess the specificities of 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 in a more natural conformational context, 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 was used to stain A549 cells for flow cytometry and A549 and T47D cells for immunofluorescence. T47D and A549 cell lines is inherently Tn-negative but can be induced to express the Tn-antigen by KO of the COSMC chaperone. When using 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3 to stain for flow cytometry, it was found that each antibody selectively stained COSMC KO A549 cells but not their wildtype counterpart, despite both cells staining positive for CMET (FIGS. 2A-3B-5). In agreement with these results, immunofluorescence showed that only CMET⁺ Tn⁺ T47D COSMC KO and CMET⁺ Tn⁺ T47D COSMC KO A549 cells stained with 15C4, 8H3, 16E12, 14E9, 19H2, or 39A3, whereas CMET⁺ Tn⁻ T47D WT cells did not (FIGS. 4A-4C).

6.3 Example 3: Tissue expression of Tn-glycosylated CMET epitope recognized by 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3

6.3.1. Overview

[0434] 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 were characterized by immunohistochemistry on various normal and cancer tissues.

6.3.2. Materials and methods

[0435] Paraffin embedded tissue micro arrays (TMAs) or tissue sections were de-paraffinized with xylene and ethanol, following antigen retrieval with citrate buffer (pH 6.0) and heated in a microwave for 18 min. TMAs were obtained from USBIOMAX and stained with Ultra Vison Quanto Detection System HRP DAB. Briefly, TMAs were washed in TBS, incubated with mAb supernatant for 2 hours. After wash in TBS x 2, the TMAs was incubated with Primary Antibody Amplifier Quanto for 10 min. After wash in TBS, TMAs were incubated with HRP polymer quanto (10 min) followed by DAB chromogen. Slides were counterstained with hematoxylin, were dehydrated, and mounted.

6.3.3. Results

[0436] When staining formalin-fixed paraffin embedded tissue sections for immunohistochemistry, positive staining was observed with 15C4, 8H3, 16E12, 4E9, 19H2, and 39A3 with staining in 8/8 colon (FIGS. 5A-5B) and 8H3 showed positive cellular surface staining on ovarian cancer (17%), pancreatic cancer (13%), lung cancer (14%), and cholangiocarcinoma (11%; FIGS. 6A-1-6A-2). This staining pattern correlated with staining for

normal CMET expression, showing that CMET expression in these carcinomas predicted reactivity to 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3. Importantly, no reactivity was observed on surface of cells when using 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 to stain healthy adjacent tissues (FIGS. 5A-5B and FIGS. 6B-1 to 6B-2). 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3 were found to positively react with several cancer tissue sections, but not their healthy counterparts.

[0437] The identity of each tissue in the TMA is set forth in Tables 8-14.

TABLE 8 (T051b; Colon cancer array)									
Position	No.	Age	Sex	Organ/ Anatomic Site	Pathology diagnosis	TNM	Grade	Stage	Type
A1	1	F	58	Colon	Adenocarcinoma	T3N1M0	2	IIIB	Malignant
A2	2	F	58	Colon	Adenocarcinoma	T3N1M0	2	IIIB	Malignant
A3	3	F	72	Colon	Adenocarcinoma	T3N0M0	2	IIA	Malignant
A4	4	F	72	Colon	Adenocarcinoma (sparse)	T3N0M0	2	IIA	Malignant
A5	5	F	58	Colon	Adenocarcinoma	T3N1M0	2	IIIB	Malignant
A6	6	F	58	Colon	Adenocarcinoma	T3N1M0	2	IIIB	Malignant
A7	7	F	72	Colon	Adenocarcinoma	T3N0M0	2	IIA	Malignant
A8	8	F	72	Colon	Adenocarcinoma	T3N0M0	2	IIA	Malignant
B1	9	M	45	Colon	Mucinous adenocarcinoma	T4N2M0	3	IIIC	Malignant
B2	10	M	45	Colon	Mucinous adenocarcinoma	T4N2M0	3	IIIC	Malignant
B3	11	M	22	Colon	Mucinous adenocarcinoma	T3N1M0	3	IIIB	Malignant
B4	12	M	22	Colon	Mucinous adenocarcinoma	T3N1M0	3	IIIB	Malignant
B5	13	M	45	Colon	Mucinous adenocarcinoma	T4N2M0	3	IIIC	Malignant
B6	14	M	45	Colon	Mucinous adenocarcinoma	T4N2M0	3	IIIC	Malignant
B7	15	M	22	Colon	Mucinous adenocarcinoma	T3N1M0	3	IIIB	Malignant
B8	16	M	22	Colon	Mucinous adenocarcinoma	T3N1M0	3	IIIB	Malignant
C1	17	M	48	Colon	Cancer adjacent colon tissue	-	-	-	AT
C2	18	M	48	Colon	Cancer adjacent colon tissue	-	-	-	AT
C3	19	M	52	Colon	Cancer adjacent colon tissue	-	-	-	AT
C4	20	M	52	Colon	Cancer adjacent colon tissue	-	-	-	AT
C5	21	M	48	Colon	Cancer adjacent colon tissue	-	-	-	AT
C6	22	M	48	Colon	Cancer adjacent colon tissue	-	-	-	AT
C7	23	M	52	Colon	Cancer adjacent colon tissue	-	-	-	AT
C8	24	M	52	Colon	Cancer adjacent colon tissue	-	-	-	AT

TABLE 9 (BN114; normal tissue array)						
Position	No.	Age	Sex	Organ/Anatomic Site	Pathology diagnosis	TNM
A1	1	2	F	Liver	Normal liver tissue	normal
A2	2	50	F	Liver	Normal liver tissue	normal
A3	3	14	F	Liver	Normal liver tissue	normal
A4	4	35	F	Liver	Normal liver tissue	normal
A5	5	24	M	Liver	Normal liver tissue	normal
A6	6	21	F	Liver	Normal liver tissue	normal
A7	7	Fetus	F	Liver	Normal fetal liver tissue	normal
A8	8	35	M	Liver	Normal liver tissue	normal
B1	9	35	M	Liver	Normal liver tissue	normal
B2	10	40	M	Liver	Normal liver tissue	normal
B3	11	40	M	Liver	Normal liver tissue	normal
B4	12	38	M	Liver	Normal liver tissue	normal
B5	13	34	M	Liver	Normal liver tissue	normal
B6	14	27	M	Liver	Normal liver tissue	normal
B7	15	25	F	Liver	Normal liver tissue	normal
B8	16	42	F	Pancreas	Normal pancreas tissue	normal
C1	17	35	F	Pancreas	Normal pancreas tissue	normal
C2	18	1 mon.	M	Pancreas	Normal pancreas tissue	normal
C3	19	35	M	Pancreas	Normal pancreas tissue	normal
C4	20	38	F	Pancreas	Normal pancreas tissue	normal
C5	21	56	M	Stomach	Normal stomach tissue	normal
C6	22	35	F	Stomach	Normal stomach tissue	normal
C7	23	35	M	Stomach	Normal stomach tissue	normal
C8	24	22	M	Stomach	Normal gastric mucosa tissue	normal
D1	25	40	M	Stomach	Normal stomach tissue	normal
D2	26	38	F	Stomach	Normal stomach tissue	normal
D3	27	35	M	Stomach	Normal stomach tissue	normal
D4	28	48	M	Stomach	Normal stomach tissue	normal
D5	29	52	F	Stomach	Normal stomach tissue	normal
D6	30	24	M	Esophagus	Normal esophagus tissue	normal
D7	31	21	F	Esophagus	Normal esophagus tissue (fibrous and connective tissue)	normal
D8	32	26	M	Esophagus	Normal esophagus tissue	normal
E1	33	22	M	Esophagus	Normal esophagus tissue	normal
E2	34	48	M	Esophagus	Normal esophagus tissue	normal
E3	35	59	M	Esophagus	Normal esophagus tissue	normal
E4	36	50	F	Colon	Normal colon tissue	normal
E5	37	49	M	Colon	Normal colon tissue	normal
E6	38	21	F	Colon	Normal colon tissue (fibrous and smooth muscle tissue)	normal
E7	39	35	M	Colon	Normal colon tissue	normal
E8	40	49	M	Intestine	Normal small intestine tissue (sparse)	normal
F1	41	35	F	Intestine	Normal small intestine tissue	normal
F2	42	40	M	Intestine	Normal small intestine tissue	normal
F3	43	38	F	Intestine	Normal small intestine tissue	normal
F4	44	42	F	Intestine	Normal small intestine tissue with necrosis	normal
F5	45	57	F	Intestine	Normal small intestine tissue	normal
F6	46	37	M	Intestine	Normal small intestine tissue	normal
F7	47	61	F	Intestine	Normal small intestine tissue	normal
F8	48	27	M	Intestine	Normal small intestine tissue	normal

TABLE 10 (OV1502; Ovarian Cancer array)						
Position	No.	Age	Sex	Organ/Anatomic Site	Pathology diagnosis	TNM
A1	1	37	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
A2	2	56	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
A3	3	30	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
A4	4	51	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
A5	5	81	F	Ovary	Serous adenocarcinoma	T1aN0M0
A6	6	34	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
A7	7	56	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
A8	8	39	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
A9	9	54	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
A10	10	66	F	Ovary	Serous papillary adenocarcinoma	-
A11	11	56	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
A12	12	30	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
A13	13	66	F	Ovary	Serous papillary adenocarcinoma	T3aN0M0
A14	14	69	F	Ovary	Serous adenocarcinoma	T2N0M0
A15	15	49	F	Ovary	Serous adenocarcinoma	T1aN0M0
B1	16	37	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
B2	17	56	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
B3	18	30	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
B4	19	51	F	Ovary	Serous papillary adenocarcinoma (ovary tissue)	T1aN0M0
B5	20	81	F	Ovary	Serous adenocarcinoma	T1aN0M0
B6	21	34	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
B7	22	56	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
B8	23	39	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
B9	24	54	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
B10	25	66	F	Ovary	Serous papillary adenocarcinoma	-
B11	26	56	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
B12	27	30	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
B13	28	66	F	Ovary	Serous papillary adenocarcinoma	T3aN0M0
B14	29	69	F	Ovary	Serous adenocarcinoma	T2N0M0
B15	30	49	F	Ovary	Serous adenocarcinoma	T1aN0M0
C1	31	59	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
C2	32	48	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
C3	33	58	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
C4	34	70	F	Ovary	Serous adenocarcinoma	T2N0M0
C5	35	54	F	Ovary	Serous adenocarcinoma	T1cN0M0
C6	36	47	F	Ovary	Serous adenocarcinoma	T1N0M0
C7	37	55	F	Ovary	Serous adenocarcinoma	T1aN0M0
C8	38	46	F	Ovary	Serous adenocarcinoma	T1N0M0
C9	39	79	F	Ovary	Serous adenocarcinoma	T1aN0M0
C10	40	56	F	Ovary	Serous adenocarcinoma	T1bN0M0
C11	41	35	F	Ovary	Serous adenocarcinoma	T1N0M0
C12	42	56	F	Ovary	Serous adenocarcinoma	T1N0M0
C13	43	49	F	Ovary	Serous adenocarcinoma	T1aN0M0
C14	44	54	F	Ovary	Serous adenocarcinoma	T2N0M0
C15	45	71	F	Ovary	Serous adenocarcinoma	T1cN0M0
D1	46	59	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
D2	47	48	F	Ovary	Serous papillary adenocarcinoma	T1N0M0

D3	48	58	F	Ovary	Serous papillary adenocarcinoma	T1aN0M0
D4	49	70	F	Ovary	Serous adenocarcinoma	T2N0M0
D5	50	54	F	Ovary	Serous adenocarcinoma	T1cN0M0
D6	51	47	F	Ovary	Serous adenocarcinoma	T1N0M0
D7	52	55	F	Ovary	Serous adenocarcinoma	T1aN0M0
D8	53	46	F	Ovary	Serous adenocarcinoma	T1N0M0
D9	54	79	F	Ovary	Serous adenocarcinoma	T1aN0M0
D10	55	56	F	Ovary	Serous adenocarcinoma	T1bN0M0
D11	56	35	F	Ovary	Serous adenocarcinoma	T1N0M0
D12	57	56	F	Ovary	Serous adenocarcinoma	T1N0M0
D13	58	49	F	Ovary	Serous adenocarcinoma	T1aN0M0
D14	59	54	F	Ovary	Serous adenocarcinoma	T2N0M0
D15	60	71	F	Ovary	Serous adenocarcinoma	T1cN0M0
E1	61	72	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
E2	62	55	F	Ovary	Serous papillary adenocarcinoma	T1bN0M0
E3	63	52	F	Ovary	Serous papillary adenocarcinoma	T1cN0M0
E4	64	41	F	Ovary	Serous adenocarcinoma	T1bN0M0
E5	65	77	F	Ovary	Serous adenocarcinoma	T1bN0M0
E6	66	50	F	Ovary	Serous adenocarcinoma	T1aN0M0
E7	67	55	F	Ovary	Serous adenocarcinoma	T1aN0M0
E8	68	60	F	Ovary	Mucinous adenocarcinoma	T1aN0M0
E9	69	52	F	Ovary	Mucinous adenocarcinoma	T1N0M0
E10	70	63	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
E11	71	66	F	Ovary	Mucinous papillary adenocarcinoma	T1cN0M0
E12	72	58	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
E13	73	48	F	Ovary	Mucinous papillary adenocarcinoma	T1bN0M0
E14	74	25	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
E15	75	52	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
F1	76	72	F	Ovary	Serous papillary adenocarcinoma	T1N0M0
F2	77	55	F	Ovary	Serous papillary adenocarcinoma	T1bN0M0
F3	78	52	F	Ovary	Serous papillary adenocarcinoma	T1cN0M0
F4	79	41	F	Ovary	Serous adenocarcinoma with necrosis (sparse)	T1bN0M0
F5	80	77	F	Ovary	Serous adenocarcinoma (tumoral necrosis)	T1bN0M0
F6	81	50	F	Ovary	Serous adenocarcinoma	T1aN0M0
F7	82	55	F	Ovary	Serous adenocarcinoma	T1aN0M0
F8	83	60	F	Ovary	Mucinous adenocarcinoma	T1aN0M0
F9	84	52	F	Ovary	Mucinous adenocarcinoma	T1N0M0
F10	85	63	F	Ovary	Mucinous papillary adenocarcinoma (mucosa and necrotic tissue)	T1aN0M0
F11	86	66	F	Ovary	Mucinous papillary adenocarcinoma	T1cN0M0
F12	87	58	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
F13	88	48	F	Ovary	Mucinous papillary adenocarcinoma	T1bN0M0
F14	89	25	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
F15	90	52	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
G1	91	34	F	Ovary	Mucinous papillary adenocarcinoma	T1bN0M0
G2	92	63	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
G3	93	48	F	Ovary	Mucinous adenocarcinoma	T1N0M0
G4	94	61	F	Ovary	Mucinous adenocarcinoma	T1aN0M0
G5	95	39	F	Ovary	Mucinous papillary adenocarcinoma	T1bN0M0
G6	96	32	F	Ovary	Mucinous papillary adenocarcinoma	T1N0M0
G7	97	50	F	Ovary	Mucinous papillary adenocarcinoma (mucinous carcinoma sparse)	T1aN0M0

G8	98	40	F	Ovary	Mucinous papillary adenocarcinoma	T1cN0M0
G9	99	45	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
G10	100	46	F	Ovary	Mucinous papillary adenocarcinoma	T1N0M0
G11	101	44	F	Ovary	Endometrioid adenocarcinoma	T1bN0M0
G12	102	48	F	Ovary	Endometrioid adenocarcinoma	T1N0M0
G13	103	41	F	Ovary	Endometrioid adenocarcinoma	T1N0M0
G14	104	70	F	Ovary	Endometrioid adenocarcinoma	T1aN0M0
G15	105	47	F	Ovary	Endometrioid adenocarcinoma	T1N0M0
H1	106	34	F	Ovary	Mucinous papillary adenocarcinoma	T1bN0M0
H2	107	63	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
H3	108	48	F	Ovary	Mucinous adenocarcinoma	T1N0M0
H4	109	61	F	Ovary	Mucinous adenocarcinoma	T1aN0M0
H5	110	39	F	Ovary	Mucinous papillary adenocarcinoma	T1bN0M0
H6	111	32	F	Ovary	Mucinous papillary adenocarcinoma	T1N0M0
H7	112	50	F	Ovary	Mucinous papillary adenocarcinoma (mucinous carcinoma sparse)	T1aN0M0
H8	113	40	F	Ovary	Mucinous papillary adenocarcinoma	T1cN0M0
H9	114	45	F	Ovary	Mucinous papillary adenocarcinoma	T1aN0M0
H10	115	46	F	Ovary	Mucinous papillary adenocarcinoma	T1N0M0
H11	116	44	F	Ovary	Endometrioid adenocarcinoma	T1bN0M0
H12	117	48	F	Ovary	Endometrioid adenocarcinoma	T1N0M0
H13	118	41	F	Ovary	Endometrioid adenocarcinoma	T1N0M0
H14	119	70	F	Ovary	Endometrioid adenocarcinoma	T1aN0M0
H15	120	47	F	Ovary	Endometrioid adenocarcinoma	T1N0M0
I1	121	39	F	Ovary	Clear cell carcinoma	T1N0M0
I2	122	49	F	Ovary	Clear cell carcinoma	T1N0M0
I3	123	25	F	Ovary	Endodermal sinus carcinoma	T1bN0M0
I4	124	56	F	Ovary	Endodermal sinus carcinoma	T1aN0M0
I5	125	25	F	Ovary	Endodermal sinus carcinoma	T1cN0M0
I6	126	56	F	Ovary	Endodermal sinus carcinoma	T1cN0M0
I7	127	49	F	Ovary	Granular cell tumor	T1N0M0
I8	128	50	F	Ovary	Granular cell tumor	T1aN0M0
I9	129	36	F	Ovary	Dysgerminoma	T1bN0M0
I10	130	11	F	Ovary	Dysgerminoma	T1N0M0
I11	131	34	F	Ovary	Normal ovary tissue	-
I12	132	19	F	Ovary	Normal ovary tissue	-
I13	133	41	F	Ovary	Cancer adjacent normal ovary tissue	-
I14	134	53	F	Ovary	Cancer adjacent normal ovary tissue	-
I15	135	48	F	Ovary	Cancer adjacent normal ovary tissue	-
J1	136	39	F	Ovary	Clear cell carcinoma	T1N0M0
J2	137	49	F	Ovary	Clear cell carcinoma	T1N0M0
J3	138	25	F	Ovary	Endodermal sinus carcinoma	T1bN0M0
J4	139	56	F	Ovary	Endodermal sinus carcinoma	T1aN0M0
J5	140	25	F	Ovary	Endodermal sinus carcinoma	T1cN0M0
J6	141	56	F	Ovary	Endodermal sinus carcinoma	T1cN0M0
J7	142	49	F	Ovary	Granular cell tumor	T1N0M0
J8	143	50	F	Ovary	Granular cell tumor	T1aN0M0
J9	144	36	F	Ovary	Dysgerminoma	T1bN0M0
J10	145	11	F	Ovary	Dysgerminoma	T1N0M0
J11	146	34	F	Ovary	Normal ovary tissue	-
J12	147	19	F	Ovary	Normal ovary tissue	-
J13	148	41	F	Ovary	Cancer adjacent normal ovary tissue	-
J14	149	53	F	Ovary	Cancer adjacent normal ovary tissue	-
J15	150	48	F	Ovary	Cancer adjacent normal ovary tissue	-

TABLE 11 (PA807; pancreatic cancer array)						
Position	No.	Age	Sex	Organ/Anatomic Site	Pathology diagnosis	TNM
A1	1	46	F	Pancreas	Adenocarcinoma (sparse)	T3N0M0
A2	2	46	F	Pancreas	Adenocarcinoma	T3N0M0
A3	3	77	M	Pancreas	Adenocarcinoma	T2N0M0
A4	4	77	M	Pancreas	Adenocarcinoma	T2N0M0
A5	5	67	F	Pancreas	Adenocarcinoma	T3N1bM0
A6	6	67	F	Pancreas	Adenocarcinoma	T3N1bM0
A7	7	42	M	Pancreas	Adenocarcinoma	T2N0M0
A8	8	42	M	Pancreas	Adenocarcinoma (fibrous tissue and blood vessel)	T2N0M0
A9	9	41	F	Pancreas	Adenocarcinoma (sparse)	T3N0M0
A10	10	41	F	Pancreas	Adenocarcinoma (sparse)	T3N0M0
A11	11	42	F	Pancreas	Adenocarcinoma	T3N0M0
A12	12	42	F	Pancreas	Adenocarcinoma	T3N0M0
A13	13	57	M	Pancreas	Adenocarcinoma	T3N0M0
A14	14	57	M	Pancreas	Carcinoma tissue (sparse)	T3N0M0
A15	15	51	M	Pancreas	Adenocarcinoma	T3N0M0
A16	16	51	M	Pancreas	Adenocarcinoma	T3N0M0
B1	17	60	M	Pancreas	Adenocarcinoma	T3N0M0
B2	18	60	M	Pancreas	Adenocarcinoma	T3N0M0
B3	19	52	M	Pancreas	Mucinous adenocarcinoma (sparse) with necrosis	T3N0M0
B4	20	52	M	Pancreas	Adenocarcinoma	T3N0M0
B5	21	54	F	Pancreas	Adenocarcinoma	T3N0M0
B6	22	54	F	Pancreas	Adenocarcinoma	T3N0M0
B7	23	62	F	Pancreas	Mucinous adenocarcinoma	T4N0M0
B8	24	62	F	Pancreas	Mucinous adenocarcinoma	T4N0M0
B9	25	47	F	Pancreas	Mucinous adenocarcinoma	T3N0M0
B10	26	47	F	Pancreas	Mucinous adenocarcinoma	T3N0M0
B11	27	50	M	Pancreas	Adenocarcinoma	TxN0M0
B12	28	50	M	Pancreas	Adenocarcinoma	TxN0M0
B13	29	43	F	Pancreas	Carcinoma tissue (sparse)	T3N0M0
B14	30	43	F	Pancreas	Adenocarcinoma	T3N0M0
B15	31	66	F	Pancreas	Adenocarcinoma (sparse)	T2N0M0
B16	32	66	F	Pancreas	Adenocarcinoma (pancreatic tissue)	T2N0M0
C1	33	50	F	Pancreas	Adenocarcinoma	T2N0M0
C2	34	50	F	Pancreas	Adenocarcinoma	T2N0M0
C3	35	55	F	Pancreas	Adenocarcinoma	T2N0M0
C4	36	55	F	Pancreas	Adenocarcinoma (pancreatic tissue)	T2N0M0
C5	37	60	M	Pancreas	Adenocarcinoma (pancreatic tissue)	T3N0M0
C6	38	60	M	Pancreas	Adenocarcinoma (fibrofatty tissue)	T3N0M0
C7	39	66	M	Pancreas	Adenocarcinoma	TxN0M0
C8	40	66	M	Pancreas	Adenocarcinoma	TxN0M0
C9	41	52	M	Pancreas	Adenocarcinoma	T3N0M0
C10	42	52	M	Pancreas	Adenocarcinoma	T3N0M0
C11	43	47	M	Pancreas	Adenocarcinoma	T3N0M0
C12	44	47	M	Pancreas	Adenocarcinoma	T3N0M0
C13	45	51	F	Pancreas	Adenocarcinoma	T4N0M0

C14	46	51	F	Pancreas	Adenocarcinoma	T4N0M0
C15	47	57	M	Pancreas	Adenocarcinoma	T3N0M0
C16	48	57	M	Pancreas	Adenocarcinoma	T3N0M0
D1	49	47	M	Pancreas	Adenocarcinoma (sparse)	T3N0M0
D2	50	47	M	Pancreas	Adenocarcinoma (chronic pancreatitis)	T3N0M0
D3	51	61	M	Pancreas	Adenocarcinoma	T3N0M0
D4	52	61	M	Pancreas	Adenocarcinoma	T3N0M0
D5	53	57	F	Pancreas	Adenocarcinoma	T3N0M0
D6	54	57	F	Pancreas	Adenocarcinoma	T3N0M0
D7	55	50	M	Pancreas	Adenocarcinoma	T3N0M0
D8	56	50	M	Pancreas	Adenocarcinoma	T3N0M0
D9	57	52	F	Pancreas	Adenocarcinoma	T2N0M0
D10	58	52	F	Pancreas	Adenocarcinoma (sparse)	T2N0M0
D11	59	75	M	Pancreas	Adenocarcinoma with necrosis	TxNxMx
D12	60	75	M	Pancreas	Adenocarcinoma	TxNxMx
D13	61	64	F	Pancreas	Adenocarcinoma with necrosis	T3N0M0
D14	62	64	F	Pancreas	Adenocarcinoma (sparse) with necrosis	T3N0M0
D15	63	68	F	Pancreas	Adenocarcinoma (sparse)	TxN0M0
D16	64	68	F	Pancreas	Adenocarcinoma	TxN0M0
E1	65	75	F	Pancreas	Adenocarcinoma	T3N0M0
E2	66	75	F	Pancreas	Adenocarcinoma	T3N0M0
E3	67	65	M	Pancreas	Adenocarcinoma	T3N0M0
E4	68	65	M	Pancreas	Adenocarcinoma	T3N0M0
E5	69	61	M	Pancreas	Adenocarcinoma	TxN0M0
E6	70	61	M	Pancreas	Adenocarcinoma	TxN0M0
E7	71	55	M	Pancreas	Adenocarcinoma with necrosis	T3N0M0
E8	72	55	M	Pancreas	Adenocarcinoma with necrosis	T3N0M0
E9	73	47	M	Pancreas	Adenocarcinoma (fibrofatty tissue and blood vessel)	T3N0M0
E10	74	47	M	Pancreas	Adenocarcinoma	T3N0M0
E11	75	49	M	Pancreas	Adenocarcinoma	T2N0M0
E12	76	49	M	Pancreas	Adenocarcinoma	T2N0M0
E13	77	49	M	Pancreas	Adenocarcinoma	T3N0M0
E14	78	49	M	Pancreas	Adenocarcinoma	T3N0M0
E15	79	48	F	Pancreas	Adenocarcinoma	T3N0M0
E16	80	48	F	Pancreas	Adenocarcinoma (fibrous tissue and blood vessel)	T3N0M0
F1	81	48	F	Pancreas	Adenocarcinoma	T3N0M0
F2	82	48	F	Pancreas	Adenocarcinoma	T3N0M0
F3	83	64	M	Pancreas	Adenocarcinoma	T3N0M0
F4	84	64	M	Pancreas	Adenocarcinoma (sparse)	T3N0M0
F5	85	65	M	Pancreas	Adenocarcinoma	T2N0M0
F6	86	65	M	Pancreas	Adenocarcinoma	T2N0M0
F7	87	48	M	Pancreas	Adenocarcinoma	T2N0M0
F8	88	48	M	Pancreas	Adenocarcinoma with necrosis	T2N0M0
F9	89	46	F	Pancreas	Adenocarcinoma	T2N0M0
F10	90	46	F	Pancreas	Adenocarcinoma	T2N0M0
F11	91	39	F	Pancreas	Adenocarcinoma	TxNxMx
F12	92	39	F	Pancreas	Adenocarcinoma (sparse)	TxNxMx
F13	93	57	M	Pancreas	Adenocarcinoma with necrosis	T3N0M0
F14	94	57	M	Pancreas	Adenocarcinoma with necrosis	T3N0M0
F15	95	44	F	Pancreas	Adenocarcinoma	T2N0M0

F16	96	44	F	Pancreas	Adenocarcinoma	T2N0M0
G1	97	64	M	Pancreas	Adenocarcinoma (sparse)	TxN0M0
G2	98	64	M	Pancreas	Adenocarcinoma (sparse)	TxN0M0
G3	99	66	M	Pancreas	Adenocarcinoma	T2N0M0
G4	100	66	M	Pancreas	Adenocarcinoma	T2N0M0
G5	101	74	M	Pancreas	Adenocarcinoma	T2N0M0
G6	102	74	M	Pancreas	Adenocarcinoma	T2N0M0
G7	103	49	M	Pancreas	Adenocarcinoma	T2N0M0
G8	104	49	M	Pancreas	Adenocarcinoma	T2N0M0
G9	105	51	F	Pancreas	Adenocarcinoma	T3N0M0
G10	106	51	F	Pancreas	Adenocarcinoma	T3N0M0
G11	107	54	F	Pancreas	Adenocarcinoma	T2N0M0
G12	108	54	F	Pancreas	Adenocarcinoma	T2N0M0
G13	109	47	F	Pancreas	Adenocarcinoma	T3N1M0
G14	110	47	F	Pancreas	Adenocarcinoma (sparse)	T3N1M0
G15	111	44	M	Pancreas	Adenocarcinoma (sparse)	T3N0M0
G16	112	44	M	Pancreas	Adenocarcinoma	T3N0M0
H1	113	60	M	Pancreas	Adenocarcinoma	T3N0M0
H2	114	60	M	Pancreas	Adenocarcinoma	T3N0M0
H3	115	64	M	Pancreas	Adenocarcinoma (sparse)	T3N0M0
H4	116	64	M	Pancreas	Adenocarcinoma (sparse)	T3N0M0
H5	117	52	M	Pancreas	Adenocarcinoma	TxN0M0
H6	118	52	M	Pancreas	Adenocarcinoma	TxN0M0
H7	119	52	M	Pancreas	Adenocarcinoma (tumoral necrosis)	T3N0M0
H8	120	52	M	Pancreas	Adenocarcinoma (tumoral necrosis)	T3N0M0
H9	121	49	M	Pancreas	Adenocarcinoma (fibrofatty tissue)	T2N0M0
H10	122	49	M	Pancreas	Adenocarcinoma	T2N0M0
H11	123	42	F	Pancreas	Adenocarcinoma	T2N0M0
H12	124	42	F	Pancreas	Adenocarcinoma	T2N0M0
H13	125	76	F	Pancreas	Adenocarcinoma	T3N0M0
H14	126	76	F	Pancreas	Adenocarcinoma	T3N0M0
H15	127	59	M	Pancreas	Adenocarcinoma	TxNxM0
H16	128	59	M	Pancreas	Adenocarcinoma	TxNxM0
I1	129	76	F	Pancreas	Adenocarcinoma	T3N0M0
I2	130	76	F	Pancreas	Adenocarcinoma	T3N0M0
I3	131	64	F	Pancreas	Adenocarcinoma (tumoral necrosis)	TxN0M0
I4	132	64	F	Pancreas	Adenocarcinoma (tumoral necrosis)	TxN0M0
I5	133	58	F	Pancreas	Adenocarcinoma	T2N0M0
I6	134	58	F	Pancreas	Adenocarcinoma	T2N0M0
I7	135	52	M	Pancreas	Adenocarcinoma	T2N0M0
I8	136	52	M	Pancreas	Adenocarcinoma	T2N0M0
I9	137	76	M	Pancreas	Adenocarcinoma	T3N0M0
I10	138	76	M	Pancreas	Adenocarcinoma	T3N0M0
I11	139	53	F	Pancreas	Adenocarcinoma	T2N0M0
I12	140	53	F	Pancreas	Adenocarcinoma	T2N0M0
I13	141	55	M	Pancreas	Adenocarcinoma	T3N0M0
I14	142	55	M	Pancreas	Adenocarcinoma	T3N0M0
I15	143	47	M	Pancreas	Adenocarcinoma	T3N0M0
I16	144	47	M	Pancreas	Adenocarcinoma	T3N0M0

J1	145	46	F	Pancreas	Adenocarcinoma	T4N0M0
J2	146	46	F	Pancreas	Adenocarcinoma (pancreatic tissue)	T4N0M0
J3	147	67	M	Pancreas	Adenocarcinoma	T3N0M0
J4	148	67	M	Pancreas	Adenocarcinoma	T3N0M0
J5	149	61	M	Pancreas	Adenocarcinoma	T3N0M0
J6	150	61	M	Pancreas	Adenocarcinoma	T3N0M0
J7	151	60	M	Pancreas	Adenocarcinoma	T2N0M0
J8	152	60	M	Pancreas	Adenocarcinoma	T2N0M0
J9	153	78	M	Pancreas	Adenocarcinoma	T3N0M0
J10	154	78	M	Pancreas	Adenocarcinoma	T3N0M0
J11	155	45	M	Pancreas	Adenocarcinoma	T2N1M0
J12	156	45	M	Pancreas	Adenocarcinoma	T2N1M0
J13	157	49	M	Pancreas	Adenocarcinoma	T1N0M0
J14	158	49	M	Pancreas	Adenocarcinoma	T1N0M0
J15	159	69	M	Pancreas	Adenocarcinoma (chronic pancreatitis)	T2N0M0
J16	160	69	M	Pancreas	Adenocarcinoma	T2N0M0
K1	161	57	F	Pancreas	Adenocarcinoma	T2N0M0
K2	162	57	F	Pancreas	Adenocarcinoma	T2N0M0
K3	163	50	M	Pancreas	Adenocarcinoma	T2N0M0
K4	164	50	M	Pancreas	Adenocarcinoma	T2N0M0
K5	165	50	M	Pancreas	Adenocarcinoma	T2N0M0
K6	166	50	M	Pancreas	Adenocarcinoma	T2N0M0
K7	167	41	M	Pancreas	Adenocarcinoma	T4N1M0
K8	168	41	M	Pancreas	Adenocarcinoma	T4N1M0
K9	169	62	M	Pancreas	Adenocarcinoma	T3N0M0
K10	170	62	M	Pancreas	Adenocarcinoma	T3N0M0
K11	171	52	M	Pancreas	Adenocarcinoma (sparse)	T2N0M0
K12	172	52	M	Pancreas	Adenocarcinoma (sparse)	T2N0M0
K13	173	62	M	Pancreas	Adenocarcinoma	T3NxM0
K14	174	62	M	Pancreas	Adenocarcinoma	T3NxM0
K15	175	49	F	Pancreas	Adenosquamous carcinoma	T3N1M0
K16	176	49	F	Pancreas	Adenosquamous carcinoma	T3N1M0
L1	177	60	F	Pancreas	Adenosquamous carcinoma	T4N0M0
L2	178	60	F	Pancreas	Adenosquamous carcinoma	T4N0M0
L3	179	50	M	Pancreas	Squamous cell carcinoma	T3N0M0
L4	180	50	M	Pancreas	Squamous cell carcinoma	T3N0M0
L5	181	62	M	Pancreas	Squamous cell carcinoma	T3N0M0
L6	182	62	M	Pancreas	Squamous cell carcinoma	T3N0M0
L7	183	66	F	Pancreas	Acinar cell carcinoma	TxN0M0
L8	184	66	F	Pancreas	Acinar cell carcinoma	TxN0M0
L9	185	53	M	Pancreas	Acinar cell carcinoma	T2N0M0
L10	186	53	M	Pancreas	Acinar cell carcinoma with calcification	T2N0M0
L11	187	42	M	Pancreas	Neuroendocrine carcinoma	T2N0M0
L12	188	42	M	Pancreas	Neuroendocrine carcinoma	T2N0M0
L13	189	56	F	Pancreas	Cancer adjacent normal pancreatic tissue	-
L14	190	56	F	Pancreas	Cancer adjacent normal pancreatic tissue	-
L15	191	52	M	Pancreas	Cancer adjacent normal pancreatic tissue of No. 119	-
L16	192	52	M	Pancreas	Cancer adjacent normal pancreatic tissue of No. 119	-

M1	193	74	F	Pancreas	Cancer adjacent normal pancreatic tissue	-
M2	194	74	F	Pancreas	Cancer adjacent normal pancreatic tissue	-
M3	195	55	F	Pancreas	Cancer adjacent normal pancreatic tissue of No. 35	-
M4	196	55	F	Pancreas	Cancer adjacent normal pancreatic tissue of No. 35	-
M5	197	65	M	Pancreas	Cancer adjacent normal pancreatic tissue (fibrous tissue and blood vessel) of No. 67	-
M6	198	65	M	Pancreas	Cancer adjacent normal pancreatic tissue of No. 67	-
M7	199	14	F	Pancreas	Normal pancreatic tissue	-
M8	200	14	F	Pancreas	Normal pancreatic tissue	-
M9	201	38	F	Pancreas	Normal pancreatic tissue	-
M10	202	38	F	Pancreas	Normal pancreatic tissue	-
M11	203	33	M	Pancreas	Normal pancreatic tissue	-
M12	204	33	M	Pancreas	Normal pancreatic tissue	-
M13	205	40	F	Pancreas	Normal pancreatic tissue	-
M14	206	40	F	Pancreas	Normal pancreatic tissue	-
M15	207	19	M	Pancreas	Normal pancreatic tissue (sparse)	-
M16	208	19	M	Pancreas	Normal pancreatic tissue	-

TABLE 12 (GA802a; Cholangiocarcinoma cancer array)

Position	No.	Age	Sex	Organ/Anatomic Site	Pathology diagnosis	TNM
A1	1	57	M	Bile duct	Adenocarcinoma	T3N0M0
A2	2	53	M	Bile duct	Adenocarcinoma (sparse)	T2N0M0
A3	3	62	M	Bile duct	Adenocarcinoma	T2N0M0
A4	4	42	F	Bile duct	Adenocarcinoma	T2N0M0
A5	5	78	M	Bile duct	Adenocarcinoma	T1N0M0
A6	6	59	F	Bile duct	Adenocarcinoma (sparse)	T3N1M0
A7	7	53	M	Bile duct	Adenocarcinoma	T3N0M0
A8	8	70	F	Common bile duct	Papillary adenocarcinoma	T1N0M0
A9	9	51	F	Common bile duct	Adenocarcinoma	T3N0M0
A10	10	67	M	Common bile duct	Adenocarcinoma	T3N0M0
B1	11	52	F	Common bile duct	Adenocarcinoma	T2N0M0
B2	12	58	M	Common bile duct	Adenocarcinoma	T3N0M0
B3	13	64	F	Bile duct	Adenocarcinoma	T3N0M0
B4	14	68	M	Common bile duct	Adenocarcinoma	T2N0M0
B5	15	62	M	Bile duct	Adenocarcinoma	T3N1M0
B6	16	65	M	Common bile duct	Adenocarcinoma	T2N0M0
B7	17	60	M	Bile duct	Adenocarcinoma	T2N0M0
B8	18	69	M	Common bile duct	Adenocarcinoma	T3N1M0
B9	19	74	F	Common bile duct	Adenocarcinoma	T2N0M0
B10	20	71	M	Bile duct	Adenocarcinoma	T2N0M0
C1	21	62	M	Bile duct	Adenocarcinoma	T3N0M0
C2	22	66	F	Common bile duct	Adenocarcinoma (sparse)	T3N0M0
C3	23	78	F	Bile duct	Adenocarcinoma	T3N1M0
C4	24	53	F	Bile duct	Adenocarcinoma	T1N0M0

C5	25	28	F	Common bile duct	Adenocarcinoma (sparse)	T2N0M0
C6	26	29	M	Bile duct	Adenocarcinoma	T1N1M0
C7	27	70	F	Common bile duct	Adenocarcinoma	T2N0M0
C8	28	65	F	Common bile duct	Adenocarcinoma	T3N0M0
C9	29	53	M	Common bile duct	Adenocarcinoma	T3N0M0
C10	30	58	F	Common bile duct	Adenocarcinoma	T3N1M0
D1	31	65	F	Common bile duct	Adenocarcinoma	T2N0M0
D2	32	70	F	Common bile duct	Adenocarcinoma	T3N1M0
D3	33	62	F	Bile duct	Adenocarcinoma	T2N0M0
D4	34	60	M	Common bile duct	Papillary adenocarcinoma	T2N1M0
D5	35	52	M	Common bile duct	Papillary adenocarcinoma	T3N1M0
D6	36	47	M	Common bile duct	Adenocarcinoma	T2N1M0
D7	37	70	F	Common bile duct	Adenocarcinoma	T3N0M0
D8	38	53	F	Common bile duct	Adenocarcinoma	T3N0M0
D9	39	50	M	Common bile duct	Adenocarcinoma	T2N0M0
D10	40	49	M	Common bile duct	Adenocarcinoma	T2N0M0
E1	41	75	F	Common bile duct	Adenocarcinoma	T2N0M0
E2	42	50	M	Common bile duct	Adenocarcinoma	T2N0M0
E3	43	73	F	Bile duct	Adenocarcinoma (sparse)	T3N1M0
E4	44	41	M	Common bile duct	Adenocarcinoma	T2N1M0
E5	45	72	M	Bile duct	Adenocarcinoma (sparse)	T3N0M0
E6	46	60	F	Bile duct	Adenocarcinoma	T2N1M0
E7	47	69	F	Bile duct	Adenocarcinoma	T3N1M0
E8	48	74	M	Common bile duct	Adenocarcinoma	T2N0M0
E9	49	36	F	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
E10	50	66	M	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
F1	51	52	M	Liver	Intrahepatic cholangiocarcinoma	T3N1M0
F2	52	51	F	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
F3	53	45	M	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
F4	54	64	M	Liver	Intrahepatic cholangiocarcinoma	T4N0M0
F5	55	65	F	Liver	Intrahepatic cholangiocarcinoma	T2N1M0
F6	56	34	M	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
F7	57	52	M	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
F8	58	66	F	Liver	Intrahepatic cholangiocarcinoma	T1N0M0
F9	59	58	F	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
F10	60	76	M	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
G1	61	49	M	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
G2	62	62	F	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
G3	63	45	M	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
G4	64	40	F	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
G5	65	52	F	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
G6	66	63	M	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
G7	67	69	F	Liver	Intrahepatic cholangiocarcinoma	T1N0M0
G8	68	49	M	Liver	Intrahepatic cholangiocarcinoma	T4N0M0
G9	69	26	M	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
G10	70	46	M	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
H1	71	45	F	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
H2	72	65	F	Liver	Intrahepatic cholangiocarcinoma	T4N0M0
H3	73	48	F	Liver	Intrahepatic cholangiocarcinoma	T2N0M0
H4	74	32	M	Liver	Intrahepatic cholangiocarcinoma	T3N0M0
H5	75	55	F	Liver	Intrahepatic cholangiocarcinoma	T3N1M0
H6	76	47	M	Liver	Adjacent normal liver tissue	-
H7	77	40	M	Liver	Adjacent normal liver tissue and portal area	-

H8	78	44	F	Liver	Adjacent normal liver tissue and portal area	-
H9	79	60	M	Liver	Adjacent normal liver tissue and portal area	-
H10	80	50	M	Liver	Adjacent normal liver tissue and portal area	-

TABLE 13 (FDA999x; normal tissue array)

Position	No.	Age	Sex	Organ/Anatomic Site	Pathology diagnosis	TNM
A1	1	42	M	Cerebrum	Cerebrum tissue	Normal
A2	2	53	M	Cerebrum	Cerebrum tissue	Normal
A3	3	37	M	Cerebrum	Cerebrum tissue	Normal
A4	4	26	M	Cerebellum	Cerebellum tissue	Normal
A5	5	45	M	Cerebellum	Cerebellum tissue	Normal
A6	6	35	M	Cerebellum	Cerebellum tissue	Normal
A7	7	43	M	Adrenal gland	Adrenal gland tissue	Normal
A8	8	18	F	Adrenal gland	Adrenal gland tissue	Normal
A9	9	23	M	Adrenal gland	Adrenal gland tissue	Normal
A10	10	25	F	Ovary	Ovary tissue	Normal
A11	11	19	F	Ovary	Ovary tissue	Normal
A12	12	40	F	Ovary	Ovary tissue	Normal
B1	13	23	M	Pancreas	Pancreas tissue (autolyzed)	Normal
B2	14	21	F	Pancreas	Pancreas tissue	Normal
B3	15	29	M	Pancreas	Pancreas tissue	Normal
B4	16	30	M	Lymph node	Lymph node tissue	Normal
B5	17	27	M	Lymph node	Lymph node tissue	Normal
B6	18	30	M	Lymph node	Lymph node tissue	Normal
B7	19	32	F	Hypophysis	Hypophysis tissue	Normal
B8	20	40	F	Hypophysis	Hypophysis tissue	Normal
B9	21	28	F	Hypophysis	Hypophysis tissue	Normal
B10	22	28	M	Testis	Testis tissue	Normal
B11	23	35	M	Testis	Testis tissue	Normal
B12	24	34	M	Testis	Testis tissue	Normal
C1	25	40	F	Thyroid gland	Thyroid gland tissue	Normal
C2	26	27	M	Thyroid gland	Thyroid gland tissue	Normal
C3	27	45	M	Thyroid gland	Thyroid gland tissue	Normal
C4	28	50	F	Breast	Breast tissue	Normal
C5	29	58	F	Breast	Breast tissue	Normal
C6	30	42	F	Breast	Breast tissue	Normal
C7	31	23	M	Spleen	Spleen tissue	Normal
C8	32	45	M	Spleen	Spleen tissue	Normal
C9	33	21	F	Spleen	Spleen tissue	Normal
C10	34	10	M	Tonsil	Tonsil tissue (Mucus gland)	Normal
C11	35	7	F	Tonsil	Tonsil tissue	Normal
C12	36	34	M	Tonsil	Tonsil tissue (fibrous tissue)	Normal
D1	37	21	F	Thymus gland	Thymus gland tissue	Normal
D2	38	15	F	Thymus gland	Thymus gland tissue	Normal
D3	39	23	M	Thymus gland	Thymus gland tissue	Normal
D4	40	21	F	Bone marrow	Bone marrow tissue	Normal
D5	41	22	M	Bone marrow	Bone marrow tissue	Normal
D6	42	21	F	Bone marrow	Bone marrow tissue	Normal
D7	43	47	M	Lung	Lung tissue	Normal
D8	44	19	M	Lung	Lung tissue	Normal

D9	45	21	F	Lung	Lung tissue	Normal
D10	46	35	M	Heart	Cardiac muscle tissue	Normal
D11	47	40	M	Heart	Cardiac muscle tissue	Normal
D12	48	21	F	Heart	Cardiac muscle tissue	Normal
E1	49	19	M	Esophagus	Esophagus tissue	Normal
E2	50	47	M	Esophagus	Esophagus tissue(smooth muscle)	Normal
E3	51	35	M	Esophagus	Esophagus tissue	Normal
E4	52	45	M	Stomach	Stomach tissue	Normal
E5	53	24	M	Stomach	Stomach tissue	Normal
E6	54	37	M	Stomach	Stomach tissue	Normal
E7	55	45	M	Small intestine	Small intestine tissue	Normal
E8	56	30	M	Small intestine	Small intestine tissue	Normal
E9	57	32	M	Small intestine	Small intestine tissue	Normal
E10	58	32	M	Colon	Colon tissue	Normal
E11	59	21	F	Colon	Colon tissue	Normal
E12	60	35	M	Colon	Colon tissue	Normal
F1	61	38	M	Liver	Liver tissue	Normal
F2	62	45	M	Liver	Liver tissue	Normal
F3	63	23	M	Liver	Liver tissue	Normal
F4	64	30	M	Salivary gland	Salivary gland tissue	Normal
F5	65	21	F	Salivary gland	Salivary gland tissue	Normal
F6	66	21	F	Salivary gland	Salivary gland tissue	Normal
F7	67	27	M	Kidney	Kidney tissue	Normal
F8	68	2	F	Kidney	Kidney tissue	Normal
F9	69	47	M	Kidney	Kidney tissue	Normal
F10	70	30	M	Prostate	Prostate tissue	Normal
F11	71	38	M	Prostate	Prostate tissue	Normal
F12	72	36	M	Prostate	Prostate tissue	Normal
G1	73	54	F	Uterus	Endometrium tissue	Normal
G2	74	40	F	Uterus	Endometrium tissue	Normal
G3	75	42	F	Uterus	Endometrium tissue	Normal
G4	76	46	F	Cervix	Cervix canals tissue	Normal
G5	77	54	F	Cervix	Cervix canals tissue	Normal
G6	78	37	F	Cervix	Cervix canals tissue	Normal
G7	79	32	M	Bladder	Bladder tissue	Normal
G8	80	35	M	Bladder	Bladder tissue	Normal
G9	81	34	M	Bladder	Bladder tissue	Normal
G10	82	21	M	Skeletal muscle	Skeletal muscle tissue	Normal
G11	83	25	M	Skeletal muscle	Skeletal muscle tissue	Normal
G12	84	40	M	Skeletal muscle	Skeletal muscle tissue	Normal
H1	85	21	M	Skin	Skin tissue(sparse)	Normal
H2	86	27	F	Skin	Skin tissue	Normal
H3	87	25	M	Skin	Skin tissue	Normal
H4	88	30	M	Nerve	Peripheral nerve tissue	Normal
H5	89	33	M	Nerve	Peripheral nerve tissue	Normal
H6	90	35	M	Nerve	Peripheral nerve tissue	Normal
H7	91	30	M	Artery	mesothelial tissue	Normal
H8	92	23	M	Pericardium	mesothelial tissue	Normal
H9	93	45	M	Pericardium	mesothelial tissue	Normal
H10	94	62	F	Eye	Cancer adjacent wall of eyeball (choroid)	AT
H11	95	42	M	Eye	Cancer adjacent wall of eyeball (choroid)	AT

H12	96	54	F	Eye	Cancer adjacent wall of eyeball	AT
I1	97	10	M	Larynx	Larynx tissue	Normal
I2	98	28	F	Larynx	Larynx tissue	Normal
I3	99	18	F	Larynx	Larynx tissue	Normal

TABLE 14 (LC121b; lung cancer array)

Position	No.	Age	Sex	Organ/Anatomic Site	Pathology diagnosis	TNM
A1	1	63	F	Lung	Squamous cell carcinoma	T3N1M0
A2	2	68	M	Lung	Squamous cell carcinoma	T2N2M0
A3	3	66	M	Lung	Squamous cell carcinoma	T2N2M0
A4	4	66	M	Lung	Squamous cell carcinoma	T3N1M0
A5	5	63	F	Lung	Squamous cell carcinoma	T2N2M0
A6	6	65	M	Lung	Squamous cell carcinoma	T2N2M0
A7	7	60	M	Lung	Squamous cell carcinoma	T4N0M0
A8	8	68	M	Lung	Squamous cell carcinoma	T2N2M0
A9	9	71	M	Lung	Squamous cell carcinoma	T2N2M0
A10	10	66	M	Lung	Squamous cell carcinoma	T2N2M0
A11	11	56	M	Lung	Squamous cell carcinoma	T3N2M0
A12	12	70	M	Lung	Squamous cell carcinoma	T4N2M0
B1	13	77	M	Lung	Squamous cell carcinoma	T4N1M0
B2	14	56	F	Lung	Squamous cell carcinoma	T4N0M0
B3	15	53	M	Lung	Squamous cell carcinoma	T4N0M0
B4	16	64	M	Lung	Squamous cell carcinoma	T3N2M0
B5	17	46	M	Lung	Squamous cell carcinoma	T2N0M0
B6	18	64	M	Lung	Squamous cell carcinoma	T4N0M0
B7	19	42	M	Lung	Squamous cell carcinoma	T3N1M0
B8	20	60	F	Lung	Squamous cell carcinoma	T2N2M0
B9	21	54	F	Lung	Large cell carcinoma	T2N1M0
B10	22	69	F	Lung	Large cell carcinoma	T3N0M0
B11	23	46	F	Lung	Large cell carcinoma	T2N0M0
B12	24	58	M	Lung	Large cell carcinoma	T2N0M0
C1	25	44	M	Lung	Large cell carcinoma	T1N0M0
C2	26	61	M	Lung	Large cell carcinoma	T4N0M0
C3	27	64	M	Lung	Large cell carcinoma	T2N0M0
C4	28	65	M	Lung	Large cell carcinoma	T2N0M0
C5	29	68	M	Lung	Large cell carcinoma (sparse)	T4N0M0
C6	30	31	M	Lung	Large cell carcinoma	T2N0M0
C7	31	58	M	Lung	Large cell carcinoma	T2N1M0
C8	32	58	M	Lung	Large cell carcinoma	T2N0M0
C9	33	60	M	Lung	Large cell carcinoma	T3N2M0
C10	34	40	M	Lung	Large cell carcinoma	T2N0M0
C11	35	60	M	Lung	Large cell carcinoma	T2N1M0
C12	36	49	M	Lung	Large cell carcinoma	T3N0M0
D1	37	65	F	Lung	Large cell carcinoma (sparse)	T2N1M0
D2	38	60	M	Lung	Large cell carcinoma	T2N0M0
D3	39	45	F	Lung	Large cell carcinoma	T2N0M0
D4	40	45	M	Lung	Large cell carcinoma	T1N0M0
D5	41	69	F	Lung	Large cell carcinoma	T2N0M0
D6	42	64	F	Lung	Large cell carcinoma	T2N0M0
D7	43	57	F	Lung	Large cell carcinoma	T2N0M0
D8	44	62	M	Lung	Large cell carcinoma	T2N0M0
D9	45	54	M	Lung	Large cell carcinoma	T2N0M0

D10	46	39	M	Lung	Large cell carcinoma	T2N0M0
D11	47	40	M	Lung	Large cell carcinoma	T2N0M0
D12	48	70	M	Lung	Large cell carcinoma	T3N1M0
E1	49	56	M	Lung	Large cell carcinoma	T2N1M0
E2	50	54	M	Lung	Large cell carcinoma	T2N1M0
E3	51	51	M	Lung	Large cell carcinoma	T2N0M0
E4	52	65	M	Lung	Large cell carcinoma	T2N0M0
E5	53	30	F	Lung	Large cell carcinoma	T2N1M0
E6	54	47	F	Lung	Large cell carcinoma	T2N2M0
E7	55	73	M	Lung	Large cell carcinoma (sparse)	T3N1M0
E8	56	51	M	Lung	Large cell carcinoma	T4N0M0
E9	57	60	M	Lung	Adenocarcinoma	T2N2M0
E10	58	49	M	Lung	Adenocarcinoma	T2N0M0
E11	59	52	M	Lung	Adenocarcinoma	T3N1M0
E12	60	46	F	Lung	Adenocarcinoma	T4N0M0
F1	61	62	F	Lung	Adenocarcinoma	T4N0M0
F2	62	56	M	Lung	Adenocarcinoma	T2N2M0
F3	63	54	F	Lung	Adenocarcinoma	T3N0M0
F4	64	61	M	Lung	Adenocarcinoma	T2N2M0
F5	65	62	M	Lung	Adenocarcinoma	T3N0M0
F6	66	36	F	Lung	Adenocarcinoma	T2N2M0
F7	67	41	M	Lung	Adenocarcinoma	T4N0M0
F8	68	66	M	Lung	Adenocarcinoma	T2N2M0
F9	69	64	M	Lung	Adenocarcinoma	T3N1M0
F10	70	68	F	Lung	Adenocarcinoma	T2N2M0
F11	71	60	M	Lung	Adenocarcinoma	T2N0M0
F12	72	42	M	Lung	Adenocarcinoma	T3N1M0
G1	73	35	M	Lung	Adenocarcinoma	T3N2M0
G2	74	62	M	Lung	Adenocarcinoma	T3N1M0
G3	75	65	F	Lung	Adenocarcinoma	T3N1M0
G4	76	54	F	Lung	Adenocarcinoma	T3N1M0
G5	77	46	M	Lung	Adenocarcinoma	T3N2M0
G6	78	72	M	Lung	Adenocarcinoma	T2N2M0
G7	79	56	M	Lung	Adenocarcinoma	T4N0M0
G8	80	30	F	Lung	Adenocarcinoma	T4N1M1
G9	81	48	M	Lung	Adenocarcinoma	T2N0M0
G10	82	47	F	Lung	Adenocarcinoma	T2N0M0
G11	83	69	M	Lung	Adenocarcinoma	T2N0M0
G12	84	65	M	Lung	Adenocarcinoma	T3N0M0
H1	85	60	F	Lung	Adenocarcinoma	T3N0M0
H2	86	22	F	Lung	Adenocarcinoma	T3N0M0
H3	87	55	F	Lung	Adenocarcinoma	T2N0M0
H4	88	43	F	Lung	Adenocarcinoma	T2N0M0
H5	89	50	F	Lung	Adenocarcinoma	T3N0M0
H6	90	49	F	Lung	Adenocarcinoma	T2N0M0
H7	91	47	F	Lung	Adenocarcinoma	T2N0M0
H8	92	50	M	Lung	Adenocarcinoma	T2N1M0
H9	93	82	M	Lung	Adenocarcinoma	T4N1M0
H10	94	54	F	Lung	Adenocarcinoma	T4N1M0
H11	95	38	M	Lung	Adenocarcinoma	T4N1M0
H12	96	57	M	Lung	Adenocarcinoma	T4N0M0
I1	97	42	M	Lung	Adenocarcinoma	T3N1M0
I2	98	65	M	Lung	Adenocarcinoma	T3N2M0

I3	99	59	M	Lung	Adenocarcinoma	T4N3M0
I4	100	62	M	Lung	Adenocarcinoma	T4N0M0
I5	101	49	F	Lung	Adenocarcinoma	T2N0M0
I6	102	39	M	Lung	Adenocarcinoma	T4N0M0
I7	103	42	F	Lung	Adenocarcinoma	T3N1M0
I8	104	60	F	Lung	Adenocarcinoma	T4N0M0
I9	105	42	F	Lung	Adenocarcinoma	T4N1M0
I10	106	54	F	Lung	Adenocarcinoma	T2N0M0
I11	107	50	M	Lung	Papillary adenocarcinoma	T3N1M0
I12	108	51	M	Lung	Papillary adenocarcinoma	T2N0M0
J1	109	47	F	Lung	Papillary adenocarcinoma	T3N1M0
J2	110	53	F	Lung	Papillary adenocarcinoma	T2N0M0
J3	111	56	M	Lung	Adjacent normal lung tissue	-
J4	112	57	M	Lung	Adjacent normal lung tissue	-
J5	113	51	F	Lung	Adjacent normal lung tissue	-
J6	114	68	M	Lung	Adjacent normal lung tissue	-
J7	115	53	F	Lung	Adjacent normal lung tissue	-
J8	116	45	M	Lung	Adjacent normal lung tissue	-
J9	117	27	M	Lung	Lung tissue	-
J10	118	15	F	Lung	Lung tissue	-
J11	119	48	M	Lung	Lung tissue	-
J12	120	16	F	Lung	Lung tissue	-

6.4 Example 4: Tn-CMET based CARs

6.4.1. Overview

[0438] Chimeric antigen receptors (CARs) having VH and VL domains of 15C4, 8H3, and 16E12 were designed. CARs were then evaluated in a target-specific cytotoxicity assay.

6.4.2. Materials and Methods

6.4.2.1 Vector Design

[0439] Various CAR constructs having scFvs having VH and VL domains of 15C4, 8H3, and 16E12 were designed (FIGS. 9A-9C). In the constructs, the VH and VL are attached together with one long linker (GGGS)₃ (SEQ ID NO:346) to the CD8a hinge followed by a second generation CAR-T (CD28 intracellular signal domain, and a CD3-zeta intracellular chain). The N-terminus of the scFvs was attached to a CD8a signal sequence. The CMET CAR-Ts were subcloned into the Virapower lentivirus vector pLENTI6.3-V5-DEST (Invitrogen).

[0440] Nucleotide sequences encoding the CARs are provided in Table 15. Amino acid sequences of the CARs are provided in Table 16.

Construct	Nucleotide Sequence	Nucleic Acid description	SEQ ID No.
15C4-CART	ATGGCTCTGCCGTTACAGCTCTGCTGCTGC CTCTGGCTCTGCTTCTGTCATGCCGCCAGACC TAACATCGTGATGACACAGAGCCCCAAGAGC ATGAGCATGTCCGTGGGCGAGAGAGTGACCC TGAGCTGTAAAGCCAGCGAGAACGTGGGCAT CTACGTGTCCTGGTATCAGCAGAAGCCCCGAG CAGAGCCCTAAGCTGCTGATCTACGGCCCCA	1-63=CD8A signal sequence 64-384=15C4 LC 385-429=Linker 430-780=15C4 HC	336

Table 15: Nucleotide sequences encoding CARs		
<p>GCAACAGATACACCGGCGTGCCCGATAGATT CACAGGCAGCGGAAGCGCCACCGACTTCACC CTGACAATCAGCTCTGTGCAGGCCGAGGACC TGGCCGATTATCACTGTGGCCAGAGCTACAG CTACCCCTTCACATTTGGCAGCGGCACCAAG CTGGAAATCAAAGGCGGCGGAGGATCTGGCG GAGGTGGAAGTGGCGGAGGCGGATCTCAAGT TCAGCTGCAGCAGTCCGATGCCGAGCTGGTT AAGCCTGGCGCCTCTGTGAAGATCAGCTGCA AGGCCAGCGGCTACACCTTCACCGATCACGC CATCCACTGGGTCAAGCAGAAACCTGAGCAG GGCCTCGAGTGGATCGGCTACTTTTTCTCCCG GCAACGGCGACATCAAGTACAACGAGAAGTT CAAGGGCAAAGCCACACTGACCGCCGACAAG AGCAGCAGCACAGCCTACATGCAGCTCAACA GCCTGACCAGCGAGGACAGCGCCGTGTA CTGCAAGAGAAGCCTGCCTGGACCTATGGAC TGTTGGGGCCAGGGAACAAGCGTGACCGTGT CCAGCACAACAACCCCTGCTCCTAGACCTCCT ACACCAGCTCCTACAATCGCCTCTCAACCTCT GTCTCTGCGGCCTGAGGCTTGTAGACCTGCT GCTGGCGGAGCCGTGCATACAAGAGGACTGG ATTTGCCTGCGACTTCTGGGTGCTCGTGGTT GTTGGCGGAGTGTGCTGGCCTGTTACTCTCTGC TGGTACCGTGGCCTTCATCATCTTTTGGGTC CGAAGCAAGCGGAGCCGGCTGCTGCACAGC GACTACATGAACATGACCCCTAGACGGCCCG GACCTACCAGAAAGCACTACCAGCCTTACGCT CCTCCTAGAGACTTCGCCGCTACCGGTCCA GAGTGAAGTTCAGCAGATCCGCCGATGCTCC CGCCTATCAGCAGGGACAGAACCAGCTGTAC AATGAGCTGAACCTGGGGCGCAGAGAAGAGT ACGACGTGCTGGATAAGCGGAGAGGCAGAGA TCCTGAGATGGGCGGCAAGCCCAGACGGAAG AATCCTCAAGAGGGCCTGTATAACGAGCTGC AGAAAGACAAGATGGCCGAGGCCTACAGCGA GATCGGAATGAAGGGCGAACGCAGAAGAGGC AAGGGCCACGATGGACTGTATCAGGGCCTGA GCACCGCCACCAAGGATACCTATGATGCCCT GCACATGCAGGCCCTGCCTCCAAGAAGAAAG AGAGGCTCTGGCGAAGGCAGAGGTAGCCTGC TGACATGTGGCGACGTGGAAGAGAACCCCGG ACCAATGGTGTCCAAGGGCGAAGAGGACAAC ATGGCCATCATCAAAGAATTCATGCGGTTCAA GGTGACATGGAAGGCAGCGTGAACGGCCAC GAGTTCGAGATTGAAGGCGAAGGCGAGGGCA GACCTTACGAGGGAACACAGACCGCCAAGCT GAAAGTGACCAAAGGCGGACCCCTGCCTTTC GCCTGGGATATCCTGTCTCCTCAGTTTATGTA CGGCAGCAAGGCCTACGTGAAGCACCCCGCC GATATTCCCGACTACCTGAAGCTGAGCTTCCC CGAGGGCTTCAAGTGGGAAAGAGTGATGAAC TTCGAGGACGGCGGCGTGGTCACAGTGACAC AAGATAGCAGTCTGCAGGACGGCGAGTTCAT CTACAAAGTGAAGCTGCGGGGCACCAACTTT CCCTCTGATGGCCCCGTGATGCAGAAAAAGA CCATGGGCTGGGAAGCCAGCAGCGAGAGAAT GTATCCTGAGGATGGCGCCCTGAAAGGCGAG ATCAAGCAGCGGCTGAAACTGAAAGGATGGCG GCCACTACGACGCCGAAGTGAAAACCACTA CAAGGCCAAGAAACCCGTGCAGCTGCCAGGC GCCTACAACGTGAACATCAAGCTGGACATTAC</p>	<p>781-915=CD8a hinge 916- 996=CD28 transmembrane domain 997-1118=intracellular signal domain 1120-1455=CD3-zeta intracellular chain 1456-2238=T2A-mCherry</p>	

Table 15: Nucleotide sequences encoding CARs			
	CAGCCACAACGAGGACTACACCATCGTGGAA CAGTACGAGAGAGCCGAAGGCAGGCACTCTA CAGGCGGAATGGACGAGCTGTATAAGTAG		
16E12- CART	ATGGCTCTGCCGTTACAGCTCTGCTGCTGC CTCTGGCTCTGCTTCTGCATGCCGCTAGACC CGACGTGCAGATCAGCCAGTCTCCTAGCTAT CTGGCCGCCTCTCCTGGCGAGACAATCACCA TCAACTGCCGGGCCAGCAAGAGCATCAACAA CTACCTCGTGTGGTATCAAGAGAAGCCCGGC AAGACCATCAAGCCCCTGATCTACAGCGGCA GCACACTGCAGACAGGCACCCCTAGCAGATT TTCCGGCAGCGGCTCTGGCACCGATTTTCAGC CTGACAATCAGCAGCCTGGAACCTGAGGACT TCGCCATGTACTACTGCCAGCAGCACAACGA GTACCCCTTCACCTTTGGAGCCGGCACCAAG CTGGAAGTCAAAGGCGGCGGAGGATCTGGC GGAGGTGGAAGTGGCGGAGGCGGATCTCAA GTTTCAGCTGCAGCAGTCCGATGCCGAGCTGG TTAAGCCTGGCGCCTCTGTGAAGATCAGCTG CAAGGCCAGCGGCTACACCTTCACCGATCAC GCCATCCACTGGGTCAAGCAGAAGCCTGAGC AGGGCCTCGAGTGGATCGGCTACTTTAGCCC CGGCAACGACGATGTGCGGTACAGCGAGAAG TTCAAGGGCAAAGCCACACTGACCGCCGACA AGAGCAGCAGCACTGCCTACATGCAGCTCAA CAGCCTGACCAGCGAGGACAGCGCCGTGTAC TTCTGCAAGAGATCCCTGCCTGGCGACTTCG ACTATTGGGGCCAGGGAACAACCCTGACCGT GTCCAGCACAAACAACCCTGCTCCTAGACCT CCTACACCAGCTCCTACAATCGCCTCTCAACC ACTGAGCCTGAGGCCAGAGGCTTGTAGACCA GCTGCTGGCGGAGCCGTGCATACAAGAGGAC TGGATTTGCGCTGCGACTTCTGGGTGCTCGT GGTTGTTGGCGGAGTGTGCTGGCCTGTTACTCT CTGCTGGTCACCGTGGCCTTCATCATCTTTTG GGTCCGAAGCAAGCGGAGCCGGCTGCTGCA CAGCGACTACATGAACATGACCCCTAGACGG CCCGGACCTACCAGAAAGCACTACCAGCCTT ACGCTCCTCCTAGAGACTTCGCCGCCTACCG GTCCAGAGTGAAGTTCAGCAGATCCGCCGAT GCTCCCGCCTATCAGCAGGGACAGAACCAGC TGTACAACGAGCTGAACCTGGGGAGAAGAGA AGAGTACGACGTGCTGGATAAGCGGAGAGGC AGAGATCCTGAGATGGGCGGCAAGCCCAGAC GGAAGAATCCTCAAGAGGGCCTGTATAATGA GCTGCAGAAAGACAAGATGGCCGAGGCCTAC TCCGAGATCGGAATGAAGGGCGAGCGCAGAA GAGGCAAGGGACACGATGGACTGTACCAGGG CCTGAGCACCGCCACCAAGGATACCTATGAT GCCCTGCACATGCAGGCCCTGCCTCCAAGAA GAAAGAGAGGCTCTGGCGAAGGCAGAGGTAG CCTGCTGACATGTGGCGACGTGGAAGAGAAC CCCGGACCAATGGTGTCCAAGGGCGAAGAGG ACAACATGGCCATCATCAAAGAATTCATGCGG TTCAAGGTGCACATGGAAGGCAGCGTGAACG GCCACGAGTTCGAGATTGAAGGCGAAGGCCGA GGGCAGACCTTACGAGGGAACACAGACCGCC AAGCTGAAAGTGACCAAAGGCGGCCCTCTGC CATTTGCCTGGGACATTCTGAGCCCTCAGTTT ATGTACGGCAGCAAGGCCCTACGTGAAGCACC CCGCCGATATTCCCGACTACCTGAAGCTGAG CTTCCCGGAGGGCTTCAAGTGGGAGAGAGTG	1-63=CD8A signal sequence 64-384=16E12 LC 385-429=Linker 430-780=16E12 HC 781-915=CD8a hinge 916-996=CD28 transmembrane domain 997-1119=intracellular signal domain 1120-1455=CD3-zeta intracellular chain 1456-2238=T2A-mCherry	337

Table 15: Nucleotide sequences encoding CARs			
	<p>ATGAACTTCGAGGACGGCGGCGTCGTGACCG TGACTCAAGATAGCTCTCTGCAGGACGGCGA GTTCATCTACAAAGTGAAGCTGCGGGGCACC AACTTTCCCTCTGATGGCCCCGTGATGCAGAA AAAGACCATGGGCTGGGAAGCCAGCAGCGAG AGAATGTACCCTGAAGATGGCGCCCTGAAAG GGGAGATCAAGCAGCGGCTGAAACTGAAGGA TGGCGGCCACTACGACGCCGAAGTGAAAACC ACCTACAAGGCCAAGAAACCCGTGCAGCTGC CAGGCGCCTACAACGTGAACATCAAGCTGGA CATCACCAGCCACAACGAGGACTACCCATC GTGGAACAGTACGAGAGAGCCGAAGGCAGG CACTCTACAGGCGGAATGGACGAGCTGTATA AGTAG</p>		
8H3-CART	<p>ATGGCTCTGCCGTTACAGCTCTGCTGCTGC CTCTGGCTCTGCTTCTGCATGCCGCTAGACC CGACATCCAGATGACCCAGACCACCAGCAGC CTGAGCGCCAGCCTGGGCGACAGAGTGACCA TCAGCTGCAGAGCCAGCCAGGACATCAGCCA CTACCTGAACTGGTACCAGCAGAAGCCCGAC GGCGCCGTGAAGCTGCTGATCTACAGCACCA GCAGACTGCACAGCGGCGTGCCAGCAGATT CAGCGGCAGCGGCAGCGGCACCGACTACAG CCTGACCATCAGCAACCTGGAGCAGGAGGAC ATCGCCACCTACTTCTGCCAGCAGGGCTACA CCCTGCCCTTACCTTCGGCAGCGGCACCAA GCTGGAGATCAAGGGCGGCGGAGGATCTGG CGGAGGTGGAAGTGGCGGAGGCGGATCTGA GGTGCAGCTGGTGGAGAGCGGCGGCGCCT GGTGAAGCCCGGCGGCAGCCTGAAGCTGAG CTGCGCCGCCAGCGGCTTACCTTCAGCAGC TACGCCATGAGCTGGGTGAGACAGAGCCCCG AGAGAAGACTGGAGTGGGTGGCCGAGATCAG CAGCGGCGGCAGCTACACCTACTACCCCGAC ACCGTGACCGGCAGATTACCATCAGCAGAG ACAACGCCAAGAACACCCTGTACCTGGAGAT GAGCAGCCTGAGAAGCGAGGACACCGCCATG TACTACTGCGCCAGAACCCTGGGCGAGGACT GGTACTTCGACGTGTGGGGCGCCGGCACCAC CGTGACCGTGAGCAGCAGCGGCACAACAACC CCTGCTCCTAGACCTCCTACACCAGCTCCTAC AATCGCCTCTCAACCTCTGTCTCTGCGGCCTG AGGCTTGTAGACCAGCTGCTGGCGGAGCCGT GCATACAAGAGGACTGGATTTGCGCTGCGAC TTCTGGGTGCTCGTGGTTGTTGGCGGAGTGC TGGCCTGTTACTCTCTGCTGGTCAACGTGGC CTTTCATCATCTTTTGGGTCCGAAGCAAGCGGA GCCGGCTGCTGCACAGCGACTACATGAACAT GACCCCTAGACGGCCCGGACCTACCAGAAAG CACTACCAGCCTTACGCTCCTCCTAGAGACTT CGCCGCTACCGGTCCAGAGTGAAGTTCAGC AGATCCGCGATGCTCCCGCCTATCAGCAGG GACAGAACCAGCTGTACAATGAGCTGAACCT GGGGCGCAGAGAAGAGTACGACGTGCTGGAT AAGCGGAGAGGCAGAGATCCTGAGATGGGC GGCAAGCCCAGACGGAAGAATCCTCAAGAGG GCCTGTATAACGAGCTGCAGAAAGACAAGAT GGCCGAGGCCTACAGCGAGATCGGAATGAAG GGCGAACGCAGAAGAGGCAAGGGCCACGAT GGACTGTATCAGGGCCTGAGCACCGCCACCA AGGATACCTATGATGCCCTGCACATGCAGGC CCTGCCTCCAAGAAGAAAGAGAGGCTCTGGC</p>	<p>1-63=CD8A signal sequence 64-384=8H3 LC 385-429=Linker 430-786=8H3 HC 787-921=CD8a hinge 922-1002=CD28 transmembrane domain 1003-1125=intracellular signal domain 1126-1460=CD3-zeta intracellular chain 1462-2244=T2A-mCherry</p>	338

Table 15: Nucleotide sequences encoding CARs		
	GAAGGCAGAGGTTAGCCTGCTGACATGTGGCG ACGTGGGAAGAGAACCCCGGACCAATGGTGTG CAAGGGCGAAGAGGACAACATGGCCATCATC AAAGAATTCATGCGGTTCAAGGTGCACATGGA AGGCAGCGTGAACGGCCACGAGTTCGAGATT GAAGGCGAAGGCGAGGGCAGACCTTACGAG GGAACACAGACCGCCAAGCTGAAAGTGACCA AAGGCGGCCCTCTGCCATTTGCCTGGGACAT TCTGAGCCCTCAGTTTATGTACGGCAGCAAG GCCTACGTGAAGCACCCCGCCGATATTCCCG ACTACCTGAAGCTGAGCTTCCCCGAGGGCTT CAAGTGGGAGAGAGTGTGAACCTTCGAGGAC GCGGGCGTCGTGACCGTGACTCAAGATAGCT CTCTGCAGGACGGCGAGTTCATCTACAAAGT GAAGCTGCGGGGCACCAACTTTCCCTCTGAT GGCCCCGTGATGCAGAAAAAGACCATGGGCT GGAAGCCAGCAGCGAGAGAATGTACCCTGA AGATGGCGCCCTGAAAGGGGAGATCAAGCAG CGGCTGAAACTGAAGGATGGCGGCCACTACG ACGCCGAAGTGAAAACCACCTACAAGGCCAA GAAACCCGTGCAGCTGCCAGGCGCCTACAAC GTGAACATCAAGCTGGACATCACCAGCCACA ACGAGGACTACACCATCGTGGAACAGTACGA GAGAGCCGAAGGCAGGCACTCTACAGGCGG AATGGACGAGCTGTATAAGTAG	

Table 16: Amino Acid sequences encoding CARs			
Construct	Amino Acid Sequence	Amino Acid Description	SEQ ID No.
15C4-CART	MALPVTALLLPLALLLHAARPNIVMTQSPKSMMS SVGERVTLSCASENVGIYVSWYQKPEQSPK LLIYGPSNRYTGVPDRFTGSGSATDFTLTISVQ AEDLADYHCGQSYSYPFTFGSGTKLEIKGGGGS GGGSGGGGSQVQLQQSDAELVKPGASVKISC KASGYTFDHAHWWKQKPEQGLEWIGYFSPG NGDIKYNEKFKGKATLTADKSSSTAYMQLNSLT SEDSAVYFCKRSLPGPMDCWGQTSVTVSSTT TPAPRPPTPAPTIASQPLSLRPEACRPAAGGAV HTRGLDFACDFWLVVVGGLVACYSLLVTVAFII FWVRSKRSLHSDYMNMTPRRPGPTRKHYQ PYAPPRDFAAYRSRVKFSRSADAPAYQQGQNG LYNELNLGRREEYDVLDKRRGRDPEMGGKPRR KNPQEGLYNELQKDKMAEAYSEIGMKGERRRG KGHDLQGLSTATKDTYDALHMQALPPRRKR GSGEGRGSLLTCGDVEENPGPMVSKGEEDNM AIIKEFMRFKVHMEGSVNGHEFEIEGEGEGRPY EGTQTAKLKVTKGGPLPFAWDILSPQFMYGSKA YVKHPADIPDYLKLSFPEGFKWERVMNFEDGG VVTVTQDSSLQDGEFIYKVKLRGTNFPDGPVM QKKTMGWEASSERMYPEDGALKGEIKQRLKLG DGGHYDAEVKTTYKAKKPVQLPGAYNVNIKLDIT SHNEDYTIVEQYERAEGRHSTGGMDELYK*	1-21=CD8a signal sequence 22-128=15C4 LC 129-143=Linker 144-260=15C4 HC 261-305=CD8a hinge 306-446=CD28 transmembrane 447-387=CD28 intracellular domain 388-499=CD3z intracellular domain 500-753=T2A mcherry	339
16E12-CART	MALPVTALLLPLALLLHAARPDVQISQSPSYLAA SPGETITINCRASKSINNYLVWYQEKPGKTIKPLI YSGSTLQTGTSPRFSGSGSDFSLTISSELEPE DFAMYYCQQHNEYPFTEGAGTKLELKGSGGSG GGGSGGGGSQVQLQQSDAELVKPGASVKISCK ASGYTFDHAHWWKQKPEQGLEWIGYFSPGN DDVRYSEKFKGKATLTADKSSSTAYMQLNSLTS	1-21=CD8a signal sequence 22-128=16E12 LC 129-143=Linker 144-260=16E12 HC	340

Table 16: Amino Acid sequences encoding CARs			
	EDSAVYFCKRSLPGDFDYWGQGTTLTVSSTTT PAPRPPTPAPTIASQPLSLRPEACRPAAGGAVH TRGLDFACDFWLVVGGVLACYSLLVTVAFIIF WRSKRSLRHSDYMNMTPRRPGPTRKHYP YAPPRDFAAYRSRVKFSRSADAPAYQQGQNL YNELNLGRREEYDVLDRRGRDPEMGGKPRRK NPQEGLYNELQKDKMAEAYSEIGMKGERRRGK GHDGLYQGLSTATKDTYDALHMQALPPRRKRG SGEGRGSLTTCGDVEENPGPMVSKGEEDNMAII KEFMRFKVHMEGSVNGHEFEIEGEGEGRPYEG TQTAKLKVTKGGPLPFAWDILSPQFMYGSKAYV KHPADIPDYLKLSFPEGFKWERVMNFEDGGVVT VTQDSSLQDGEFIYKVKLRGTNFPDGPVMQKK TMGWEASSERMYPEDGALKGEIKQRLKLDGG HYDAEVKTTYKAKKPVQLPGAYNVNIKLDITSHN EDYTIVEQYERAEGRHSTGGMDELYK*	261-305=CD8a hinge 306-446=CD28 transmembrane 447-387=CD28 intracellular domain 388-499=CD3z intracellular domain 500-753=T2A mcherry	
8H3-CART	MALPVTALLLPLALLLHAARPDVQITQSPSYLAA SPGETITINCRASKSVSEYLAWYQEKPGKTNKLL IYSGSTLHSGIPSRFSGSGSGTDFTLTITSLAPED FAMYFCQQHNEYPFTFGAGTKLELKGSGSGG GGSGGGGSQVQLQQSDAELVKPGASVKISCKA SGYTFDHAHWWKQRPEQGLEWIGYFSPGNG DIKYNEKFKDKATLTADKSSSTAYMQLNSLTSED SAVYFCKRSLPGDFDYWGQGTTLTVSSTTTTAP RPPTPAPTIASQPLSLRPEACRPAAGGAVHTRG LDFACDFWLVVGGVLACYSLLVTVAFIIFWVR SKRSRLLHSDYMNMTPRRPGPTRKHYPYAPP RDFAAAYRSRVKFSRSADAPAYQQGQNL YNELNLGRREEYDVLDRRGRDPEMGGKPRRK NPQEGLYNELQKDKMAEAYSEIGMKGERRRGK GHDGLYQGLSTATKDTYDALHMQALPPRRKRG SGEGRGSLTTCGDVEENPGPMVSKGEEDNMAII KEFMRFKVHMEGSVNGHEFEIEGEGEGRPYEG TQTAKLKVTKGGPLPFAWDILSPQFMYGSKAYV KHPADIPDYLKLSFPEGFKWERVMNFEDGGVVT VTQDSSLQDGEFIYKVKLRGTNFPDGPVMQKK TMGWEASSERMYPEDGALKGEIKQRLKLDGG HYDAEVKTTYKAKKPVQLPGAYNVNIKLDITSHN EDYTIVEQYERAEGRHSTGGMDELYK*	1-21=CD8a signal sequence 22-128=8H3 LC 129-143=Linker 144-262=8H3 HC 263-307=CD8a hinge 308-446=CD28 transmembrane 447-387=CD28 intracellular domain 388-499=CD3z intracellular domain 500-753=T2A mcherry	341

6.4.3. Results

[0441] CAR constructs were expressed in human T cells. Surface expression of CAR constructs was confirmed by flow cytometry using Alexa488-ProteinL. 8H3-CART specifically killed Tn+ COSMC-KO HaCaTs and Tn+ COSMC-KO A673s, but neither Tn- HaCaTs nor Tn-A673 (FIGS. 7A-7C). Table 17 summarizes the time to kill 50% Tn+ COSMC-KO HaCaTs. The time to kill 50% Tn+ COSMC-KO A673 was 9 hrs for 8H3-CART at the 3:1 ratio of T cells to HaCaTs. The time to kill 50% Tn+ COSMC-KO A673 was 5.73 hrs for 8H3-CART at the 5:1 ratio and 4.98 at the 10:1 ratio. The data indicates that 8H3-CART selectively targets CMET-Tn.

Table 17: Time to kill 50% of cells (KT50)			
Target Cell	T cell ratio	8H3 (KT50)	T Cells (KT50)
COSMC-KO HaCaT	3:1	9 hrs	N/A
COSMC-KO A673	5:1	5.73 hrs	N/A
COSMC-KO A673	10:1	4.98 hrs	N/A

6.5 Example 5: In vivo activity of cMET-CART in solid tumor mouse models

[0408] A CDx A549 solid tumor model was established by flank injection. The tumor volume at CART injection was 88 mm³. Mice were treated with 2nd generation 8H3-CAR-T by IT injection (2 doses at 1x10⁷ cells). Tumor volume was measured by caliper, 8H3-CAR-T treatment resulted in approximately 70% inhibition of tumor growth (FIG. 8A). No clinical signs indicating adverse events was observed in treated mice.

[0409] A Lung cancer solid tumor model (PDX) was established by flank injection (Champions model CTG-2823). The tumor volume at CART injection was 200 mm³ and CART was delivered by IV injection (4 doses at 1x10⁷ cells). Tumor volume was measured by caliper, 8H3-CAR-T treatment resulted in approximately 50% inhibition of tumor growth (FIG. 8B). No clinical signs indicating adverse events was observed in treated mice.

6.6 Example 6: Sequence Analysis of Anti-Glyco-CMET Antibodies

[0410] Rapid Amplification of cDNA Ends (RACE) was performed to determine the heavy chain and light chain nucleotide sequences for 15C4, 8H3, 16E12, 14E9, 19H2, and 39A3. The nucleotide sequences encoding the heavy and light chain variable regions of 15C4 are set forth in SEQ ID NO:21 and SEQ ID NO:22, respectively. The heavy and light chain variable regions encoded by SEQ ID NO:21 and SEQ ID NO:22 are set forth in SEQ ID NO:1 and SEQ ID NO:2, respectively. The predicted heavy chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:3-5, respectively, and the predicted light chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:6-8, respectively. The predicted heavy chain CDR sequences (Kabat definition) are set forth in SEQ ID NO:9-11, respectively, and the predicted light chain CDR sequences (Kabat definition) are set forth in SEQ ID NO:12-14, respectively. The predicted heavy chain CDR sequences (Chothia definition) are set forth in SEQ ID NO:15-17, respectively, and the predicted light chain CDR sequences (Chothia definition) are set forth in SEQ ID NO:18-20, respectively.

[0411] The nucleotide sequences encoding the heavy and light chain variable regions of 8H3 are set forth in SEQ ID NO:43 and SEQ ID NO:44, respectively. The heavy and light chain variable regions encoded by SEQ ID NO:43 and SEQ ID NO:44 are set forth in SEQ ID NO:23 and SEQ ID NO:24, respectively. The predicted heavy chain CDR sequences (IMGT definition)

are set forth in SEQ ID NOS:25-27, respectively, and the predicted light chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:28-30, respectively. The predicted heavy chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:31-33, respectively, and the predicted light chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:34-36, respectively. The predicted heavy chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:37-39, respectively, and the predicted light chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:40-42, respectively.

[0412] The nucleotide sequences encoding the heavy and light chain variable regions of 16E12 are set forth in SEQ ID NO:65 and SEQ ID NO:66, respectively. The heavy and light chain variable regions encoded by SEQ ID NO:65 and SEQ ID NO:66 are set forth in SEQ ID NO:45 and SEQ ID NO:46, respectively. The predicted heavy chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:47-49, respectively, and the predicted light chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:50-52, respectively. The predicted heavy chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:53-55, respectively, and the predicted light chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:56-58, respectively. The predicted heavy chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:59-61, respectively, and the predicted light chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:62-64, respectively.

[0413] The nucleotide sequences encoding the heavy and light chain variable regions of 14E9 are set forth in SEQ ID NO:87 and SEQ ID NO:88, respectively. The heavy and light chain variable regions encoded by SEQ ID NO:87 and SEQ ID NO:88 are set forth in SEQ ID NO:67 and SEQ ID NO:68, respectively. The predicted heavy chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:69-71, respectively, and the predicted light chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:72-74, respectively. The predicted heavy chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:75-77, respectively, and the predicted light chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:78-80, respectively. The predicted heavy chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:81-83, respectively, and the predicted light chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:84-86, respectively.

[0414] The nucleotide sequences encoding the heavy and light chain variable regions of 19H2 are set forth in SEQ ID NO:109 and SEQ ID NO:110, respectively. The heavy and light chain variable regions encoded by SEQ ID NO:109 and SEQ ID NO:110 are set forth in SEQ ID NO:89 and SEQ ID NO:90, respectively. The predicted heavy chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:91-93, respectively, and the predicted light chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:94-96, respectively. The predicted heavy chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:97-99, respectively, and the predicted light chain CDR sequences (Kabat definition) are set forth in

SEQ ID NOS:100-102, respectively. The predicted heavy chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:103-105, respectively, and the predicted light chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:106-108, respectively.

[0415] The nucleotide sequences encoding the heavy and light chain variable regions of 19H2 are set forth in SEQ ID NO:131 and SEQ ID NO:132, respectively. The heavy and light chain variable regions encoded by SEQ ID NO:131 and SEQ ID NO:132 are set forth in SEQ ID NO:111 and SEQ ID NO:112, respectively. The predicted heavy chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:113-115, respectively, and the predicted light chain CDR sequences (IMGT definition) are set forth in SEQ ID NOS:116-118, respectively. The predicted heavy chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:119-121, respectively, and the predicted light chain CDR sequences (Kabat definition) are set forth in SEQ ID NOS:122-124, respectively. The predicted heavy chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:125-127, respectively, and the predicted light chain CDR sequences (Chothia definition) are set forth in SEQ ID NOS:128-130, respectively.

6.7 Example 7: Humanized Antibodies

6.7.1. Overview

[0416] The murine antibody 8H3 was humanized using standard CDR-grafting technology. For the heavy chain, four templates, IGHV1-3*01, IGHV5-51*01, IGHV7-4-1*02, and IGHV1-69*06 were employed in order to generate CDR-grafted versions containing successively aggressive levels of humanization, *i.e.*, identity to the human acceptor germline. Similarly for the light chain, three templates, IGKV1-9*01, IGKV3-15*01, and IGKV6-21*01, were employed to generate CDR-grafted versions containing successively aggressive levels of humanization.

[0417] Expression constructs were designed for expression in Expi-293 cells. IL2 secretion signals were added to both heavy and light chain constructs. Antibodies were purified with ProteinA beads using conventional methods. Humanized candidates were evaluated for their ability to binding to the non-glycosylated and Tn-glycosylated cMET peptides using ELISA. The humanized candidates were also compared to the parental antibody by size exclusion chromatography, Octet to determine binding affinity to the peptide antigen, and cell binding to target positive cells using flow cytometry.

6.7.2. Materials and Methods

6.7.2.1 Vector Design

[0418] For each germline, three humanize versions were created: a conservative "A" sequence, a less conservative "B" sequence, and an "aggressive" "C" sequence (see Tables 4A-4G). Consensus sequences of all three of the A, B, and C sequences for each germline were also created that reflect the most common amino acid residue at each position.

[0419] These humanized templates were assembled and assayed for optimal biophysical and functional properties in two phases. In the first phase, up to 12 pairs of the conservative “A” designs were constructed and assayed for binding to the cMET glycopeptide. After selection of the most optimal combination based upon the “A” designs, the conservative “A” designs were iteratively replaced with the less conservative “B” designs and ultimately with the least conservative “C” designs.

6.7.2.2 ELISA

[0420] 96-well Corning high bind ELISA microplates plates were coated with cMET peptides titrated in 0.2 M bicarbonate buffer, pH 9.4 overnight at 4 °C in concentrations ranging from 0.08 µg/ml to 10 µg/ml. BSA was used as a control/measure of background. The plates were then blocked with SuperBlock™ (Thermo Fisher) for 1 hr at room temperature. After plate washing, the humanized variants of 8H3 were incubated on the ELISA plate for 1 hour. All tested variants were expressed and purified using conventional methods. Briefly, Expi-293 cells were transiently transfected with heavy and light chain constructs, antibodies were secreted into supernatant and purified using Protein A agarose beads. The plates were then washed, and then incubated with secondary antibody (1/3000 Goat Anti-mouse IgG (H+L) HRP (Abcam 62-6520)) for 1 hour. The plate was then washed and color was developed with 1-Step™ Ultra TMB (Thermo Fisher) for 2 minutes. Color development was then stopped with 2 N Sulfuric Acid. Absorbance at 450 nm was then measured.

6.7.2.3 Bio-Layer Interferometry (Octet)

[0421] Antibody affinity of the humanized candidates of 8H3 can be assessed against specific antigens using BLI. In a BLI assay, the antigen can be immobilized onto a biosensor (e.g., the cMET-Tn peptide Biotin-PTKSFISGGSTITGVGKLN (the amino acid portion of which is SEQ ID NO:285) or a negative control analyte such as an unglycosylated cMET peptide (Biotin-PTKSFISGGSTITGVGKLN (the amino acid portion of which is SEQ ID NO:286)) and presented to one antibody candidate for affinity measurements or two competing antibodies in tandem (or consecutive steps) for epitope binning. The binding to non-overlapping epitopes occurs if saturation with the first antibody does not block the binding of the second antibody. The affinity is determined by fitting the binding curve to a specific model: a 1:1 monovalent model or a 2:1 bivalent model. The error (>95% confidence) is calculated by how close the generated curve matches the model.

6.7.2.4 Size Exclusion Chromatography

[0422] The humanized candidates for 8H3 were tested for the presence of soluble protein aggregates using size exclusion chromatography (SEC). Briefly, purified antibodies were loaded on an HPLC silica TSK-GEL G3000SW column (TOSOH Biosciences, Montgomeryville, PA) and associated UV detector (166 Detector). The mobile phase composition was PBS and

flow rate was 1.0 mL/min. Concentrations of protein species were determined by monitoring the absorbance of column eluate at 280 nm.

6.8 Example 8: Humanized 8H3-based CAR

6.8.1. Overview

[0423] A chimeric antigen receptor (CAR) having VH and VL domains of humanized 8H3 was designed. The CAR was then evaluated in a target-specific cytotoxicity assay.

6.8.2. Materials and Methods

[0424] A CAR construct having a scFv with 8H3-HV1-3-A VH (SEQ ID NO:264) and 8H3-KV1-A VL (SEQ ID NO:276) domains was designed (hu8H3-CART). In the construct, the VH and VL are attached together with one long linker (GGGS)₃ (SEQ ID NO:346) to the CD8a hinge followed by a second generation CAR-T (CD28 intracellular signal domain, and a CD3-zeta intracellular chain). The N-terminus of the scFvs was attached to a CD8a signal sequence. Nucleotide and amino acid sequences for the CAR are provided in Table 18.

Table 18: Nucleotide and amino acid sequences of hu8H3-CART			
Construct	Sequence	Description	SEQ ID No.
hu8H3-CART (nucleotide sequence)	ATGGCTCTGCCCGTTACAGCTCTGCTGCTGC CTCTGGCTCTGCTTCTGCATGCCGCTAGACC CGACGTGCAGATTACCCAGTCTCCTAGCTTT CTGAGCGCCAGCGTGGGCGACAGAGTGACC ATTACATGCAGGGCCAGCAAGAGCGTGTCC GAGTACCTGGCCTGGTATCAAGAGAAGCCC GGCAAGGCCAACCAAGCTGCTGATCTACAGC GGCAGCACACTGCACAGCGGAGTGCCTAGC AGATTTTCCGGCAGCGGCTCTGGCACCGAG TTCACCCTGACCATATCTAGCCTGCAGCCTG AGGACTTTGCCACCTACTTTTGGCAGCAGCA CAACGAGTACCCCTTACCTTTGGCCAGGGC ACCAAGCTGGAATCAAAGGCGGCGGAGGA TCTGGCGGAGGTGGAAGTGGCGGAGGCGG ATCTCAAGTTCAGCTGGTTCAGTCTGGCGCC GAAGTGAAGAAACCTGGCGCCTCTGTGAAG GTGTCCCTGCAAGGCCAGCGGCTACACCTTTA CCGATCACGCCATCCACTGGGTCCGACAGG CTCCAGGACAACGGCTGGAATGGATCGGCT ACTTCAGCCCCGGCAACGGCGACATCAAGTA CAACGAGAAGTTCAAGGACCGGGCCACACT GACCGCCGATAAGTCTGCCAGCACCGCCTA CATGGAAGTGTCCAGCCTGAGAAGCGAGGA TACCGCCGTGTAATTCTGCAAGAGATCCCTG CCTGGCGACTTCGACTATTGGGGCCAGGGA ACACTGGTCACCGTGTCCAGCACAAACAACC CTGCTCCTAGACCTCCTACACCAGCTCCTAC AATCGCCTCTCAACCTCTGTCTCTGCGCCT GAGGCTTGTAGACCAGCTGCTGGCGGAGCC GTGCATACAAGAGGACTGGATTTGCGCTGCG ACTTCTGGGTGCTCGTGGTTGTTGGCGGAGT GCTGGCCTGTTACTCTCTGCTGGTCCACAGTG GCCTTCATCATCTTTTGGGTCCGAAGCAAGC GGAGCCGGCTGCTGCACTCCGACTACATGA ACATGACCCCTAGACGGCCCGGACCTACCA GAAAGCACTACCAGCCTTACGCTCCTCCTAG	1-63=CD8a signal sequence 64-384= 8H3-KV1-A LC 385-429=Linker 430-780= 8H3-HV1-3-A HC 781-915=CD8a hinge 916-996=CD28 transmembrane 997-1119=CD28 intracellular domain 1120-1455=CD3z intracellular domain 1456-2235=T2A mcherry	347

Table 18: Nucleotide and amino acid sequences of hu8H3-CART			
	<p>AGACTTCGCCGCCTACCGGTCCAGAGTGAA GTTCAGCAGATCCGCCGATGCTCCCGCCTAT CAGCAGGGACAGAATCAGCTGTACAATGAGC TGAACCTGGGGCGCAGAGAAGAGTACGACG TGCTGGATAAGCGGAGAGGCAGAGATCCTG AGATGGGCGGCAAGCCCAGACGGAAGAATC CTCAAGAGGGCCTGTATAACGAGCTGCAGAA AGACAAGATGGCCGAGGCCTACAGCGAGAT CGGAATGAAGGGCGAACGCAGAAGAGGCAA GGGCCACGATGGACTGTATCAGGGCCTGAG CACCGCCACCAAGGATACCTATGATGCCCTG CACATGCAGGCCCTGCCTCCAAGAAGAAAGA GAGGCTCTGGCGAAGGCAGAGGTAGCCTGC TGACATGTGGCGACGTGGAAGAGAACCCCG GACCAATGGTGTCCAAGGGCGAAGAGGACA ACATGGCCATCATCAAAGAATTCATGCGGTT CAAGGTGCACATGGAAGGCAGCGTGAACGG CCACGAGTTCGAGATTGAAGGCGAAGGCGA GGGCAGACCTTACGAGGGAACACAGACCGC CAAGCTGAAAGTGACCAAAGGCGGACCCCT GCCTTTCGCCTGGGATATCCTGTCTCCTCAG TTTATGTACGGCAGCAAGGCCTACGTGAAGC ACCCGCCGATATTCCCGACTACCTGAAGCT GAGCTTCCCCGAGGGCTTCAAGTGGGAGAG AGTGATGAACTTCGAGGACGGCGGCGTCGT GACCGTGACTCAAGATAGCTCTCTGCAGGAC GCGGAGTTCATCTACAAAGTGAAGCTGCGG GGCACCAACTTTCCCTCTGATGGCCCCGTGA TGCAGAAAAAGACCATGGGCTGGGAAGCCA GCAGCGAGAGAAATGTACCCTGAAGATGGCG CCCTGAAAGGCGAGATCAAGCAGCGGCTGA AACTGAAAGGATGGCGGCCACTACGACGCTG AAGTGA AAAACCACCTACAAGGCCAAGAAACC CGTG CAGCTGCCAGGCGCCTACAACGTGAA CATCAAGCTGGACATTACCAGCCACAACGAG GACTACACCATCGTGGAACAGTACGAGAGAG CCGAAGGCAGGCACTCTACAGGCGGAATGG ACGAGCTGTATAAGTAG</p>		
hu8H3-CART (amino acid sequence)	<p>MALPVTALLLPLALLLHAARPDVQITQSPSFLSA SVGDRVITICRASKSVSEYLAWYQEKPGKANK LLIYSGSTLHSGVPSRFSGSGSTEFTLTISSLQ PEDFATYFCQQHNEYPTFFGQGTKLEIKGGGG SGGGSGGGGSQVLVQSGAEVKKPGASVK VSCKASGYTFTDHAHWVRQAPGQRLEWIGYF SPGNQDIKYNEKFKDRATLTADKSASTAYMEL SSLRSEDTAVYFCKRSLPGLDFDYWGQTLVTV SSTTTPAPRPPTPAPTIASQPLSLRPEACRPAA GGAVHTRGLDFACDFWLVVVGVLACYLLV TVAFIIFWVRSKRSRLLHSDYMNMTPRRPGPT RKHYQPYAPPRDFAAYRSRVKFSRSADAPAY QQGQNQLYNELNLGRREEYDVLDKRRGRDPE MGKPRRKNPQEGLYNELQKDKMAEAYSEIG MGERRRGKGHDGLYQGLSTATKDTYDALHM QALPPRRKRGSGEGRGSLLTCGDVEENPGPM VSKGEEDNMAIIEFMRFKVHMEGSVNGHEFEI EGEGERPYEGTQTAKLKVTKGGPLPFAWDIL SPQFMYGSKAYVKHPADIPDYLLKLSFPEGFKW ERVMNFEDGGVTVTQDSSLQDGEFIYKVKLR GTNFPSDGPVMQKKTMGWEASSERMYPEDG ALKGEIKQRLKLDGGHYDAEVKTTYKAKKPV</p>	<p>1-21=CD8a signal sequence 22-128= 8H3-KV1-A LC 129-143=Linker 144-260= 8H3-HV1-3-A HC 261-305=CD8a hinge 306-332=CD28 transmembrane 333-373=CD28 intracellular domain 374-485=CD3z intracellular domain 486-746=T2A mcherry</p>	348

Table 18: Nucleotide and amino acid sequences of hu8H3-CART			
	QLPGAYNVNIKLDITSHNEDYTIVEQYERAEGR		
	HSTGGMDELYK		

[0425] hu8H3-CART cells were incubated with A673 (Tn+) and A673 (Tn-) cells at an effector cell:target cell (E:T) ratio of 2:1 and cytotoxicity was monitored in real-time using noninvasive electrical impedance on a RTCA iCELLigence™ instrument.

6.8.3. Results

[0426] 100% of Tn+ cells were specifically killed over six hours with hu8H3-CART (FIG. 10). KT50 (time to kill 50% of target cells) for hu8H3-CART on A673 (Tn+) cells was determined to be 1 hour and 15 minutes.

7. SPECIFIC EMBODIMENTS, CITATION OF REFERENCES

[0427] While various specific embodiments have been illustrated and described, it will be appreciated that various changes can be made without departing from the spirit and scope of the disclosure(s). The present disclosure is exemplified by the numbered embodiments set forth below.

1. An anti-glyco-cMET antibody or antigen binding fragment that specifically binds to a cMET peptide PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285) that has been glycosylated with GalNAc on the serine and threonine residues shown with bold and underlined text (“the cMET glycopeptide”).

2. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNGDIKYNE KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWQGQTSVTVSS(SEQ ID NO:1) and a light chain variable (VL) sequence of NIVMTQSPKSMMSGVGERVTLCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF TGSGSATDFTLTISVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2) for binding to the cMET glycopeptide.

3. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQRPEQGLEWIGYFSPGNGDIKYNE KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID NO:23) and a light chain variable (VL) sequence of DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPGKTNKLLIYSGSTLHSGIPSRFSG

SGSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24) for binding to the cMET glycopeptide.

4. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNDVRYSEKFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTTLTVSS (SEQ ID NO:45) and a light chain variable (VL) sequence of DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKPGKTIKPLIYSGSTLQTGTPSRFSGSGSGTDFSLTISLEPEDFAMYQCQQHNEYPTFGAGTKLELK (SEQ ID NO:46) for binding to the cMET glycopeptide.

5. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWGCITYTSGGGNTYYATWAKGRFTVSETSSSTTVLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGLTVTVSSGQPK (SEQ ID NO:67) and a light chain variable (VL) sequence of DVVMTQTPASVGA AVGGTVTIKCQASQSISNWLAWYQQKPGQPPKLLIYSASYLESQVPSRFSGSGSGTEFTLTISDLECAATYYCQCTYGSSGDSGSWDFGGGTEVVVKGDPV (SEQ ID NO:68) for binding to the cMET glycopeptide.

6. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDYANWARGRFTISKSSPTVDLKMSTLTGADTGTYFCARMGYEDGYVGGVYITVGAFNLWGQGLTVTVSSGQPK (SEQ ID NO:89) and a light chain variable (VL) sequence of DVVMTQTASPVSA AVGGTVTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESQVPSRFGSGSGTQFTLTISGVQCDDAATYYCQCSYSGSGYSWTFGGGTEVVVKGDPV (SEQ ID NO:90) for binding to the cMET glycopeptide.

7. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKGLEWIACMDNRVYATWAKGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNLWGQGLTVTVSSGQPK (SEQ ID NO:111) and a light chain variable (VL) sequence of DPVLTQTPPSVSA AVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPSRFKSGSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFSGGTEVVVKGDPV (SEQ ID NO:112) for binding to the cMET glycopeptide.

8. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPQKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

9. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

10. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPKAPKLLIYASSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:278) for binding to the cMET glycopeptide.

11. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:279) for binding to the cMET glycopeptide.

12. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody

or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:280) for binding to the cMET glycopeptide.

13. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:281) for binding to the cMET glycopeptide.

14. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

15. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

16. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN

EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSGSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

17. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYSQKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRVITICRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

18. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYSQKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVITICRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

19. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYSQKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVITICRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:278) for binding to the cMET glycopeptide.

20. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYSQKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of

EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWEYQEKPGQANRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:279) for binding
to the cMET glycopeptide.

21. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWEYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:280) for binding
to the cMET glycopeptide.

22. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWEYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:281) for binding
to the cMET glycopeptide.

23. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of
EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWEYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding
to the cMET glycopeptide.

24. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWEYQQKPDQSPKLLIYSGSTLHSGVPSRFS

GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

25. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYS QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSEYLAHWYQQKPDQSPKLLIYSGSTLFSGVPSRFSGSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

26. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRTITCRASKSVSEYLAHWYQEKPGKANKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTFGQGKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

27. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAHWYQQKPGKAPKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

28. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSISEYLAHWYQQKPGKAPKLLIYASSTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGKLEIK (SEQ ID NO:278) for binding to the cMET glycopeptide.

29. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:279) for binding to the cMET glycopeptide.

30. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:280) for binding to the cMET glycopeptide.

31. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:281) for binding to the cMET glycopeptide.

32. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

33. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody

or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

34. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

35. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYFSPGNNGDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:267) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

36. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYFSPGNNGDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:267) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

37. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYFSPGNNGDIKYN

EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:278) for binding to the cMET glycopeptide.

38. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of EWITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:279) for binding to the cMET glycopeptide.

39. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:280) for binding to the cMET glycopeptide.

40. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:281) for binding to the cMET glycopeptide.

41. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding
to the cMET glycopeptide.

42. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLVTVSS(SEQ ID
NO:267) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for
binding to the cMET glycopeptide.

43. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLVTVSS(SEQ ID
NO:267) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
SGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:284) for binding
to the cMET glycopeptide.

44. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
QKFKGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLVTVSS(SEQ ID
NO:268) and a light chain variable (VL) sequence of
DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPQKANKLLIYSGSTLHSGVPSRFS
GSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:276) for binding
to the cMET glycopeptide.

45. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
QKFKGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLVTVSS(SEQ ID
NO:268) and a light chain variable (VL) sequence of
DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS

GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:277) for binding to the cMET glycopeptide.

46. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGRVTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSGSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:278) for binding to the cMET glycopeptide.

47. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:279) for binding to the cMET glycopeptide.

48. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:280) for binding to the cMET glycopeptide.

49. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:281) for binding to the cMET glycopeptide.

50. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNNGDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:268) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

51. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNNGDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:268) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

52. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNNGDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:268) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

53. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFSDHAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPQKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

54. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody

or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFS D HAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of DIQLTQSPSFLSASV GDRVTITCRASKSVSEYLA WYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISLQPEDFATYYCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

55. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFS D HAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of DIQLTQSPSFLSASV GDRVTITCRASKSISEYLA WYQQKPGKAPKLLIY SASTLHSGVPSRFS GSGSGTEFTLTISLQPEDFATYYCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:278) for binding to the cMET glycopeptide.

56. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFS D HAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLA WYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:279) for binding to the cMET glycopeptide.

57. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFS D HAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:280) for binding to the cMET glycopeptide.

58. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSCASGYTFS D HAIHWWRQAPGQGLEWIGYFSPGNADINYA

QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:281) for binding to the cMET glycopeptide.

59. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFSDHAIHWVRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EWITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQKPDQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

60. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFSDHAIHWVRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

61. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFSDHAIHWVRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFS GSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

62. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFSDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of

DVQITQSPSFLSASVGDRVTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS
GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:276) for binding
to the cMET glycopeptide.

63. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:277) for binding
to the cMET glycopeptide.

64. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIY SASTLHSGVPSRFSG
S GSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:278) for binding
to the cMET glycopeptide.

65. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:279) for binding
to the cMET glycopeptide.

66. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS

GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:280) for binding to the cMET glycopeptide.

67. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:281) for binding to the cMET glycopeptide.

68. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLA WYQEKPQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

69. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLA WYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

70. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLA WYQQKPDQSPKLLIYSGSTLFSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:284) for binding to the cMET glycopeptide.

71. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRVTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:276) for binding to the cMET glycopeptide.

72. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:277) for binding to the cMET glycopeptide.

73. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSG SSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:278) for binding to the cMET glycopeptide.

74. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:279) for binding to the cMET glycopeptide.

75. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody

or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:280) for binding to the cMET glycopeptide.

76. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:281) for binding to the cMET glycopeptide.

77. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

78. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNGDIKYNE KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYCCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

79. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNGDIKYNE

KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:271) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSGSGSGTDFLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:284) for binding to the cMET glycopeptide.

80. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSEKFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:272) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRVTITCRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:276) for binding to the cMET glycopeptide.

81. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSEKFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:272) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:277) for binding to the cMET glycopeptide.

82. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSEKFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:272) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:278) for binding to the cMET glycopeptide.

83. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSEKFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:272) and a light chain variable (VL) sequence of

EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWEYQEKPGQANRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:279) for binding
to the cMET glycopeptide.

84. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:272) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWEYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:280) for binding
to the cMET glycopeptide.

85. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:272) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWEYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:281) for binding
to the cMET glycopeptide.

86. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:272) and a light chain variable (VL) sequence of
EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWEYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTIINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding
to the cMET glycopeptide.

87. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:272) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWEYQQKPDQSPKLLIYSGSTLHSGVPSRFS

GS GSGTDFTLTINSLEAEDAATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

88. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSEKFQGVQVTSADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:272) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSIS SYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFS GSGTDFTLTINSLEAEDAATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

89. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKV SCKASGYTFTDHAIHWWRQAPGQGLEWIGYFSPGN GDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRVTITCRASKSVSEYLA WYQEKPGKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYFCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

90. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKV SCKASGYTFTDHAIHWWRQAPGQGLEWIGYFSPGN GDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLA WYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

91. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKV SCKASGYTFTDHAIHWWRQAPGQGLEWIGYFSPGN GDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSISEYLA WYQQKPGKAPKLLIYSASTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:278) for binding to the cMET glycopeptide.

92. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLA WYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYFCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:279) for binding to the cMET glycopeptide.

93. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:280) for binding to the cMET glycopeptide.

94. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:281) for binding to the cMET glycopeptide.

95. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLA WYQEKPQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

96. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody

or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

97. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG SSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

98. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYISTGNGDIKYNQ KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRVTITCRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLPEDFATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:276) for binding to the cMET glycopeptide.

99. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYISTGNGDIKYNQ KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLPEDFATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:277) for binding to the cMET glycopeptide.

100. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYISTGNGDIKYNQ

KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTLTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:278) for binding to the cMET glycopeptide.

101. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCKASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTLTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of EWITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:279) for binding to the cMET glycopeptide.

102. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCKASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTLTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:280) for binding to the cMET glycopeptide.

103. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCKASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTLTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:281) for binding to the cMET glycopeptide.

104. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCKASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTLTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282) for binding
to the cMET glycopeptide.

105. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ
KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:274) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for
binding to the cMET glycopeptide.

106. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ
KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:274) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
SGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:284) for binding
to the cMET glycopeptide.

107. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of
DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPQKANKLLIYSGSTLHSGVPSRFS
GSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:276) for binding
to the cMET glycopeptide.

108. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1,
wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody
or antigen binding fragment comprising a heavy chain variable (VH) sequence of
QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of
DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS

GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:277) for binding to the cMET glycopeptide.

109. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSG GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:278) for binding to the cMET glycopeptide.

110. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EVWITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:279) for binding to the cMET glycopeptide.

111. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:280) for binding to the cMET glycopeptide.

112. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:281) for binding to the cMET glycopeptide.

113. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAIHWRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding to the cMET glycopeptide.

114. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAIHWRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to the cMET glycopeptide.

115. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAIHWRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG SSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:284) for binding to the cMET glycopeptide.

116. The anti-glyco-cMET antibody or antigen binding fragment of any one of embodiments 1 to 115, which specifically binds to COSMC knock-out T47D cells.

117. The anti-glyco-cMET antibody or antigen binding fragment of any one of embodiments 1 to 115, which specifically binds to COSMC knock-out A549 cells.

118. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQKPEQGLEWIGYFSPGNGDIKYNE KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVTVSS (SEQ ID NO:1) and a light chain variable (VL) sequence of

NIVMTQSPKSMSSVGERVTLSCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF TSGSATDFLTISSVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

119. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWKQRPEQGLEWIGYFSPGNGDIKYNE KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLVSS(SEQ ID NO:23) and a light chain variable (VL) sequence of DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKP GKTKNLLIYSGSTLHSGIPSRFSG SGGTDFLTITSLAPEDFAMYFCQQHNEY PFTFGAGTKLELK (SEQ ID NO:24) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

120. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWKQKPEQGLEWIGYFSPGNDVRYSE KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLVSS(SEQ ID NO:45) and a light chain variable (VL) sequence of DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKP GKTIKPLIYSGSTLQTGTSPSRFSGS GSGTDFSLTISLEPEDFAMYFCQQHNEY PFTFGAGTKLELK(SEQ ID NO:46) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

121. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAIHWVRQAPGQRLEWIGYFSPGNGDIKYN EKFKDRATLTADKSASTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:264) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISLQPEDFATYFCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:276) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

122. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAIHWVRQAPGQRLEWIGYFSPGNGDIKYN

EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:264) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

123. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYN
 EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:264) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSG
 SSGSGTEFTLTISSLQPEDFATYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

124. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYN
 EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:264) and a light chain variable (VL) sequence of
 EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

125. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYN
 EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:264) and a light chain variable (VL) sequence of
 EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:280) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

126. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable

(VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN
EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:264) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:281) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

127. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN
EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:264) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

128. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN
EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:264) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for
binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

129. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYN
EKFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
NO:264) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
SGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:284) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

130. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
 QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:276) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

131. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
 QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

132. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
 QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSG
 SSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

133. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNGDIKYS
 QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of
 EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS

GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

134. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAIHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:280) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

135. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAIHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:281) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

136. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAIHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:265) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

137. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAIHWVRQAPGQRLEWIGYFSPGNGDIKYS
QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID

NO:265) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
 GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for
 binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

138. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNQDIKYS
 QKFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 SSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:284) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

139. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS
 QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:276) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

140. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS
 QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

141. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:266) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSG SSGSGTEFTLTISLQPEDFATYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

142. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:266) and a light chain variable (VL) sequence of EVWITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

143. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:266) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:280) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

144. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:266) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK(SEQ ID NO:281) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

145. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment

competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS
 QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:266) and a light chain variable (VL) sequence of
 EVWITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS
 GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

146. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS
 QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:266) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
 GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:283) for
 binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

147. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGASVKVSCASGYTFTDHAHWVRQAPGQRLEWIGYFSPGNADTKYS
 QKFQGRVTITADKSASTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:266) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 SSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:284) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

148. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYFSPGNNGDIKYN
 EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:267) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPQKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:276) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

149. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFTD HAIHW RQAPGQGLEWIGYFSPGN GDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of DIQLTQSPSFLSASV GDRVTITCRASKSVSEYLA WYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

150. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFTD HAIHW RQAPGQGLEWIGYFSPGN GDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of DIQLTQSPSFLSASV GDRVTITCRASKSISEYLA WYQQKPGKAPKLLIY SASTLHSGVPSRFSG SSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

151. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFTD HAIHW RQAPGQGLEWIGYFSPGN GDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLA WYQEKPGQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYFCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

152. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFTD HAIHW RQAPGQGLEWIGYFSPGN GDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS

GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:280) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

153. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:281) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

154. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

155. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:267) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

156. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN EKFKDRATLTADKSTSTAYMELSSLRSEDTAVYFCKRSLPGDFDYWGQGLTVTVSS(SEQ ID

NO:267) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 SGGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGKLEIK(SEQ ID NO:284) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

157. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
 QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFGQGKLEIK(SEQ ID NO:276) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

158. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
 QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGKLEIK(SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

159. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVCKASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN
 QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSG
 SGGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGKLEIK(SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

160. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQGANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

161. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:280) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

162. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK(SEQ ID NO:281) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

163. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNQDIKYN QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID NO:268) and a light chain variable (VL) sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

164. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment

competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNNGDIKYN
 QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:268) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
 GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for
 binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

165. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNNGDIKYN
 QKFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:268) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 SSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:284) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

166. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVSCASGYTFSDHAHWWRQAPGQGLEWIGYFSPGNADINYA
 QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
 NO:269) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:276) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

167. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGAEVKKPGSSVKVSCASGYTFSDHAHWWRQAPGQGLEWIGYFSPGNADINYA
 QKFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
 NO:269) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:277) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

168. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFS DHAHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGRVTITCRASKSISEYLA WYQQKPGKAPKLLIYSASTLHSGVPSRFSG SSGSGTEFTLTISLQPEDFATYYCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

169. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFS DHAHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EVWITQSPATLSVSPGERATLSCRASKSVSEYLA WYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

170. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFS DHAHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPFTFGQGTKLEIK (SEQ ID NO:280) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

171. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKV SCKASGYTFS DHAHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLA WYQQKPGQAPRLLIYSGSTLHSGIPARFS

GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:281) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

172. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFS DHAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EVWITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

173. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFS DHAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

174. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFS DHAIHWWRQAPGQGLEWIGYFSPGNADINYA QKFQGRVTITADKSTSTAYMELSSLRSEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269) and a light chain variable (VL) sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:284) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

175. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTD HAIHWWRQMPGKGLEWIGYFSPGNNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID

NO:270) and a light chain variable (VL) sequence of DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:276) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

176. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

177. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIY SASTLHSGVPSRFSG SSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

178. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAHWVRQMPGKGLEWIGYFSPGNGDIKYNE KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGT LVTVSS (SEQ ID NO:270) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

179. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:280) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

180. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:281) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

181. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

182. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of
EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:270) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGTKLEIK (SEQ ID NO:283) for
binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

183. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment

competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
 KFKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGTLVTVSS (SEQ ID
 NO:270) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 SSGSGTDFTLTINSLEAEDAATYYCQQHNEYPFQGGTKLEIK(SEQ ID NO:284) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

184. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
 KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID
 NO:271) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRVTITCRASKSVSEYLAWYQEKPGKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPFQGGTKLEIK(SEQ ID NO:276) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

185. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
 KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID
 NO:271) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPFQGGTKLEIK(SEQ ID NO:277) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

186. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
 KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS (SEQ ID
 NO:271) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSG
 SSGSGTEFTLTISSLQPEDFATYYCQQHNEYPFQGGTKLEIK(SEQ ID NO:278) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

187. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPAGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:271) and a light chain variable (VL) sequence of
EVIWQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:279) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

188. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPAGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:271) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:280) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

189. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPAGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:271) and a light chain variable (VL) sequence of
EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:281) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

190. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPAGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:271) and a light chain variable (VL) sequence of
EVIWQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS

GS GSGTDFTLTINSLEAEDAATYFCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:282) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

191. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNNDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:271) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLA WYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GS GSGTDFTLTINSLEAEDAATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:283) for
binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

192. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNNDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:271) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLA WYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
SGSGTDFTLTINSLEAEDAATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:284) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

193. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
KFQGGVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:272) and a light chain variable (VL) sequence of

DVQITQSPSFLSASVGDRTITCRASKSVSEYLA WYQEKPGKANKLLIYSGSTLHSGVPSRFS
GS GSGTEFTLTISSLQPEDFATYFCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:276) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

194. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
KFQGGVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID

NO:272) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:277) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

195. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSG
 SSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:278) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

196. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 EVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:279) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

197. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of
 EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSLKASDTAMYYCKRSLPGDFDYWGQGLTVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:280) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

198. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:281) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

199. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 EVWITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS
 GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:282) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

200. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
 GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK (SEQ ID NO:283) for
 binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

201. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of

EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWWRQMPGKGLEWIGYFSPGNADIRYSE
 KFQGQVTISADKSISTAYLQWSSSLKASDTAMYYCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
 NO:272) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 SSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTFGQGTKLEIK(SEQ ID NO:284) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

202. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment

competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
 EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS(SEQ ID
 NO:273) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKP GKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYFCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:276) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

203. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
 EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS(SEQ ID
 NO:273) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:277) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

204. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
 EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS(SEQ ID
 NO:273) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIY SASTLHSGVPSRFSG
 SSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:278) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

205. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
 EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGT LVTVSS(SEQ ID
 NO:273) and a light chain variable (VL) sequence of
 EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKP GQANRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISSLQSEDFAVYFCQQHNEY PFTFGQGTKLEIK(SEQ ID NO:279) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

206. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
NO:273) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:280) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

207. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
NO:273) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:281) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

208. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
NO:273) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

209. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYFSPGNGDIKYN
EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGTLVTVSS(SEQ ID
NO:273) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS

GS GSGTDFTLTINSLEAEDAATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:283) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

210. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKV SCKASGYTFTD HAIHWWRQAPGQGLEWIGYFSPGNGDIKYN
 EKFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:273) and a light chain variable (VL) sequence of
 EIVLTQSPDFQSVTPKEKVTITCRASKSIS SYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
 S GSGTDFTLTINSLEAEDAATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:284) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

211. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKV SCKASGYTFTD HAIHWWRQAPGQGLEWIGYISTGNGDIKYNQ
 KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of
 DVQITQSPSFLSASV GDRVTITCRASKSVSEY LAWYQEKP GKANKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYFCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:276) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

212. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKV SCKASGYTFTD HAIHWWRQAPGQGLEWIGYISTGNGDIKYNQ
 KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:274) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASV GDRVTITCRASKSVSEY LAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS
 GSGSGTEFTLTISSLQPEDFATYYCQQHNEY PFTFGQGTKLEIK (SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

213. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of
 QVQLVQSGSELKKPGASVKV SCKASGYTFTD HAIHWWRQAPGQGLEWIGYISTGNGDIKYNQ
 KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID

NO:274) and a light chain variable (VL) sequence of
 DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYSASTLHSGVPSRFSG
 SSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:278) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

214. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYISTGNGDIKYNQ
 KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
 NO:274) and a light chain variable (VL) sequence of
 EWITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPQANRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISLQSEDFAVYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:279) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

215. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYISTGNGDIKYNQ
 KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
 NO:274) and a light chain variable (VL) sequence of
 EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:280) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

216. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of
 QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWWRQAPGQGLEWIGYISTGNGDIKYNQ
 KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
 NO:274) and a light chain variable (VL) sequence of
 EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
 GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:281) for binding
 to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

217. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
 competes with an antibody or antigen binding fragment comprising a heavy chain variable
 (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ
KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:274) and a light chain variable (VL) sequence of
EVWITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

218. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ
KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:274) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:283) for
binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

219. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNGDIKYNQ
KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:274) and a light chain variable (VL) sequence of
EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG
SGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:284) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

220. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment
competes with an antibody or antigen binding fragment comprising a heavy chain variable
(VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of
DVQITQSPSFLSASVGDRTITCRASKSVSEYLAWYQEKPQKANKLLIYSGSTLHSGVPSRFS
GSGSGTEFTLTISSLQPEDFATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:276) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

221. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116
or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment

competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSVSEYLAWYQQKPGKAPKLLIYSGSTLHSGVPSRFS GSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:277) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

222. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of DIQLTQSPSFLSASVGDRTITCRASKSISEYLAWYQQKPGKAPKLLIYASSTLHSGVPSRFSG SSGSGTEFTLTISSLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:278) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

223. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWYQEKPGQANRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:279) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

224. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:275) and a light chain variable (VL) sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS GSGSGTEFTLTISSLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:280) for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

225. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of

EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWYQQKPGQAPRLLIYSGSTLHSGIPARFS
GSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:281) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

226. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of

EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQEKPDQSNKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:282) for binding
to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

227. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFS
GSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFTGQGTKLEIK (SEQ ID NO:283) for
binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

228. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 116 or embodiment 117, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of

QVQLVQSGSELKKPGASVKVSCASGYTFTDHAHWVRQAPGQGLEWIGYISTGNANITYAQ
GFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID
NO:275) and a light chain variable (VL) sequence of

EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSG

SGSGTDFTLTINSLEAEDAATYYCQQHNEYPTFGQGKLEIK(SEQ ID NO:284) for binding to the cMET glycopeptide.

229. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen binding fragment according to any one of embodiments 1 to 228, comprising:

- (a) a complementarity determining region (CDR) H1 comprising the amino acid sequence of a CDR-H1 of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:133, SEQ ID NO:139, SEQ ID NO:145, SEQ ID NO:205, or SEQ ID NO:253);
- (b) a CDR-H2 comprising the amino acid sequence of a CDR-H2 of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:134, SEQ ID NO:140, SEQ ID NO:146, SEQ ID NO:206, or SEQ ID NO:254);
- (c) a CDR-H3 comprising the amino acid sequence of a CDR-H3 of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, or SEQ ID NO:255);
- (d) a CDR-L1 comprising the amino acid sequence of a CDR-L1 of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, or SEQ ID NO:256);
- (e) a CDR-L2 comprising the amino acid sequence of a CDR-L2 of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:137, SEQ ID NO:143, SEQ ID NO:149, SEQ ID NO:209, or SEQ ID NO:257); and
- (f) a CDR-L3 comprising the amino acid sequence of a CDR-L3 of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, or SEQ ID NO:258).

230. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 229, wherein the amino acid designated X₁ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:134, SEQ ID NO:140, SEQ ID NO:146, SEQ ID NO:206, and/or SEQ ID NO:254) is G.

231. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 229, wherein the amino acid designated X₁ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:134, SEQ ID NO:140, SEQ ID NO:146, SEQ ID NO:206, and/or SEQ ID NO:254) is D.

232. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 231, wherein the amino acid designated X₂ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:134, SEQ ID NO:140, and/or SEQ ID NO:206) is I.

233. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 231, wherein the amino acid designated X₂ in a CDR sequence of any

one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:134, SEQ ID NO:140, and/or SEQ ID NO:206) is V.

234. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 233, wherein the amino acid designated X₃ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:140 and/or SEQ ID NO:206) is K.

235. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 233, wherein the amino acid designated X₃ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:140 and/or SEQ ID NO:206) is R.

236. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 235, wherein the amino acid designated X₄ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:140 and/or SEQ ID NO:206) is N.

237. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 235, wherein the amino acid designated X₄ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:140 and/or SEQ ID NO:206) is S.

238. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 237, wherein the amino acid designated X₅ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:140 and/or SEQ ID NO:206) is G.

239. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 237, wherein the amino acid designated X₅ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:140 and/or SEQ ID NO:206) is D.

240. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 239, wherein the amino acid designated X₆ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, and/or SEQ ID NO:255) is P.

241. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 239, wherein the amino acid designated X₆ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, and/or SEQ ID NO:255) is D.

242. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 241, wherein the amino acid designated X₇ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, and/or SEQ ID NO:255) is M.

243. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 241, wherein the amino acid designated X₇ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, and/or SEQ ID NO:255) is F.

244. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 243, wherein the amino acid designated X₈ in a CDR sequence of any

one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, and/or SEQ ID NO:255) is C.

245. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 243, wherein the amino acid designated X₈ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, and/or SEQ ID NO:255) is Y.

246. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 245, wherein the amino acid designated X₉ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is K.

247. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 245, wherein the amino acid designated X₉ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is R.

248. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 247, wherein the amino acid designated X₁₀ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is E.

249. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 247, wherein the amino acid designated X₁₀ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is K.

250. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 249, wherein the amino acid designated X₁₁ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is N.

251. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 249, wherein the amino acid designated X₁₁ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is S.

252. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 251, wherein the amino acid designated X₁₂ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is V.

253. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 251, wherein the amino acid designated X₁₂ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is I.

254. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 253, wherein the amino acid designated X_{13} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is G.

255. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 253, wherein the amino acid designated X_{13} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is S.

256. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 253, wherein the amino acid designated X_{13} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is N.

257. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 256, wherein the amino acid designated X_{14} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is I.

258. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 256, wherein the amino acid designated X_{14} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is E.

259. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 256, wherein the amino acid designated X_{14} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, and/or SEQ ID NO:256) is N.

260. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 259, wherein the amino acid designated X_{15} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is V.

261. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 259, wherein the amino acid designated X_{15} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is L.

262. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 261, wherein the amino acid designated X_{16} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is S.

263. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 261, wherein the amino acid designated X_{16} in a CDR sequence of any

one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is A.

264. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 261, wherein the amino acid designated X₁₆ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:142, SEQ ID NO:148, and/or SEQ ID NO:208) is V.

265. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 264, wherein the amino acid designated X₁₇ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:137, SEQ ID NO:143, SEQ ID NO:209, and/or SEQ ID NO:257) is G.

266. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 264, wherein the amino acid designated X₁₇ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:137, SEQ ID NO:143, SEQ ID NO:209, and/or SEQ ID NO:257) is S.

267. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 266, wherein the amino acid designated X₁₈ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:137, SEQ ID NO:143, SEQ ID NO:209, and/or SEQ ID NO:257) is P.

268. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 266, wherein the amino acid designated X₁₈ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:137, SEQ ID NO:143, SEQ ID NO:209, and/or SEQ ID NO:257) is G.

269. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 268, wherein the amino acid designated X₁₉ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is N.

270. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 268, wherein the amino acid designated X₁₉ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is T.

271. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 270, wherein the amino acid designated X₂₀ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is R.

272. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 270, wherein the amino acid designated X₂₀ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is L.

273. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 272, wherein the amino acid designated X_{21} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is Y.

274. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 272, wherein the amino acid designated X_{21} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is H.

275. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 272, wherein the amino acid designated X_{21} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is Q.

276. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 275, wherein the amino acid designated X_{22} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is T.

277. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 275, wherein the amino acid designated X_{22} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:143, SEQ ID NO:149, and/or SEQ ID NO:209) is S.

278. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 277, wherein the amino acid designated X_{23} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is G.

279. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 277, wherein the amino acid designated X_{23} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is Q.

280. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 279, wherein the amino acid designated X_{24} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is S.

281. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 279, wherein the amino acid designated X_{24} in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is H.

282. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 281, wherein the amino acid designated X_{25} in a CDR sequence of any

one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is Y.

283. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 281, wherein the amino acid designated X₂₅ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is N.

284. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 283, wherein the amino acid designated X₂₆ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is S.

285. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 283, wherein the amino acid designated X₂₆ in a CDR sequence of any one of Tables 1G, 1H, 1I, 2G, and 3G (e.g., SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, and/or SEQ ID NO:258) is E.

286. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 285, wherein CDR-H1 comprises the amino acid sequence of GYTFTDHA (SEQ ID NO:133).

287. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 285, wherein CDR-H1 comprises the amino acid sequence of DHAIH (SEQ ID NO:139).

288. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 285, wherein CDR-H1 comprises the amino acid sequence of GYTFTDH (SEQ ID NO:145).

289. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 285, wherein CDR-H1 comprises the amino acid sequence of GYTFTDHAIH (SEQ ID NO:205).

290. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 285, wherein CDR-H1 comprises the amino acid sequence of DH (SEQ ID NO:253).

291. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 290, wherein CDR-H2 comprises the amino acid sequence of FSPGNX₁DX₂ (SEQ ID NO:134).

292. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 290, wherein CDR-H2 comprises the amino acid sequence of YFSPGNX₁DX₂X₃YX₄EKFKX₅ (SEQ ID NO:140).

293. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 290, wherein CDR-H2 comprises the amino acid sequence of SPGNX₁D (SEQ ID NO:146).

294. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 290, wherein CDR-H2 comprises the amino acid sequence of YFSPGNX₁DX₂X₃YX₄EKF₅ (SEQ ID NO:206).

295. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 290, wherein CDR-H2 comprises the amino acid sequence of SPGNX₁D (SEQ ID NO:254).

296. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 295, wherein CDR-H3 comprises the amino acid sequence of KRSLPGX₆X₇DX₈ (SEQ ID NO:135).

297. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 295, wherein CDR-H3 comprises the amino acid sequence of SLPGX₆X₇DX₈ (SEQ ID NO:141).

298. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 295, wherein CDR-H3 comprises the amino acid sequence of SLPGX₆X₇DX₈ (SEQ ID NO:147).

299. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 295, wherein CDR-H3 comprises the amino acid sequence of KRSLPGX₆X₇DX₈ (SEQ ID NO:207).

300. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 295, wherein CDR-H3 comprises the amino acid sequence of SLPGX₆X₇DX₈ (SEQ ID NO:255).

301. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 300, wherein CDR-L1 comprises the amino acid sequence of X₁₀X₁₁X₁₂X₁₃X₁₄Y (SEQ ID NO:136).

302. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 300, wherein CDR-L1 comprises the amino acid sequence of X₉ASX₁₀X₁₁X₁₂X₁₃X₁₄YX₁₅X₁₆ (SEQ ID NO:142).

303. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 300, wherein CDR-L1 comprises the amino acid sequence of X₉ASX₁₀X₁₁X₁₂X₁₃X₁₄YX₁₅X₁₆ (SEQ ID NO:148).

304. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 300, wherein CDR-L1 comprises the amino acid sequence of X₉ASX₁₀X₁₁X₁₂X₁₃X₁₄YX₁₅X₁₆ (SEQ ID NO:208).

305. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 300, wherein CDR-L1 comprises the amino acid sequence of X₁₀X₁₁X₁₂X₁₃X₁₄Y (SEQ ID NO:256).

306. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 305, wherein CDR-L2 comprises the amino acid sequence of $X_{17}X_{18}S$ (SEQ ID NO:137).

307. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 305, wherein CDR-L2 comprises the amino acid sequence of $X_{17}X_{18}SX_{19}X_{20}X_{21}X_{22}$ (SEQ ID NO:143).

308. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 305, wherein CDR-L2 comprises the amino acid sequence of $X_{17}X_{18}SX_{19}X_{20}X_{21}X_{22}$ (SEQ ID NO:149).

309. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 305, wherein CDR-L2 comprises the amino acid sequence of $X_{17}X_{18}SX_{19}X_{20}X_{21}X_{22}$ (SEQ ID NO:209).

310. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 305, wherein CDR-L2 comprises the amino acid sequence of $X_{17}X_{18}S$ (SEQ ID NO:257).

311. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 310, wherein CDR-L3 comprises the amino acid sequence of $X_{23}QX_{24}X_{25}X_{26}YPFT$ (SEQ ID NO:138).

312. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 310, wherein CDR-L3 comprises the amino acid sequence of $X_{23}QX_{24}X_{25}X_{26}YPFT$ (SEQ ID NO:144).

313. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 310, wherein CDR-L3 comprises the amino acid sequence of $X_{23}QX_{24}X_{25}X_{26}YPFT$ (SEQ ID NO:150).

314. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 310, wherein CDR-L3 comprises the amino acid sequence of $X_{23}QX_{24}X_{25}X_{26}YPFT$ (SEQ ID NO:210).

315. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 229 to 310, wherein CDR-L3 comprises the amino acid sequence of $X_{23}QX_{24}X_{25}X_{26}YPFT$ (SEQ ID NO:258).

316. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 15C4 as defined by IMGT (e.g., SEQ ID NOS:3-5) and a VL comprising CDRs of 15C4 as defined by IMGT (e.g., SEQ ID NOS:6-8).

317. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 15C4 as defined by Kabat (e.g., SEQ ID NOS:9-11) and a VL comprising CDRs of 15C4 as defined by Kabat (e.g., SEQ ID NOS:12-14).

318. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 15C4 as defined by Chothia (*e.g.*, SEQ ID NOS:15-17) and a VL comprising CDRs of 15C4 as defined by Chothia (*e.g.*, SEQ ID NOS:18-20).

319. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 8H3 as defined by IMGT (*e.g.*, SEQ ID NOS:25-27) and a VL comprising CDRs of 8H3 as defined by IMGT (*e.g.*, SEQ ID NOS:28-30).

320. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 8H3 as defined by Kabat (*e.g.*, SEQ ID NOS:31-33) and a VL comprising CDRs of 8H3 as defined by Kabat (*e.g.*, SEQ ID NOS:34-36).

321. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 8H3 as defined by Chothia (*e.g.*, SEQ ID NOS:37-39) and a VL comprising CDRs of 8H3 as defined by Chothia (*e.g.*, SEQ ID NOS:40-42).

322. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 16E12 as defined by IMGT (*e.g.*, SEQ ID NOS:47-49) and a VL comprising CDRs of 16E12 as defined by IMGT (*e.g.*, SEQ ID NOS:50-52).

323. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 16E12 as defined by Kabat (*e.g.*, SEQ ID NOS:53-55) and a VL comprising CDRs of 16E12 as defined by Kabat (*e.g.*, SEQ ID NOS:56-58).

324. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 16E12 as defined by Chothia (*e.g.*, SEQ ID NOS:59-61) and a VL comprising CDRs of 16E12 as defined by Chothia (*e.g.*, SEQ ID NOS:62-64).

325. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GYTFTDHAIH (SEQ ID NO:169), YFSPGNGDIKYNEKFKG (SEQ ID NO:170), and KRSLPGPMDC (SEQ ID NO:171); and a VL comprising CDRs of KASENVGIYVS (SEQ ID NO:172), GPSNRYT (SEQ ID NO:173), and GQSYSYPFT (SEQ ID NO:174).

326. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228,

which comprises a VH comprising CDRs of GYTFTDHAIH (SEQ ID NO:175), YFSPGNGDIKYNEKFKD (SEQ ID NO:176), and KRSLPGDFDY (SEQ ID NO:177); and a VL comprising CDRs of RASKSVSEYLA (SEQ ID NO:178), SGSTLHS (SEQ ID NO:179), and QQHNEYPPFT (SEQ ID NO:180).

327. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GYTFTDHAIH (SEQ ID NO:181), YFSPGNDDVRYSEKFKG (SEQ ID NO:182), and KRSLPGDFDY (SEQ ID NO:183); and a VL comprising CDRs of RASKSINNYLV (SEQ ID NO:184), SGSTLQT (SEQ ID NO:185), and QQHNEYPPFT (SEQ ID NO:186).

328. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of DH (SEQ ID NO:217), SPGNGD (SEQ ID NO:218), and SLPGPMD (SEQ ID NO:219); and a VL comprising CDRs of ENVGIY (SEQ ID NO:220), GPS (SEQ ID NO:221), and GQSYSYPFT (SEQ ID NO:222).

329. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of DH (SEQ ID NO:223), SPGNGD (SEQ ID NO:224), and SLPGDFDY (SEQ ID NO:225); and a VL comprising CDRs of KSVSEY (SEQ ID NO:226), SGS (SEQ ID NO:227), and QQHNEYPPFT (SEQ ID NO:228).

330. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of DH (SEQ ID NO:229), SPGNDD (SEQ ID NO:230), and SLPGDFDY (SEQ ID NO:231); and a VL comprising CDRs of KSINNY (SEQ ID NO:232), SGS (SEQ ID NO:233), and QQHNEYPPFT (SEQ ID NO:234).

331. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, comprising:

- (a) a complementarity determining region (CDR) H1 comprising the amino acid sequence of a CDR-H1 of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211, or SEQ ID NO:259);
- (b) a CDR-H2 comprising the amino acid sequence of a CDR-H2 of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212, or SEQ ID NO:260);
- (c) a CDR-H3 comprising the amino acid sequence of a CDR-H3 of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, or SEQ ID NO:261);

(d) a CDR-L1 comprising the amino acid sequence of a CDR-L1 of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, or SEQ ID NO:262);

(e) a CDR-L2 comprising the amino acid sequence of a CDR-L2 of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, or SEQ ID NO:263); and

(f) a CDR-L3 comprising the amino acid sequence of a CDR-L3 of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, or SEQ ID NO:342).

332. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 331, wherein the amino acid designated X_{27} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:163, and/or SEQ ID NO:211) is I.

333. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 331, wherein the amino acid designated X_{27} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:163, and/or SEQ ID NO:211) is V.

334. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 331, wherein the amino acid designated X_{27} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:163, and/or SEQ ID NO:211) is L.

335. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 335, wherein the amino acid designated X_{28} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:163, and/or SEQ ID NO:211) is D.

336. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 335, wherein the amino acid designated X_{28} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:163, and/or SEQ ID NO:211) is A.

337. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 336, wherein the amino acid designated X_{29} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211 and/or SEQ ID NO:259) is absent.

338. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 336, wherein the amino acid designated X_{29} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211 and/or SEQ ID NO:259) is G.

339. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 338, wherein the amino acid designated X_{30} in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211 and/or SEQ ID NO:259) is S.

340. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 338, wherein the amino acid designated X₃₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211 and/or SEQ ID NO:259) is I.

341. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 340, wherein the amino acid designated X₃₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211 and/or SEQ ID NO:259) is Y.

342. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 340, wherein the amino acid designated X₃₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211 and/or SEQ ID NO:259) is Q.

343. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 342, wherein the amino acid designated X₃₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:157 and/or SEQ ID NO:211) is I.

344. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 342, wherein the amino acid designated X₃₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:157 and/or SEQ ID NO:211) is A.

345. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 344, wherein the amino acid designated X₃₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, and/or SEQ ID NO:212) is I.

346. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 344, wherein the amino acid designated X₃₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, and/or SEQ ID NO:212) is M.

347. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 346, wherein the amino acid designated X₃₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is Y.

348. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 346, wherein the amino acid designated X₃₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is D.

349. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 348, wherein the amino acid designated X₃₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is T.

350. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 348, wherein the amino acid designated X₃₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is N.

351. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 350, wherein the amino acid designated X₃₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is G.

352. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 350, wherein the amino acid designated X₃₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is R.

353. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 352, wherein the amino acid designated X₃₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is S.

354. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 352, wherein the amino acid designated X₃₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is V.

355. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 352, wherein the amino acid designated X₃₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is absent.

356. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 355, wherein the amino acid designated X₃₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is G.

357. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 355, wherein the amino acid designated X₃₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is S.

358. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 355, wherein the amino acid designated X₃₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is absent.

359. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 358, wherein the amino acid designated X₃₉ in a CDR sequence of any

one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is G.

360. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 358, wherein the amino acid designated X₃₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is A.

361. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 358, wherein the amino acid designated X₃₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is absent.

362. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 361, wherein the amino acid designated X₄₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is N.

363. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 361, wherein the amino acid designated X₄₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is T.

364. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 361, wherein the amino acid designated X₄₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212 and/or SEQ ID NO:260) is absent.

365. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 364, wherein the amino acid designated X₄₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, and/or SEQ ID NO:212) is T.

366. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 364, wherein the amino acid designated X₄₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, and/or SEQ ID NO:212) is D.

367. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 364, wherein the amino acid designated X₄₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:152, SEQ ID NO:158, and/or SEQ ID NO:212) is absent.

368. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 367, wherein the amino acid designated X₄₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:158 and/or SEQ ID NO:212) is Y.

369. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 367, wherein the amino acid designated X₄₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:158 and/or SEQ ID NO:212) is absent.

370. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 369, wherein the amino acid designated X₄₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:158 and/or SEQ ID NO:212) is T.

371. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 369, wherein the amino acid designated X₄₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:158 and/or SEQ ID NO:212) is N.

372. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 371, wherein the amino acid designated X₄₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:158 and/or SEQ ID NO:212) is K.

373. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 371, wherein the amino acid designated X₄₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:158 and/or SEQ ID NO:212) is R.

374. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 373, wherein the amino acid designated X₄₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is M.

375. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 373, wherein the amino acid designated X₄₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is G.

376. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 375, wherein the amino acid designated X₄₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is S.

377. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 375, wherein the amino acid designated X₄₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is E.

378. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 375, wherein the amino acid designated X₄₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is G.

379. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 378, wherein the amino acid designated X₄₇ in a CDR sequence of any

one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is A.

380. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 378, wherein the amino acid designated X₄₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is D.

381. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 378, wherein the amino acid designated X₄₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

382. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 381, wherein the amino acid designated X₄₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is Y.

383. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 381, wherein the amino acid designated X₄₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is R.

384. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 383, wherein the amino acid designated X₄₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is I.

385. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 383, wherein the amino acid designated X₄₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is V.

386. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 383, wherein the amino acid designated X₄₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

387. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 386, wherein the amino acid designated X₅₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is A.

388. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 386, wherein the amino acid designated X₅₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is G.

389. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 386, wherein the amino acid designated X₅₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

390. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 389, wherein the amino acid designated X₅₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is T.

391. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 389, wherein the amino acid designated X₅₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is V.

392. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 389, wherein the amino acid designated X₅₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

393. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 392, wherein the amino acid designated X₅₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is Y.

394. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 392, wherein the amino acid designated X₅₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

395. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 394, wherein the amino acid designated X₅₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is I.

396. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 394, wherein the amino acid designated X₅₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is T.

397. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 394, wherein the amino acid designated X₅₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

398. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 397, wherein the amino acid designated X₅₄ in a CDR sequence of any

one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is T.

399. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 397, wherein the amino acid designated X₅₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is I.

400. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 397, wherein the amino acid designated X₅₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is L.

401. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 400, wherein the amino acid designated X₅₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is G.

402. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 400, wherein the amino acid designated X₅₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

403. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 402, wherein the amino acid designated X₅₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is A.

404. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 402, wherein the amino acid designated X₅₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, and/or SEQ ID NO:261) is absent.

405. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 404, wherein the amino acid designated X₅₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:160, SEQ ID NO:166, and/or SEQ ID NO:214) is A.

406. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 404, wherein the amino acid designated X₅₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:160, SEQ ID NO:166, and/or SEQ ID NO:214) is S.

407. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 406, wherein the amino acid designated X₅₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is S.

408. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 406, wherein the amino acid designated X₅₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is T.

409. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 408, wherein the amino acid designated X₅₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is I.

410. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 408, wherein the amino acid designated X₅₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is V.

411. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 410, wherein the amino acid designated X₆₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is S.

412. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 410, wherein the amino acid designated X₆₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is Y.

413. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 412, wherein the amino acid designated X₆₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is N.

414. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 412, wherein the amino acid designated X₆₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is S.

415. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 414, wherein the amino acid designated X₆₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is W.

416. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 414, wherein the amino acid designated X₆₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is Y.

417. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 414, wherein the amino acid designated X₆₂ in a CDR sequence of any

one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is N.

418. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 417, wherein the amino acid designated X₆₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is N.

419. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 417, wherein the amino acid designated X₆₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is absent.

420. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 419, wherein the amino acid designated X₆₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is E.

421. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 419, wherein the amino acid designated X₆₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, and/or SEQ ID NO:262) is absent.

422. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 421, wherein the amino acid designated X₆₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:160, SEQ ID NO:166, and/or SEQ ID NO:214) is A.

423. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 421, wherein the amino acid designated X₆₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:160, SEQ ID NO:166, and/or SEQ ID NO:214) is S.

424. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 423, wherein the amino acid designated X₆₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, and/or SEQ ID NO:263) is S.

425. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 423, wherein the amino acid designated X₆₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, and/or SEQ ID NO:263) is A.

426. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 423, wherein the amino acid designated X₆₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, and/or SEQ ID NO:263) is D.

427. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 426, wherein the amino acid designated X₆₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, and/or SEQ ID NO:263) is A.

428. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 426, wherein the amino acid designated X₆₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, and/or SEQ ID NO:263) is T.

429. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 428, wherein the amino acid designated X₆₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:161, SEQ ID NO:167, and/or SEQ ID NO:215) is Y.

430. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 428, wherein the amino acid designated X₆₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:161, SEQ ID NO:167, and/or SEQ ID NO:215) is T.

431. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 430, wherein the amino acid designated X₆₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:161, SEQ ID NO:167, and/or SEQ ID NO:215) is E.

432. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 430, wherein the amino acid designated X₆₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:161, SEQ ID NO:167, and/or SEQ ID NO:215) is A.

433. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 432, wherein the amino acid designated X₇₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is C.

434. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 432, wherein the amino acid designated X₇₀ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is G.

435. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 434, wherein the amino acid designated X₇₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is T.

436. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 434, wherein the amino acid designated X₇₁ in a CDR sequence of any

one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is S.

437. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 434, wherein the amino acid designated X₇₁ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is I.

438. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 437, wherein the amino acid designated X₇₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is G.

439. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 437, wherein the amino acid designated X₇₂ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is Y.

440. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 438, wherein the amino acid designated X₇₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is S.

441. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 438, wherein the amino acid designated X₇₃ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is I.

442. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 441, wherein the amino acid designated X₇₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is S.

443. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 441, wherein the amino acid designated X₇₄ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is absent.

444. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 443, wherein the amino acid designated X₇₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is D.

445. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 443, wherein the amino acid designated X₇₅ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is Y.

446. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 445, wherein the amino acid designated X₇₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is S.

447. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 445, wherein the amino acid designated X₇₆ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is W.

448. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 447, wherein the amino acid designated X₇₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is G.

449. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 447, wherein the amino acid designated X₇₇ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is Y.

450. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 449, wherein the amino acid designated X₇₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is W.

451. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 449, wherein the amino acid designated X₇₈ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is A.

452. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 451, wherein the amino acid designated X₇₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is D.

453. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 451, wherein the amino acid designated X₇₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is T.

454. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 451, wherein the amino acid designated X₇₉ in a CDR sequence of any one of Tables 1J, 1K, 1L, 2H, and 3H (e.g., SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, and/or SEQ ID NO:342) is absent.

455. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 454, wherein CDR-H1 comprises the amino acid sequence of $GX_{27}X_{28}FSX_{29}X_{30}X_{31}W$ (SEQ ID NO:151).

456. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 454, wherein CDR-H1 comprises the amino acid sequence of $X_{29}X_{30}X_{31}WX_{32}C$ (SEQ ID NO:157).

457. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 454, wherein CDR-H1 comprises the amino acid sequence of $GX_{27}X_{28}FSX_{29}X_{30}X_{31}$ (SEQ ID NO:163).

458. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 454, wherein CDR-H1 comprises the amino acid sequence of $GX_{27}X_{28}FSX_{29}X_{30}X_{31}WX_{32}C$ (SEQ ID NO:211).

459. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 454, wherein CDR-H1 comprises the amino acid sequence of $X_{29}X_{30}X_{31}$ (SEQ ID NO:259).

460. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 459, wherein CDR-H2 comprises the amino acid sequence of $X_{33}X_{34}X_{35}X_{36}X_{37}X_{38}X_{39}X_{40}X_{41}$ (SEQ ID NO:152).

461. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 459, wherein CDR-H2 comprises the amino acid sequence of $CX_{33}X_{34}X_{35}X_{36}X_{37}X_{38}X_{39}X_{40}X_{41}X_{42}YAX_{43}WAX_{44}G$ (SEQ ID NO:158).

462. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 459, wherein CDR-H2 comprises the amino acid sequence of $X_{34}X_{35}X_{36}X_{37}X_{38}X_{39}X_{40}$ (SEQ ID NO:164).

463. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 459, wherein CDR-H2 comprises the amino acid sequence of $CX_{33}X_{34}X_{35}X_{36}X_{37}X_{38}X_{39}X_{40}X_{41}X_{42}YAX_{43}WAX_{44}G$ (SEQ ID NO:212).

464. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 459, wherein CDR-H2 comprises the amino acid sequence of $X_{34}X_{35}X_{36}X_{37}X_{38}X_{39}X_{40}$ (SEQ ID NO:260).

465. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 464, wherein CDR-H3 comprises the amino acid sequence of $ARX_{45}GYX_{46}X_{47}GX_{48}X_{49}GX_{50}X_{51}X_{52}X_{53}X_{54}VX_{55}X_{56}FNL$ (SEQ ID NO:153).

466. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 464, wherein CDR-H3 comprises the amino acid sequence of $X_{45}GYX_{46}X_{47}GX_{48}X_{49}GX_{50}X_{51}X_{52}X_{53}X_{54}VX_{55}X_{56}FNL$ (SEQ ID NO:159).

467. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 464, wherein CDR-H3 comprises the amino acid sequence of X₄₅GYX₄₆X₄₇GX₄₈X₄₉GX₅₀X₅₁X₅₂X₅₃X₅₄VX₅₅X₅₆FNL (SEQ ID NO:165).

468. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 464, wherein CDR-H3 comprises the amino acid sequence of ARX₄₅GYX₄₆X₄₇GX₄₈X₄₉GX₅₀X₅₁X₅₂X₅₃X₅₄VX₅₅X₅₆FNL (SEQ ID NO:213).

469. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 464, wherein CDR-H3 comprises the amino acid sequence of X₄₅GYX₄₆X₄₇GX₄₈X₄₉GX₅₀X₅₁X₅₂X₅₃X₅₄VX₅₅X₅₆FNL (SEQ ID NO:261).

470. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 469, wherein CDR-L1 comprises the amino acid sequence of QX₅₈X₅₉X₆₀X₆₁X₆₂X₆₃X₆₄ (SEQ ID NO:154).

471. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 469, wherein CDR-L1 comprises the amino acid sequence of QX₅₇SQX₅₈X₅₉X₆₀X₆₁X₆₂X₆₃X₆₄LX₆₅ (SEQ ID NO:160).

472. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 469, wherein CDR-L1 comprises the amino acid sequence of QX₅₇SQX₅₈X₅₉X₆₀X₆₁X₆₂X₆₃X₆₄LX₆₅ (SEQ ID NO:166).

473. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 469, wherein CDR-L1 comprises the amino acid sequence of QX₅₇SQX₅₈X₅₉X₆₀X₆₁X₆₂X₆₃X₆₄LX₆₅ (SEQ ID NO:214).

474. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 469, wherein CDR-L1 comprises the amino acid sequence of QX₅₈X₅₉X₆₀X₆₁X₆₂X₆₃X₆₄ (SEQ ID NO:262).

475. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 474, wherein CDR-L2 comprises the amino acid sequence of X₆₆X₆₇S (SEQ ID NO:155).

476. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 474, wherein CDR-L2 comprises the amino acid sequence of X₆₆X₆₇SX₆₈LX₆₉S (SEQ ID NO:161).

477. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 474, wherein CDR-L2 comprises the amino acid sequence of X₆₆X₆₇SX₆₈LX₆₉S (SEQ ID NO:167).

478. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 474, wherein CDR-L2 comprises the amino acid sequence of X₆₆X₆₇SX₆₈LX₆₉S (SEQ ID NO:209).

479. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 474, wherein CDR-L2 comprises the amino acid sequence of X₆₆X₆₇S (SEQ ID NO:263).

480. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 331 to 480, wherein CDR-L3 comprises the amino acid sequence of QX₇₀X₇₁YX₇₂X₇₃X₇₄GX₇₅X₇₆X₇₇SX₇₈X₇₉ (SEQ ID NO:156).

481. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 14E9 as defined by IMGT (*e.g.*, SEQ ID NOS:69-71) and a VL comprising CDRs of 14E9 as defined by IMGT (*e.g.*, SEQ ID NOS:72-74).

482. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 14E9 as defined by Kabat (*e.g.*, SEQ ID NOS:75-77) and a VL comprising CDRs of 14E9 as defined by Kabat (*e.g.*, SEQ ID NOS:78-80).

483. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 14E9 as defined by Chothia (*e.g.*, SEQ ID NOS:81-83) and a VL comprising CDRs of 14E9 as defined by Chothia (*e.g.*, SEQ ID NOS:84-86).

484. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 19H2 as defined by IMGT (*e.g.*, SEQ ID NOS:91-93) and a VL comprising CDRs of 19H2 as defined by IMGT (*e.g.*, SEQ ID NOS:94-96).

485. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 19H2 as defined by Kabat (*e.g.*, SEQ ID NOS:97-99) and a VL comprising CDRs of 19H2 as defined by Kabat (*e.g.*, SEQ ID NOS:100-102).

486. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 19H2 as defined by Chothia (*e.g.*, SEQ ID NOS:103-105) and a VL comprising CDRs of 19H2 as defined by Chothia (*e.g.*, SEQ ID NOS:106-108).

487. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 39A3 as defined by IMGT (*e.g.*, SEQ ID NOS:113-115) and a VL comprising CDRs of 39A3 as defined by IMGT (*e.g.*, SEQ ID NOS:116-118).

488. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228,

which comprises a VH comprising CDRs of 39A3 as defined by Kabat (e.g., SEQ ID NOS:119-121) and a VL comprising CDRs of 39A3 as defined by Kabat (e.g., SEQ ID NOS:122-124).

489. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of 39A3 as defined by Chothia (e.g., SEQ ID NOS:125-127) and a VL comprising CDRs of 39A3 as defined by Chothia (e.g., SEQ ID NOS:128-130).

490. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GIDFSSYWIC (SEQ ID NO:187), CIYTGSSGNTYYATWAKG (SEQ ID NO:188), and ARMGYSAGYIGATYITVGAFNL (SEQ ID NO:189); and a VL comprising CDRs of QASQISINWLA (SEQ ID NO:190), SASYLES (SEQ ID NO:191), and QCTYGSSGDSGSWD (SEQ ID NO:192).

491. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GVAFSGSQWIC (SEQ ID NO:193), CIYTGSSATDYANWARG (SEQ ID NO:194), and ARMGYEDGYVGGVYITVGAFNL (SEQ ID NO:195); and a VL comprising CDRs of QASQTISSYLA (SEQ ID NO:196), ATSYLES (SEQ ID NO:197), and QCSYGSGYSGSWT (SEQ ID NO:198).

492. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GLDFSGIYWAC (SEQ ID NO:199), CMDNRVTYATWAKG (SEQ ID NO:200), and ARGGYGGRGLVFNL (SEQ ID NO:201); and a VL comprising CDRs of QSSQSVYNNNELS (SEQ ID NO:202), DASTLAS (SEQ ID NO:203), and QGIYYIGDWYSA (SEQ ID NO:204).

493. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of SY (SEQ ID NO:235), YTGSSGN (SEQ ID NO:236), and MGYSAGYIGATYITVGAFNL (SEQ ID NO:237); and a VL comprising CDRs of QSISNW (SEQ ID NO:238), SAS (SEQ ID NO:239), and QCTYGSSGDSGSWD (SEQ ID NO:240).

494. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GSQ (SEQ ID NO:241), YTGSSAT (SEQ ID NO:242), and MGYEDGYVGGVYITVGAFNL (SEQ ID NO:243); and a VL comprising CDRs of QTISSY (SEQ ID NO:244), ATS (SEQ ID NO:245), and QCSYGSGYSGSWT (SEQ ID NO:246).

495. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising CDRs of GIY (SEQ ID NO:247), DNR (SEQ ID NO:248), and GYGGRGLVFNL (SEQ ID NO:249); and a VL comprising CDRs of QSVYNNNE (SEQ ID NO:250), DAS (SEQ ID NO:251), and QGIYYIGDWYSA (SEQ ID NO:252).

496. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 495, which is a chimeric or humanized antibody or antigen-binding fragment of a chimeric or humanized antibody.

497. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNGDIKYNE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVTVSS(SEQ ID
NO:1) and a VL comprising an amino acid sequence having at least 95% sequence identity to
NIVMTQSPKSMSSVGERVTLCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF
TGSGSATDFTLTISVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2).

498. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNGDIKYNE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVTVSS(SEQ ID
NO:1) and a VL comprising an amino acid sequence having at least 97% sequence identity to
NIVMTQSPKSMSSVGERVTLCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF
TGSGSATDFTLTISVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2).

499. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNGDIKYNE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVTVSS(SEQ ID
NO:1) and a VL comprising an amino acid sequence having at least 99% sequence identity to
NIVMTQSPKSMSSVGERVTLCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF
TGSGSATDFTLTISVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2).

500. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising the amino acid sequence of

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQKPEQGLEWIGYFSPGNGDIKYNE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVSS(SEQ ID
NO:1) and a VL comprising the amino acid sequence of
NIVMTQSPKSMMSVGERVTLCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF
TGSGSATDFTLTISVQAEDLADYHCGQSYSYPFTFGSGTKLEIK(SEQ ID NO:2).

501. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQRPEQGLEWIGYFSPGNGDIKYNE
KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:23) and a VL comprising an amino acid sequence having at least 95% sequence identity to
DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPCKTNKLLIYSGSTLHSGIPSRFSG
SGSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24).

502. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQRPEQGLEWIGYFSPGNGDIKYNE
KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:23) and a VL comprising an amino acid sequence having at least 97% sequence identity to
DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPCKTNKLLIYSGSTLHSGIPSRFSG
SGSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24).

503. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQRPEQGLEWIGYFSPGNGDIKYNE
KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:23) and a VL comprising an amino acid sequence having at least 99% sequence identity to
DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPCKTNKLLIYSGSTLHSGIPSRFSG
SGSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24).

504. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising the amino acid sequence of
QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQRPEQGLEWIGYFSPGNGDIKYNE
KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:23) and a VL comprising the amino acid sequence of

DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKP GKTNKLLIYSGSTLHSGIPSRFSG
SGSGTDFTLTITSLAPEDFAMYFCQQHNEYPFTFGAGTKLELK (SEQ ID NO:24).

505. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWKQKPEQGLEWIGYFSPGNDDVRYSE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:45) and a VL comprising an amino acid sequence having at least 95% sequence identity to
DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKP GKTIKPLIYSGSTLQTGTSPSRFSGS
GSGTDFSLTISLEPEDFAMYQCQQHNEYPFTFGAGTKLELK(SEQ ID NO:46).

506. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWKQKPEQGLEWIGYFSPGNDDVRYSE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:45) and a VL comprising an amino acid sequence having at least 97% sequence identity to
DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKP GKTIKPLIYSGSTLQTGTSPSRFSGS
GSGTDFSLTISLEPEDFAMYQCQQHNEYPFTFGAGTKLELK(SEQ ID NO:46).

507. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWKQKPEQGLEWIGYFSPGNDDVRYSE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:45) and a VL comprising an amino acid sequence having at least 99% sequence identity to
DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKP GKTIKPLIYSGSTLQTGTSPSRFSGS
GSGTDFSLTISLEPEDFAMYQCQQHNEYPFTFGAGTKLELK(SEQ ID NO:46).

508. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising the amino acid sequence of

QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWKQKPEQGLEWIGYFSPGNDDVRYSE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:45) and a VL comprising the amino acid sequence of
DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKP GKTIKPLIYSGSTLQTGTSPSRFSGS
GSGTDFSLTISLEPEDFAMYQCQQHNEYPFTFGAGTKLELK(SEQ ID NO:46).

509. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to

QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWGCITYTGSGGNTYY
ATWAKGRFTVSETSSTTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGT
LVTVSSGQPK (SEQ ID NO:67) and a VL comprising an amino acid sequence having at least 95% sequence identity to

DVVMTPASVGA AVGGTVTIKCQASQSI SNWLAWYQQKPGQPPKLLIYSASYLESVPSRF
SGSGSGTEFTLTISDLECADAAATYYCQCTYGSSGDSGSWDFGGGTEVVKGD PV (SEQ ID
NO:68).

510. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to

QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWGCITYTGSGGNTYY
ATWAKGRFTVSETSSTTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGT
LVTVSSGQPK (SEQ ID NO:67) and a VL comprising an amino acid sequence having at least 97% sequence identity to

DVVMTPASVGA AVGGTVTIKCQASQSI SNWLAWYQQKPGQPPKLLIYSASYLESVPSRF
SGSGSGTEFTLTISDLECADAAATYYCQCTYGSSGDSGSWDFGGGTEVVKGD PV (SEQ ID
NO:68).

511. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to

QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWGCITYTGSGGNTYY
ATWAKGRFTVSETSSTTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGT
LVTVSSGQPK (SEQ ID NO:67) and a VL comprising an amino acid sequence having at least 99% sequence identity to

DVVMTPASVGA AVGGTVTIKCQASQSI SNWLAWYQQKPGQPPKLLIYSASYLESVPSRF
SGSGSGTEFTLTISDLECADAAATYYCQCTYGSSGDSGSWDFGGGTEVVKGD PV (SEQ ID
NO:68).

512. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising the amino acid sequence of

QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWGCITYTGSGGNTYY
ATWAKGRFTVSETSSTTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGT

LVTVSSGQPK (SEQ ID NO:67) and a VL comprising the amino acid sequence of DVVMTQTPASVGA AVGGTVTIKCQASQSISNWLAWYQQKPGQPPKLLIYSASYLESGVPSRF SGSGSGTEFTLTISDLECADAAATYYCQCTYGSSGDSGSWDFGGGTEVVVKGDPV (SEQ ID NO:68).

513. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to

QQQLEESGGGLVKPGASLTLCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDY YANWARGRFTISKGSSPTVDLKMTSLTGADTGTYFCARMGYEDGYVGGVYTIVGAFNLWGQ GTLVTVSSGQPK (SEQ ID NO:89) and a VL comprising an amino acid sequence having at least 95% sequence identity to

DVVMTQTASPVSAAVGGTVTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV (SEQ ID NO:90).

514. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to

QQQLEESGGGLVKPGASLTLCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDY YANWARGRFTISKGSSPTVDLKMTSLTGADTGTYFCARMGYEDGYVGGVYTIVGAFNLWGQ GTLVTVSSGQPK (SEQ ID NO:89) and a VL comprising an amino acid sequence having at least 97% sequence identity to

DVVMTQTASPVSAAVGGTVTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV (SEQ ID NO:90).

515. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to

QQQLEESGGGLVKPGASLTLCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDY YANWARGRFTISKGSSPTVDLKMTSLTGADTGTYFCARMGYEDGYVGGVYTIVGAFNLWGQ GTLVTVSSGQPK (SEQ ID NO:89) and a VL comprising an amino acid sequence having at least 99% sequence identity to

DVVMTQTASPVSAAVGGTVTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV (SEQ ID NO:90).

516. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising the amino acid sequence of
 QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDY
 YANWARGRFTISKGSSPTVDLKMTSLTGADTGTYFCARMGYEDGYVGGVYIVGAFNLWGQ
 GTLVTVSSGQPK (SEQ ID NO:89) and a VL comprising the amino acid sequence of
 DVVMTQTASPVSAAVGGTVTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESQVPSRFK
 GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV (SEQ ID
 NO:90).

517. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to
 QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKGLEWIACMDNRVTYATWA
 KGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNLWGQGTTLVTVSSGQPK
 (SEQ ID NO:111) and a VL comprising an amino acid sequence having at least 95% sequence identity to
 DPVLTQTTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPS
 RFKGSGSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVVKGDPV (SEQ ID
 NO:112).

518. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to
 QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKGLEWIACMDNRVTYATWA
 KGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNLWGQGTTLVTVSSGQPK
 (SEQ ID NO:111) and a VL comprising an amino acid sequence having at least 97% sequence identity to
 DPVLTQTTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPS
 RFKGSGSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVVKGDPV (SEQ ID
 NO:112).

519. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to
 QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKGLEWIACMDNRVTYATWA
 KGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNLWGQGTTLVTVSSGQPK
 (SEQ ID NO:111) and a VL comprising an amino acid sequence having at least 99% sequence

identity to

DPVLTQTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPS
RFKGS GSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVKGDPV (SEQ ID
NO:112).

520. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising the amino acid sequence of

QSLEEYGGDLVKPGASLTCTASGLDFSGIYWACWVRQAPGKGLEWACMDNRVTYATWA
KGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNLWGQGLTVTVSSGQPK

(SEQ ID NO:111) and a VL comprising the amino acid sequence of

DPVLTQTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPS
RFKGS GSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVKGDPV (SEQ ID
NO:112).

521. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 228, which comprises a VH comprising an amino acid sequence having at least 95% sequence identity to any one of SEQ ID NOS:264-275 (the "VH reference sequence") and a VL comprising an amino acid sequence having at least 95% sequence identity to any one of SEQ ID NOS:276-284 (the "VL reference sequence").

522. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 521, which comprises a VH comprising an amino acid sequence having at least 97% sequence identity to the VH reference sequence and a VL comprising an amino acid sequence having at least 97% sequence identity to the VL reference sequence.

523. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 521, which comprises a VH comprising an amino acid sequence having at least 99% sequence identity to the VH reference sequence and a VL comprising an amino acid sequence having at least 99% sequence identity to the VL reference sequence.

524. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 521, which comprises a VH comprising an amino acid sequence having 100% sequence identity to the VH reference sequence and a VL comprising an amino acid sequence having 100% sequence identity to the VL reference sequence.

525. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:264.

526. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:265.

527. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:266.

528. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:267.

529. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:268.

530. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:269.

531. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:270.

532. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:271.

533. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:272.

534. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:273.

535. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:274.

536. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 524, wherein the VH reference sequence is SEQ ID NO:275.

537. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:276.

538. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:277.

539. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:278.

540. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:279.

541. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:280.

542. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:281.

543. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:282.

544. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:283.

545. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 521 to 536, wherein the VH reference sequence is SEQ ID NO:284.

546. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-cMET antibody or antigen-binding fragment according to any one of embodiments 1 to 545, that competes with a reference antibody or antigen binding fragment comprising:

(a) a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQKPEQGLEWIGYFSPGNNGDIKYNEKFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVTVSS(SEQ ID NO:1) and a light chain variable (VL) sequence of NIVMTQSPKSMMSVGERVTLSCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRFTGSGSATDFTLTISSVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2);

(b) a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQRPEQGLEWIGYFSPGNNGDIKYNEKFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID NO:23) and a light chain variable (VL) sequence of DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPGKTNKLLIYSGSTLHSGIPSRFSGSGSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24);

(c) a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWWKQKPEQGLEWIGYFSPGNDDVRYSEKFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID NO:45) and a light chain variable (VL) sequence of DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKPGKTIKPLIYSGSTLQTGTSPSRFSGSGSGTDFSLTISLEPEDFAMYYCQQHNEYPTFGAGTKLELK(SEQ ID NO:46);

(d) a heavy chain variable (VH) sequence of QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWVGCITYTGSSGNTYYATWAKGRFTVSETSTTVTLRMTSLTAADTATYFCARMGYASAGYIGATYITVGAFNLWGQGTLVTVSSGQPK (SEQ ID NO:67) and a light chain variable (VL) sequence of DVVMTQTPASVGA AVGGT VTIKQASQSISNWLAWYQQKPGQPPKLLIYSASYLESVPSRFSGSGSGTEFTLTISDLECAATYYCQCTYGSSGDSGSWDFGGGTEVVVKGDPV (SEQ ID NO:68);

(e) a heavy chain variable (VH) sequence of QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDYYANWARGRFTISKSSPTVDLKMSTLTGADTGTYFCARMGYEDGYVGGVYTVGAFNLWGQGT LVTVSSGQPK (SEQ ID NO:89) and a light chain variable (VL) sequence of DVVMTQTASPVSA AVGGT VTIKQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV (SEQ ID NO:90);

(f) a heavy chain variable (VH) sequence of QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKGLEWIACMDNRVYATWAKGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNWLWGQGTLVTVSSGQPK

(SEQ ID NO:111) and a light chain variable (VL) sequence of DPVLTQTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPSR FKGSGSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVVKGDPV (SEQ ID NO:112); or

(g) a humanized heavy chain variable (VH) sequence of 8H3 (e.g., any one of SEQ ID NOS:264-275) and a humanized light chain variable (VL) sequence of 8H3 (e.g., SEQ ID NOS:276-284),

for binding to a cMET peptide PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285) that has been glycosylated with GalNAc on the serine and threonine residues shown with bold and underlined text (“the cMET glycopeptide”), the anti-glyco-cMET antibody or antigen-binding fragment comprising:

(a) a VH sequence with first, second and third CDR means within the VH sequence; and

(b) a VL sequence with fourth, fifth and sixth CDR means within the VL sequence,

wherein the first, second, third, fourth, fifth, and sixth CDR means cooperate to effect binding of the anti-glyco-cMET antibody or antigen-binding fragment to the cMET glycopeptide.

547. An anti-glyco-cMET antibody or antigen-binding fragment that competes with a reference antibody or antigen binding fragment comprising:

(a) a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQKPEQGLEWIGYFSPGNQDIKYNEK FKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSTVSS(SEQ ID NO:1) and a light chain variable (VL) sequence of NIVMTQSPKSMMSVGERVTLSCASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRFT GSGSATDFTLTISSVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2);

(b) a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQRPEQGLEWIGYFSPGNQDIKYNEK FKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID NO:23) and a light chain variable (VL) sequence of DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPGKTNKLLIYSGSTLHSGIPSRFSGS GSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24);

(c) a heavy chain variable (VH) sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQKPEQGLEWIGYFSPGNDDVRYSE KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID NO:45) and a light chain variable (VL) sequence of DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKPGKTIKPLIYSGSTLQTGTPSRFSGS GSGTDFSLTISLEPEDFAMYYCQQHNEYPTFGAGTKLELK(SEQ ID NO:46);

(d) a heavy chain variable (VH) sequence of
 QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWVGCYITGSSGNTYYA
 TWAKGRFTVSETSSTTVTLRMTSLTAADTATYFCARMGY SAGYIGATYITVGAFNLWGQGTLV
 TVSSGQPK (SEQ ID NO:67) and a light chain variable (VL) sequence of
 DVWMTQTPASVGA AVGGT VTIKCQASQSISNWLAWYQQKPGQPPKLLIYSASYLESVPSRFS
 GSGSGTEFTLTISDLECA DAATYYCQCTY GSSGDSGSWDFGGGTEVWVKGDPV (SEQ ID
 NO:68);

(e) a heavy chain variable (VH) sequence of
 QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDYY
 ANWARGRFTISKSSPTVDLKMTSLTGADTGT YFCARMGYEDGYVGGVYTVGAFNLWGQGT
 LVTVSSGQPK (SEQ ID NO:89) and a light chain variable (VL) sequence of
 DVWMTQTASPVSA AVGGT VTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK
 GSGSGTQFTLTISGVQCDDAATYYCQCSYGSYSGSWTFGGGTEVWVKGDPV (SEQ ID
 NO:90);

(f) a heavy chain variable (VH) sequence of
 QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKLEWIACMDNRVYATWAK
 GRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGG RGLVFN LWGQGT LVTVSSGQPK
 (SEQ ID NO:111) and a light chain variable (VL) sequence of
 DPVLTQTPPSVSA AVGGT VTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPSR
 FKSGSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVWVKGDPV (SEQ ID
 NO:112); or

(g) a humanized heavy chain variable (VH) sequence of 8H3 (*e.g.*, any one
 of SEQ ID NOS:264-275) and a humanized light chain variable (VL) sequence of 8H3 (*e.g.*,
 SEQ ID NOS:276-284),

for binding to a cMET peptide PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285) that has been
 glycosylated with GalNAc on the serine and threonine residues shown with bold and
 underlined text (“the cMET glycopeptide”), the anti-glyco-cMET antibody or antigen-binding
 fragment comprising a means for binding the cMET glycopeptide.

548. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 547,
 wherein the means for binding the cMET glycopeptide comprises a heavy chain variable (VH)
 domain and a light chain variable (VL) domain

549. The anti-glyco-cMET antibody or antigen-binding fragment of any one of
 embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment
 competes with a reference antibody or antigen binding fragment comprising a VH sequence of
 QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNGDIKYNE
 KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSVTVSS (SEQ ID
 NO:1) and a VL sequence of

NIVMTQSPKSMSSVGERVTLSCASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRF
TGSGSATDFLTISSVQAEDLADYHCGQSYSPFTFGSGTKLEIK(SEQ ID NO:2).

550. The anti-glyco-cMET antibody or antigen-binding fragment of any one of
embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment
competes with a reference antibody or antigen binding fragment comprising a VH sequence of
QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQRPEQGLEWIGYFSPGNNGDIKYNE
KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:23) and a VL sequence of
QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQRPEQGLEWIGYFSPGNNGDIKYNE
KFKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:23).

551. The anti-glyco-cMET antibody or antigen-binding fragment of any one of
embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment
competes with a reference antibody or antigen binding fragment comprising a VH sequence of
QVQLQQSDAELVKPGASVKISCKASGYTFTDHAHWKQKPEQGLEWIGYFSPGNDDVRYSE
KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS(SEQ ID
NO:45) and a VL sequence of
DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKPGKTIKPLIYSGSTLQTGTSPSRFSGS
GSGTDFSLTISLEPEDFAMYQCQHNEYPTFGAGTKLELK(SEQ ID NO:46).

552. The anti-glyco-cMET antibody or antigen-binding fragment of any one of
embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment
competes with a reference antibody or antigen binding fragment comprising a VH sequence of
QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWIGCIYTGSGGNTYY
ATWAKGRFTVSETSSSTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGT
LTVSSGQPK (SEQ ID NO:67) and a VL sequence of
DVVMTQTPASVGAAVGGTVTIKCQASQISNWLAWYQQKPGQPPKLLIYSASYLESQVPSRF
SGSGSGTEFTLTISDLECAATYYCQCTYGSSGDSGSWDFGGGTEVVVKGDPV (SEQ ID
NO:68).

553. The anti-glyco-cMET antibody or antigen-binding fragment of any one of
embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment
competes with a reference antibody or antigen binding fragment comprising a VH sequence of
QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDY
YANWARGRFTISKSSPTVDLKMTSLTGADTGTYFCARMGYEDGYVGGVYITVGAFNLWGQ
GTLVTVSSGQPK (SEQ ID NO:89) and a VL sequence of
DVVMTQTASPVSAAVGGTVTIKCQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK
GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSYSGSWTFGGGTEVVVKGDPV (SEQ ID
NO:90).

554. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment competes with a reference antibody or antigen binding fragment comprising a VH sequence of QSL E E Y G G D L V K P G A S L T L T C T A S G L D F S G I Y W A C W V R Q A P G K G L E W I A C M D N R V T Y A T W A K G R F T S S K T S S T T V T L Q M T S L T A A D T A T Y F C A R G G Y G G R G L V F N L W G Q G T L V T V S S G Q P K (SEQ ID NO:111) and a VL sequence of D P V L T Q T P P S V S A A V G G T V T I K C Q S S Q S V Y N N N E L S W Y Q Q K P G Q P P K L L I Y D A S T L A S G V P S R F K G S G S G T Q F T L T I S G V Q C D D A A T Y Y C Q G I Y Y I G D W Y S A F G G G T E V V K G D P V (SEQ ID NO:112).

555. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 546 to 548, wherein the anti-glyco-cMET antibody or antigen-binding fragment competes with a reference antibody or antigen binding fragment comprising a humanized heavy chain variable (VH) sequence of 8H3 (e.g., any one of SEQ ID NOS:264-275) and a humanized light chain variable (VL) sequence of 8H3 (e.g., any one of SEQ ID NOS:276-284).

556. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 555, which preferentially binds to a glyco-cMET epitope that is overexpressed on cancer cells as compared to normal cells.

557. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 556, which specifically binds to a cMET peptide P T K S F I S G G **ST** I T G V G K N L N (SEQ ID NO:285) that has been glycosylated with STn on the serine and threonine residues shown with bold and underlined text.

558. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 556, which does not specifically bind to a cMET peptide P T K S F I S G G **ST** I T G V G K N L N (SEQ ID NO:285) that has been glycosylated with STn on the serine and threonine residues shown with bold and underlined text.

559. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 1 nM to 200 nM as measured by surface plasmon resonance or bio-layer interferometry.

560. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 1 nM to 150 nM as measured by surface plasmon resonance or bio-layer interferometry.

561. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 1 nM to 100 nM as measured by surface plasmon resonance or bio-layer interferometry.

562. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 1 nM to 50 nM as measured by surface plasmon resonance or bio-layer interferometry.

563. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 5 nM to 200 nM as measured by surface plasmon resonance or bio-layer interferometry.

564. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 5 nM to 100 nM as measured by surface plasmon resonance or bio-layer interferometry.

565. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 5 nM to 50 nM as measured by surface plasmon resonance or bio-layer interferometry.

566. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 5 nM to 25 nM as measured by surface plasmon resonance or bio-layer interferometry.

567. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 5 nM to 10 nM as measured by surface plasmon resonance or bio-layer interferometry.

568. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 10 nM to 200 nM as measured by surface plasmon resonance or bio-layer interferometry.

569. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 10 nM to 100 nM as measured by surface plasmon resonance or bio-layer interferometry.

570. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 10 nM to 150 nM as measured by surface plasmon resonance or bio-layer interferometry.

571. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 10 nM to 100 nM as measured by surface plasmon resonance or bio-layer interferometry.

572. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 10 nM to 50 nM as measured by surface plasmon resonance or bio-layer interferometry.

573. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 10 nM to 25 nM as measured by surface plasmon resonance or bio-layer interferometry.

574. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 50 nM to 200 nM as measured by surface plasmon resonance or bio-layer interferometry.

575. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 50 nM to 150 nM as measured by surface plasmon resonance or bio-layer interferometry.

576. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 50 nM to 100 nM as measured by surface plasmon resonance or bio-layer interferometry.

577. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 100 nM to 200 nM as measured by surface plasmon resonance or bio-layer interferometry.

578. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 558, which binds to the cMET glycopeptide with a binding affinity (KD) of 100 nM to 150 nM as measured by surface plasmon resonance or bio-layer interferometry.

579. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 578, in which the binding affinity to the cMET glycopeptide is as measured by surface plasmon resonance.

580. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 578, in which the binding affinity to the cMET glycopeptide is as measured by bio-layer interferometry.

581. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 580, which does not specifically bind to the unglycosylated cMET peptide PTKSFISGGSTITGVGKNLN (SEQ ID NO:286) (the "unglycosylated cMET peptide").

582. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 581, which has a binding affinity to the cMET glycopeptide which is at least 3 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the unglycosylated cMET peptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the unglycosylated cMET peptide (*e.g.*, about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

583. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 582, which has a binding affinity to the cMET glycopeptide which is at least 5 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the unglycosylated cMET peptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the unglycosylated cMET peptide (*e.g.*, about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

584. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 583, which has a binding affinity to the cMET glycopeptide which is at least 10 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the unglycosylated cMET peptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the unglycosylated cMET peptide (*e.g.*, about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

585. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 584, which has a binding affinity to the cMET glycopeptide which is at least 20 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the unglycosylated cMET peptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the unglycosylated cMET peptide (*e.g.*, about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

586. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 585, which has a binding affinity to the cMET glycopeptide which is at least 50 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the unglycosylated cMET peptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the unglycosylated cMET peptide (*e.g.*, about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

587. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 586, which has a binding affinity to the cMET glycopeptide which is at least 100 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the unglycosylated cMET peptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the unglycosylated cMET peptide (*e.g.*, about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

588. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 587, which does not specifically bind to the MUC1 tandem repeat (VTSAPDTRPAPGSTAPPAHG)₃ (SEQ ID NO:288) that has been glycosylated *in vitro* using purified recombinant human glycosyltransferases GalNAc-T1, GalNAc-T2, and GalNAc-T4 (“the first MUC1 glycopeptide”).

589. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 588, which has a binding affinity to the cMET glycopeptide which is at least 3 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the first MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the first MUC1 peptide (*e.g.*, about 1 μM , about 1.5 μM , or about 2 μM of either peptide).

590. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 589, which has a binding affinity to the cMET glycopeptide which is at least 5 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the first MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the first MUC1 peptide (*e.g.*, about 1 μM , about 1.5 μM , or about 2 μM of either peptide).

591. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 590, which has a binding affinity to the cMET glycopeptide which is at least 10 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the first MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the first MUC1 peptide (*e.g.*, about 1 μM , about 1.5 μM , or about 2 μM of either peptide).

592. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 591, which has a binding affinity to the cMET glycopeptide which is at least 20 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the first MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the first MUC1 peptide (*e.g.*, about 1 μM , about 1.5 μM , or about 2 μM of either peptide).

593. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 592, which has a binding affinity to the cMET glycopeptide which is at least 50 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the first MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the first MUC1 peptide (*e.g.*, about 1 μM , about 1.5 μM , or about 2 μM of either peptide).

594. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 593, which has a binding affinity to the cMET glycopeptide which is at least 100 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to

the first MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the first MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

595. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 594, which does not specifically bind to the MUC1 peptide TAPPAHGVTSAPDTRPAPGSTAPPAHGVT (SEQ ID NO:289) that has been glycosylated in vitro with GalNAc on the serine and threonine residues shown with bold and underlined text (the “second MUC1 glycopeptide”).

596. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 595, which has a binding affinity to the cMET glycopeptide which is at least 3 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the second MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the second MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

597. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 596, which has a binding affinity to the cMET glycopeptide which is at least 5 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the second MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the second MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

598. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 597, which has a binding affinity to the cMET glycopeptide which is at least 10 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the second MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the second MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

599. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 598, which has a binding affinity to the cMET glycopeptide which is at least 20 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the second MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the second MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

600. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 599, which has a binding affinity to the cMET glycopeptide which is at least 50 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the second MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the second MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

601. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 600, which has a binding affinity to the cMET glycopeptide which is at least 100 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the second MUC1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the second MUC1 peptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

602. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 601, which does not specifically bind to the CD44v6 peptide GYRQIPKEDSHSTTGTAAA (SEQ ID NO:345) that has been glycosylated in vitro with GalNAc on the threonine and serine residues shown with bold and underlined text (the "CD44v6 glycopeptide").

603. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 602, which has a binding affinity to the cMET glycopeptide which is at least 3 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the CD44v6 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the CD44v6 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

604. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 603, which has a binding affinity to the cMET glycopeptide which is at least 5 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the CD44v6 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the CD44v6 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

605. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 604, which has a binding affinity to the cMET glycopeptide which is at least 10 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the CD44v6 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the

presence of saturating amounts of either the cMET glycopeptide or the CD44v6 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

606. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 605, which has a binding affinity to the cMET glycopeptide which is at least 20 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the CD44v6 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the CD44v6 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

607. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 606, which has a binding affinity to the cMET glycopeptide which is at least 50 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the CD44v6 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the CD44v6 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

608. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 607, which has a binding affinity to the cMET glycopeptide which is at least 100 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the CD44v6 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the CD44v6 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

609. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 608, which does not specifically bind to the MUC4 peptide CTIPSTAMHTRSTAAPIILP (SEQ ID NO:291) that has been glycosylated in vitro with GalNAc on the serine and threonine residues shown with bold and underlined text (the "MUC4 glycopeptide").

610. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 609, which has a binding affinity to the cMET glycopeptide which is at least 3 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the MUC4 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the MUC4 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

611. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 610, which has a binding affinity to the cMET glycopeptide which is at least 5 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the

MUC4 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the MUC4 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

612. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 611, which has a binding affinity to the cMET glycopeptide which is at least 10 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the MUC4 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the MUC4 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

613. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 612, which has a binding affinity to the cMET glycopeptide which is at least 20 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the MUC4 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the MUC4 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

614. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 613, which has a binding affinity to the cMET glycopeptide which is at least 50 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the MUC4 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the MUC4 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

615. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 614, which has a binding affinity to the cMET glycopeptide which is at least 100 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the MUC4 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the MUC4 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

616. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 615, which does not specifically bind to the LAMP1 peptide CEQDRP**SPTT**APPAPPSPSP (SEQ ID NO:292) that has been glycosylated in vitro with GalNAc on the serine and threonine residues shown with bold and underlined text (the "LAMP1 glycopeptide").

617. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 616, which has a binding affinity to the cMET glycopeptide which is at least 3 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the LAMP1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the LAMP1 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

618. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 617, which has a binding affinity to the cMET glycopeptide which is at least 5 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the LAMP1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the LAMP1 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

619. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 618, which has a binding affinity to the cMET glycopeptide which is at least 10 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the LAMP1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the LAMP1 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

620. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 619, which has a binding affinity to the cMET glycopeptide which is at least 20 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the LAMP1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the LAMP1 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

621. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 620, which has a binding affinity to the cMET glycopeptide which is at least 50 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to the LAMP1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the LAMP1 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

622. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 621, which has a binding affinity to the cMET glycopeptide which is at least 100 times the binding affinity of the anti-glyco-cMET antibody or antigen-binding fragment to

the LAMP1 glycopeptide, optionally wherein the binding affinity is measured by surface plasmon resonance, and further optionally where in the surface plasmon resonance is measured in the presence of saturating amounts of either the cMET glycopeptide or the LAMP1 glycopeptide (e.g., about 1 μ M, about 1.5 μ M, or about 2 μ M of either peptide).

623. An anti-glyco-cMET antibody or antigen-binding fragment comprising a means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells.

624. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 623, wherein the means for binding the cMET epitope comprises a heavy chain variable (VH) domain and a light chain variable (VL) domain.

625. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 624, which is multivalent.

626. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 625, which is an antigen-binding fragment.

627. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 626, wherein the antigen-binding fragment is in the form of a single-chain variable fragment (scFv).

628. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 627, wherein the scFv comprises the heavy chain variable fragment N-terminal to the light chain variable fragment.

629. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 627, wherein the scFv comprises the heavy chain variable fragment C-terminal to the light chain variable fragment.

630. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 627 to 629, wherein the scFv heavy chain variable fragment and light chain variable fragment are covalently bound to a linker sequence, which is optionally 4-15 amino acids.

631. The anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 625, which is in the form of a multispecific antibody.

632. An anti-glyco-cMET antibody or antigen-binding fragment comprising a means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells.

633. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 632, wherein the means for binding the cMET epitope comprises a heavy chain variable (VH) domain and a light chain variable (VL) domain.

634. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 631 to 633, wherein the multispecific antibody is a bispecific antibody that binds to a second epitope that is different from the first epitope.

635. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 634, wherein the bispecific antibody is a bottle opener, mAb-Fv, mAb-scFv, central-scFv, one-armed central-scFv, or dual scFv format bispecific antibody.

636. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 635, wherein the bispecific antibody is a bottle opener format bispecific antibody.

637. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 635, wherein the bispecific antibody is a mAb-Fv format bispecific antibody.

638. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 635, wherein the bispecific antibody is a mAb-scFv format bispecific antibody.

639. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 635, wherein the bispecific antibody is a central-scFv format bispecific antibody.

640. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 635, wherein the bispecific antibody is a one-armed central-scFv format bispecific antibody.

641. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 635, wherein the bispecific antibody is a dual scFv format bispecific antibody.

642. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 634, wherein the bispecific antibody is a bispecific domain-exchanged antibody (*e.g.*, a CrossMab), a Fab-arm exchange antibody, a bispecific T-cell engager (BiTE), or a dual-affinity retargeting molecule (DART).

643. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 642, wherein the bispecific antibody is a bispecific domain-exchanged antibody (*e.g.*, a CrossMab).

644. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 643, wherein the bispecific antibody is a bispecific IgG comprising a Fab-arm having a domain crossover between heavy and light chains (*e.g.*, a CrossMabFAB).

645. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 643, wherein the bispecific antibody is a bispecific IgG comprising a Fab-arm having a domain crossover between variable heavy and variable light chains (*e.g.*, a CrossMabVH-VL).

646. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 643, wherein the bispecific antibody is a bispecific IgG comprising a Fab-arm having a domain crossover between constant heavy and constant light chains (*e.g.*, a CrossMabCH1-CL).

647. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 642, wherein the bispecific antibody is a Fab-arm exchange antibody.

648. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 642, wherein the bispecific antibody is a dual-affinity retargeting molecule (DART).

649. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 642, wherein the bispecific antibody is a bispecific T-cell engager (BiTE).

650. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 634 to 649, wherein the second epitope is a cMET epitope.

651. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 634 to 649, wherein the second epitope is a cMET epitope that is overexpressed on cancer cells as compared to normal cells.

652. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 634 to 649, wherein the second epitope is a T-cell epitope.

653. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 652, wherein the T-cell epitope comprises a CD3 epitope, a CD8 epitope, a CD16 epitope, a CD25 epitope, a CD28 epitope, or an NKG2D epitope.

654. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 653, wherein the T-cell epitope comprises a CD3 epitope, which is optionally an epitope present in human CD3.

655. The anti-glyco-cMET antibody or antigen-binding fragment of embodiment 654, wherein the CD3 epitope comprises a CD3 gamma epitope, a CD3 delta epitope, a CD3 epsilon epitope, or a CD3 zeta epitope.

656. The anti-glyco-cMET antibody or antigen-binding fragment of any one of embodiments 1 to 655 which is conjugated to a detectable moiety.

657. The anti-glyco-cMET antibody or antigen binding fragment of embodiment 656 in which the detectable moiety is an enzyme, a radioisotope, or a fluorescent label.

658. A bispecific antibody comprising (a) a means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells and (b) a means for binding a T-cell epitope, optionally wherein the bispecific antibody has the features described in any one of embodiments 634 to 657.

659. The bispecific antibody of embodiment 658, wherein the means for binding the cMET epitope comprises a heavy chain variable (VH) domain and a light chain variable (VL) domain.

660. The bispecific antibody of embodiment 658 or embodiment 659, wherein the means for binding the T-cell epitope comprises a heavy chain variable (VH) domain and a light chain variable (VL) domain.

661. The bispecific antibody of any one of embodiments 658 to 660, wherein the T-cell epitope comprises a CD3 epitope, a CD8 epitope, a CD16 epitope, a CD25 epitope, a CD28 epitope, or an NKG2D epitope.

662. The bispecific antibody of embodiment 661, wherein the T-cell epitope comprises a CD3 epitope, which is optionally an epitope present in human CD3.

663. The bispecific antibody of embodiment 662, wherein the CD3 epitope comprises a CD3 gamma epitope, a CD3 delta epitope, a CD3 epsilon epitope, or a CD3 zeta epitope.

664. A fusion protein comprising the amino acid sequence of the anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 657 or the bispecific antibody of any one of embodiments 658 to 663, operably linked to at least a second amino acid sequence.

665. The fusion protein of embodiment 664, wherein the second amino acid sequence is that of 4-1BB, CD2, CD3-zeta, or a fragment thereof.

666. The fusion protein of embodiment 664, wherein the second amino acid sequence is that of a fusion peptide.

667. The fusion protein of embodiment 666, wherein the fusion peptide is a CD28-CD3-zeta, a 4-1BB (CD137)-CD3-zeta fusion peptide, a CD2-CD3-zeta fusion peptide, a CD28-CD2-CD3-zeta fusion peptide, or a 4-1BB (CD137)-CD2-CD3-zeta fusion peptide.

668. The fusion protein of embodiment 664, wherein the second amino acid sequence is that of a modulator of T cell activation or a fragment thereof.

669. The fusion protein of embodiment 668, wherein the modulator of T cell activation is IL-15 or IL-15R α .

670. The fusion protein of embodiment 664, wherein the second amino acid sequence is that of a MIC protein domain.

671. The fusion protein of embodiment 670, wherein the MIC protein domain is an α 1- α 2 domain.

672. The fusion protein of embodiment 671, wherein the α 1- α 2 domain is a MICA, MICB, ULBP1, ULBP2, ULBP3, ULBP4, ULBP5, ULBP6, or OMCP α 1- α 2 domain.

673. The fusion protein of any one of embodiments 670 to 672, wherein the MIC protein domain is an engineered MIC protein domain.

674. The fusion protein of embodiment 664, wherein the second amino acid sequence is that of a neuraminidase (EC 3.2.1.18 or EC 3.2.1.129).

675. The fusion protein of embodiment 674, wherein the neuraminidase amino acid sequence is derived from *Micromonospora viridifaciens*.

676. The fusion protein of embodiment 674 or 675, wherein the neuraminidase comprises an amino acid sequence having at least 95% sequence identity to
GGSPVPPGGEPLYTEQDLAVNGREGFPNYRIPALVTPDGDLLASYDGRPTGIDAPGPNSILQ
RRSTDGGRTWGEQQVVSAGQTTAPIKGFSDPSYLVRETGTIFNFHVYSQRQGFAGSRPGT
DPADPNVLHANVATSTDGGLTWSHRTITADITPDGWRSRFAASGEGIQRLRYGPHAGRLIQQ
YTIINAAGAFQAVSVYSDDHGRTWRAGEAVGVGMENKTVELSDGRVLLNSRDSARSGYRK
VAVSTDGGHSYGPVTIDRDLPDPTNNASIIRAFDPAPAGSARAKVLLFSNAASQTSRSQGTIR
MSCDDGQTPVSKVFQPGSMSYSTLTALPDGTYGLLYEPGTGIRYANFNLAWLGG (SEQ ID
NO:318).

677. The fusion protein of any one of embodiments 674 to 676, wherein the neuraminidase comprises an amino acid sequence having at least 97% sequence identity to
GGSPVPPGGEPLYTEQDLAVNGREGFPNYRIPALVTPDGDLLASYDGRPTGIDAPGPNSILQ
RRSTDGGRTWGEQQVVSAGQTTAPIKGFSDPSYLVRETGTIFNFHVYSQRQGFAGSRPGT
DPADPNVLHANVATSTDGGLTWSHRTITADITPDGWRSRFAASGEGIQRLRYGPHAGRLIQQ
YTIINAAGAFQAVSVYSDDHGRTWRAGEAVGVGMENKTVELSDGRVLLNSRDSARSGYRK
VAVSTDGGHSYGPVTIDRDLPDPTNNASIIRAFDPAPAGSARAKVLLFSNAASQTSRSQGTIR

MSCDDGQTPVSKVFQPGSMSYSTLTALPDGTYGLLYEPGTGIRYANFNLAWLGG (SEQ ID NO:318).

678. The fusion protein of any one of embodiments 674 to 677, wherein the neuraminidase comprises an amino acid sequence having at least 98% sequence identity to GGSPVPPGGEPLYTEQDLAVNGREGFPNYRIPALTVTPDGDLLASYDGRPTGIDAPGPNSILQ RRSTDGGRTWGEQQVVSAGQTTAPIKGFSDPSYLVRETGTIFNFHVYSQRQGFAGSRPGT DPADPNVLHANVATSTDGGLTWSHRTITADITPDPGWRSRFAASGEGIQRLRYGPHAGRLIQQ YTIINAAGAFQAVSVYSDDHGRTWRAGEAVGVGMNDENKTVELSDGRVLLNSRDSARSGYRK VAVSTDGGHSYGPVTIDRDLDPPTNNASIIRAFDPAPAGSARAKVLLFSNAASQTSRSQGTIR MSCDDGQTPVSKVFQPGSMSYSTLTALPDGTYGLLYEPGTGIRYANFNLAWLGG (SEQ ID NO:318).

679. The fusion protein of any one of embodiments 674 to 678, wherein the neuraminidase comprises an amino acid sequence having at least 99% sequence identity to GGSPVPPGGEPLYTEQDLAVNGREGFPNYRIPALTVTPDGDLLASYDGRPTGIDAPGPNSILQ RRSTDGGRTWGEQQVVSAGQTTAPIKGFSDPSYLVRETGTIFNFHVYSQRQGFAGSRPGT DPADPNVLHANVATSTDGGLTWSHRTITADITPDPGWRSRFAASGEGIQRLRYGPHAGRLIQQ YTIINAAGAFQAVSVYSDDHGRTWRAGEAVGVGMNDENKTVELSDGRVLLNSRDSARSGYRK VAVSTDGGHSYGPVTIDRDLDPPTNNASIIRAFDPAPAGSARAKVLLFSNAASQTSRSQGTIR MSCDDGQTPVSKVFQPGSMSYSTLTALPDGTYGLLYEPGTGIRYANFNLAWLGG (SEQ ID NO:318).

680. The fusion protein of any one of embodiments 674 to 679, wherein the neuraminidase comprises the amino acid GGSPVPPGGEPLYTEQDLAVNGREGFPNYRIPALTVTPDGDLLASYDGRPTGIDAPGPNSILQ RRSTDGGRTWGEQQVVSAGQTTAPIKGFSDPSYLVRETGTIFNFHVYSQRQGFAGSRPGT DPADPNVLHANVATSTDGGLTWSHRTITADITPDPGWRSRFAASGEGIQRLRYGPHAGRLIQQ YTIINAAGAFQAVSVYSDDHGRTWRAGEAVGVGMNDENKTVELSDGRVLLNSRDSARSGYRK VAVSTDGGHSYGPVTIDRDLDPPTNNASIIRAFDPAPAGSARAKVLLFSNAASQTSRSQGTIR MSCDDGQTPVSKVFQPGSMSYSTLTALPDGTYGLLYEPGTGIRYANFNLAWLGG (SEQ ID NO:318).

681. The fusion protein of any one of embodiments 674 to 680, which comprises a signal sequence.

682. The fusion protein of embodiment 681, wherein the signal sequence is a granulysin signal sequence.

683. The fusion protein of embodiment 681, wherein the signal sequence is a granzymeK signal sequence.

684. The fusion protein of embodiment 681, wherein the signal sequence is an NPY signal sequence.

685. The fusion protein of embodiment 681, wherein the signal sequence is an IFN signal sequence.

686. The fusion protein of any one of embodiments 674 to 685, which comprises a self-cleaving peptide sequence.

687. The fusion protein of embodiment 686, wherein the self-cleaving peptide sequence is a 2A peptide.

688. The fusion protein of embodiment 687, wherein the 2A peptide is T2A.

689. A chimeric antigen receptor (CAR) comprising one or more antigen-binding fragments according to any one of embodiments 626 to 630.

690. The CAR of embodiment 689, which comprises one or more scFvs according to any one of embodiments 627 to 630.

691. The CAR of embodiment 690, which comprises one scFv according to any one of embodiments 627 to 630.

692. The CAR of embodiment 691, which comprises two scFvs according to any one of embodiments 627 to 630.

693. The CAR of embodiment 692, wherein the two scFvs have the same amino acid sequence.

694. The CAR of embodiment 692 or 693, wherein the two scFvs are covalently bound by a linker sequence, which is optionally 4-15 amino acids.

695. The CAR of any one of embodiments 689 to 694, comprising in amino- to carboxy-terminal order: (i) the one or more antigen-binding fragments, (ii) a transmembrane domain, and (iii) an intracellular signaling domain.

696. A chimeric antigen receptor (CAR) comprising in amino- to carboxy-terminal order: (i) one or more means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells, (ii) a transmembrane domain, and (iii) an intracellular signaling domain.

697. The CAR of embodiment 696, wherein the means for binding the cMET epitope comprises a heavy chain variable (VH) domain and a light chain variable (VL) domain.

698. The CAR of any one of embodiments 695 to 697, wherein the transmembrane domain comprises a CD28 transmembrane domain.

699. The CAR of embodiment 698, wherein the CD28 transmembrane domain comprises the amino acid sequence FWLVVVGGLACYSLLVTVAFIIFWV (SEQ ID NO:296).

700. The CAR of any one of embodiments 695 to 699, wherein the intracellular signaling domain comprises a co-stimulatory signaling region.

701. The CAR of embodiment 700, wherein the co-stimulatory signaling region comprises a signaling portion of, or the entire, cytoplasmic domain of CD27, CD28, 4-1BB, OX40, CD30, CD40, PD-1, ICOS, lymphocyte function-associated antigen-1 (LFA-1), CD2,

CD7, LIGHT, NKG2C, B7-H3, a ligand that specifically binds with CD83, DAP10, GITR, or a combination thereof.

702. The CAR of embodiment 701, wherein the CD27, CD28, 4-1BB, OX40, CD30, CD40, PD-1, ICOS, lymphocyte function-associated antigen-1 (LFA-1), CD2, CD7, LIGHT, NKG2C, B7-H3, a ligand that specifically binds with CD83, DAP10, or GITR a human CD27, CD28, 4-1BB, OX40, CD30, CD40, PD-1, ICOS, lymphocyte function-associated antigen-1 (LFA-1), CD2, CD7, LIGHT, NKG2C, B7-H3, a ligand that specifically binds with CD83, DAP10, or GITR.

703. The CAR of embodiment 701 or embodiment 702, wherein a signaling portion of, or the entire co-stimulatory signaling domain comprises the cytoplasmic domain of CD2.

704. The CAR of embodiment 703, wherein the cytoplasmic domain of CD2 comprises the amino acid sequence
TKRKKQRSRRNDEELETRAHRVATEERGRKPHQIPASTPQNPATSQHPPPPPGHRSQAPSH
RPPPPGHRVQHQPQKRPPAPSGTQVHQKGPPLPRPRVQPKPPHGAENSLSPSSN (SEQ
ID NO:303).

705. The CAR of any one of embodiments 701 to 704, wherein the co-stimulatory signaling domain comprises a signaling portion of, or the entire, cytoplasmic domain of CD28.

706. The CAR of embodiment 705, wherein the cytoplasmic domain of CD28 comprises the amino acid sequence
RSKRSRLLHSDYMNMPRRPGPTRKHYPYAPPRDFAAYRS (SEQ ID NO:302).

707. The CAR of any one of embodiments 694 to 706, wherein the intracellular signaling domain comprises a T cell signaling domain.

708. The CAR of embodiment 707, wherein the T cell signaling domain is C-terminal to the co-stimulatory signaling region.

709. The CAR of embodiment 707 or embodiment 708, wherein the T cell signaling domain comprises a CD3-zeta signaling domain.

710. The CAR of embodiment 709, wherein the CD3-zeta signaling domain comprises the amino acid sequence
RVKFERSADAPAYQQGQNQLYNELNLGRREEYDVLDKRRGRDPEMGGKPRRKNPQEGLYN
ELQKDKMAEAYSEIGMKGERRRGKGDGLYQGLSTATKDTYDALHMQALPPR (SEQ ID
NO:301).

711. The CAR of any one of embodiments 695 to 710, which further comprises a signal peptide N-terminal to the one or more antibody fragments, one or more scFvs, or one or more means for binding a cMET epitope.

712. The CAR of embodiment 710, wherein the signal peptide is a human CD8 signal peptide.

713. The CAR of embodiment 712, wherein the human CD8 signal peptide comprises the amino acid sequence MALPVTALLLPLALLLHAARP (SEQ ID NO:294).

714. The CAR of any one of embodiments 695 to 713, which further comprises a hinge between the one or more antigen-binding fragments and the transmembrane domain.

715. The CAR of embodiment 714, wherein the hinge comprises a human CD8a hinge.

716. The CAR of embodiment 715, wherein the human CD8a hinge comprises the amino acid sequence TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFAC (SEQ ID NO:297).

717. The CAR of embodiment 715, wherein the human CD8a hinge comprises the amino acid sequence TTTPAPRPPTPAPTIASPLSLRPEACRPAAGGAVHTRGLDFACD (SEQ ID NO:298).

718. The CAR of embodiment 715, wherein the human CD8a hinge comprises the amino acid sequence TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFACD (SEQ ID NO:349).

719. The CAR of embodiment 714, wherein the hinge comprises a human IgG4-short hinge comprising the amino acid sequence ESKYGPPCPSCP (SEQ ID NO:299).

720. The CAR of embodiment 714, wherein the hinge comprises a human IgG4-long hinge comprising the amino acid sequence
ESKYGPPCPPCPAPPVAGPSVFLFPPKPKDTLMISRTPEVTCVVDVVSQEDPEVQFNWYVDG
VEVHNAKTKPREEQFQSTYRVVSVLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQP
REPQVYTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTPPVLDSDGSFF
LYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLSLGLGKM (SEQ ID NO:300).

721. A chimeric antigen receptor (CAR), whose amino acid sequence comprises the amino acid sequence of hu8H3-CART of Table 18 (SEQ ID NO:348).

722. A chimeric antigen receptor (CAR), whose amino acid sequence comprises the amino acid sequence of 15C4-CART of Table 16 (SEQ ID NO:339).

723. A chimeric antigen receptor (CAR), whose amino acid sequence comprises the amino acid sequence of 16E12-CART of Table 16 (SEQ ID NO:340).

724. A chimeric antigen receptor (CAR), whose amino acid sequence comprises the amino acid sequence of 8H3-CART of Table 16 (SEQ ID NO:341).

725. An antibody-drug conjugate comprising the anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 657 or the fusion protein of any one of embodiments 658 to 688 conjugated to a cytotoxic agent.

726. The antibody-drug conjugate of embodiment 725, wherein the cytotoxic agent is an auristatin, a DNA minor groove binding agent, an alkylating agent, an enediyne, a lexitropsin, a duocarmycin, a taxane, a dolastatin, a maytansinoid, or a vinca alkaloid.

727. The antibody-drug conjugate of embodiment 726, wherein the anti-glyco-cMET antibody or antigen-binding fragment or bispecific antibody is conjugated to the cytotoxic agent via a linker.

728. The antibody-drug conjugate of embodiment 727, wherein the linker is cleavable under intracellular conditions.
729. The antibody-drug conjugate of embodiment 728, wherein the cleavable linker is cleavable by an intracellular protease.
730. The antibody-drug conjugate of embodiment 729, wherein the linker comprises a dipeptide.
731. The antibody-drug conjugate of embodiment 730, wherein the dipeptide is val-cit or phe-lys.
732. The antibody-drug conjugate of embodiment 728, wherein the cleavable linker is hydrolyzable at a pH of less than 5.5.
733. The antibody-drug conjugate of embodiment 732, wherein the hydrolyzable linker is a hydrazone linker.
734. The antibody-drug conjugate of embodiment 728, wherein the cleavable linker is a disulfide linker.
735. A chimeric T cell receptor (TCR) comprising
- (a) an antigen-binding fragment according to any one of embodiments 626 to 630
 - (b) a first polypeptide chain comprising a first TCR domain comprising a first TCR transmembrane domain from a first TCR subunit; and
 - (c) a second polypeptide chain comprising a second TCR domain comprising a second TCR transmembrane domain from a second TCR subunit.
736. The chimeric TCR of embodiment 735, which comprises one or more scFvs according to any one of embodiments 627 to 630.
737. The chimeric TCR of embodiment 735 or 563, which comprises one scFv according to any one of embodiments 627 to 630.
738. A chimeric T cell receptor (TCR) comprising:
- (a) a means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells;
 - (b) a first polypeptide chain comprising a first TCR domain comprising a first TCR transmembrane domain from a first TCR subunit; and
 - (c) a second polypeptide chain comprising a second TCR domain comprising a second TCR transmembrane domain from a second TCR subunit.
739. The chimeric TCR of embodiment 738, wherein the means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells comprises an scFv.

740. The chimeric TCR of embodiment 737 or 739, wherein the first polypeptide chain further comprises the scFv, and optionally further comprises a linker between the first TCR domain and the scFv.

741. The chimeric TCR of embodiment 737 or 739, wherein the second polypeptide chain further comprises the scFv, and optionally further comprises a linker between the second TCR domain and the scFv.

742. The chimeric TCR of embodiment 735 or 736, which comprises two scFvs according to any one of embodiments 627 to 630.

743. The chimeric TCR of embodiment 738, wherein the means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells comprises two scFvs.

744. The chimeric TCR of embodiment 742 or 743, wherein the two scFvs have the same amino acid sequence.

745. The chimeric TCR of embodiment 742 or 743, wherein the two scFvs have different amino acid sequences.

746. The chimeric TCR of any one of embodiments 742 to 745, wherein the two scFvs are covalently bound by a linker sequence, which is optionally 4-15 amino acids in length.

747. The chimeric TCR of any one of embodiments 742 to 746, wherein the first polypeptide chain further comprises the two scFvs, and optionally further comprises a linker between the first TCR domain and a first scFv of the two scFvs.

748. The chimeric TCR of any one of embodiments 742 to 746, wherein the second polypeptide chain further comprises the two scFvs, and optionally further comprises a linker between the second TCR domain and a first scFv of the two scFvs.

749. The chimeric TCR of any one of embodiments 742 to 746, wherein the first polypeptide chain comprises a first scFv of the two scFvs, and the second polypeptide chain comprises a second scFv of the two scFvs, and optionally wherein (i) the first polypeptide chain comprises a first linker between the first TCR domain and the first scFv, and (ii) the second polypeptide chain comprises a second linker between the second TCR domain and the second scFv.

750. The chimeric TCR of embodiment 735, wherein the antigen-binding fragment is an anti-glyco-cMET Fv fragment.

751. The chimeric TCR of embodiment 738, wherein the means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells is an anti-glyco-cMET Fv fragment.

752. The chimeric TCR of embodiment 750 or 751, wherein the Fv fragment comprises an anti-glyco-cMET variable heavy chain (VH) and an anti-glyco-cMET variable light

chain (VL), optionally wherein the VH and VL are a VH and a VL of an anti-glyco-cMET antibody or binding fragment according to any one of embodiments 1 to 657.

753. The chimeric TCR of embodiment 752, wherein the first polypeptide chain further comprises the anti-glyco-cMET VH and the second polypeptide chain further comprises the anti-glyco-cMET VL, optionally wherein (i) the first polypeptide chain further comprises a linker between the first TCR domain and the anti-glyco-cMET VH, and (ii) the second polypeptide chain further comprises a linker between the second TCR domain and the anti-glyco-cMET VL.

754. The chimeric TCR of embodiment 752, wherein the first polypeptide chain further comprises the anti-glyco-cMET VL and the second polypeptide chain further comprises the anti-glyco-cMET VH, optionally wherein (i) the first polypeptide chain further comprises a linker between the first TCR domain and the anti-glyco-cMET VL, and (ii) the second polypeptide chain further comprises a linker between the second TCR domain and the anti-glyco-cMET VH.

755. The chimeric TCR of any one of embodiments 735 and 750 to 754, wherein the first polypeptide chain further comprises a common heavy chain 1 (CH1) domain.

756. The chimeric TCR of any one of embodiments 735 and 750 to 755, wherein the second polypeptide chain further comprises a common light chain (CL) domain.

757. The chimeric TCR of embodiment 735, wherein the antigen-binding fragment is an anti-glyco-cMET Fab domain.

758. The chimeric TCR of embodiment 738, wherein the means for binding a cMET epitope that is overexpressed on cancer cells as compared to normal cells is an anti-glyco-cMET Fab domain.

759. The chimeric TCR of embodiment 757 or 758 which comprises one anti-glyco-cMET Fab domain.

760. The chimeric TCR of embodiment 757 or 758 which comprises two anti-glyco-cMET Fab domain.

761. The chimeric TCR of embodiment 760, wherein the two Fab domains have the same amino acid sequence.

762. The chimeric TCR of embodiment 760, wherein the two Fab domains have different amino acid sequences.

763. The chimeric TCR of any one of embodiments 757 to 762, wherein the Fab domain or each Fab domain comprises an anti-glyco-cMET variable heavy chain (VH) and an anti-glyco-cMET variable light chain (VL), optionally wherein the VH and VL are a VH and a VL of an anti-glyco-cMET antibody or binding fragment according to any one of embodiments 1 to 657.

764. The chimeric TCR of embodiment 763, wherein the first polypeptide chain comprises the anti-glyco-cMET VH and a CH1 domain or a CL domain, optionally wherein the first polypeptide chain comprises a linker between the first TCR domain and the CH1 domain or the CL domain.

765. The chimeric TCR of embodiment 764, wherein the second polypeptide chain comprises the anti-glyco-cMET VL and a CL domain or a CH1 domain, optionally wherein the second polypeptide chain comprises a linker between the second TCR domain and the CL domain or the CH1 domain.

766. The chimeric TCR of embodiment 764, comprising a third polypeptide chain comprising the anti-glyco-cMET VL and a CL domain or a CH1 domain, the third polypeptide chain being capable of associating with the anti-glyco-cMET VH and the CH1 domain or the CL domain of the first polypeptide chain.

767. The chimeric TCR of embodiment 763, wherein the second polypeptide chain comprises the anti-glyco-cMET VH and a CH1 domain or a CL domain, optionally wherein the second polypeptide chain comprises a linker between the second TCR domain and the CH1 domain or the CL domain.

768. The chimeric TCR of embodiment 767, wherein the first polypeptide chain comprises the anti-glyco-cMET VL and a CL or a CH1 domain, optionally wherein the first polypeptide chain comprises a linker between the second TCR domain and the CL domain or the CH1.

769. The chimeric TCR of embodiment 767, comprising a third polypeptide chain comprising the anti-glyco-cMET VL and a CL domain or a CH1 domain, the third polypeptide chain being capable of associating with the anti-glyco-cMET VH and the CH1 domain or the CL domain of the second polypeptide chain.

770. The chimeric TCR of embodiment 763, wherein the first polypeptide chain comprises a first anti-glyco-cMET VH and a first chain CH1 domain or a first chain CL domain and the second polypeptide chain comprises a second anti-glyco-cMET VH and a second chain CH1 domain or a second chain CL domain, optionally wherein the first polypeptide chain comprises a linker between the first TCR domain and the first chain CH1 domain or the first chain CL domain, and optionally wherein the second polypeptide chain comprises a linker between the second TCR domain and the second chain CH1 domain or the second chain CL domain.

771. The chimeric TCR of embodiment 770, comprising:

- (a) a third polypeptide chain comprising a first anti-glyco-cMET VL and a third chain CL domain or a third chain CH1 domain, capable of associating with the first anti-glyco-cMET VH and the first chain CH1 domain or the first chain CL domain of the first polypeptide; and
- (b) a fourth polypeptide chain comprising a second anti-glyco-cMET VL and a fourth chain CL domain or a fourth chain CH1 domain, capable of associating with the second anti-glyco-cMET VH and the second chain CH1 domain or the second chain CL domain of the second polypeptide.

772. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQKPEQGLEWIGYFSPGNNGDIKYNEK FKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGPMDCWGQGTSVTVSS (SEQ ID NO:1).

773. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQRPEQGLEWIGYFSPGNNGDIKYNEK FKDKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID NO:23).

774. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLQQSDAELVKPGASVKISCKASGYTFTDHAIHWWKQKPEQGLEWIGYFSPGNDDVRYSE KFKGKATLTADKSSSTAYMQLNSLTSEDSAVYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID NO:45).

775. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QEQLVESGGGLVEPGASLTLTCKASGIDFSSYWICWVRQAPGKGLEWIGCIYTGSGGNTYY ATWAKGRFTVSETSTTVTLRMTSLTAADTATYFCARMGYSAGYIGATYITVGAFNLWGQGT LVTVSSGQPK (SEQ ID NO:67).

776. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QQQLEESGGGLVKPGASLTLTCAASGVAFSGSQWICWVRQAPGKGLEWIGCIYTGSSATDY YANWARGRFTISKSSPTVDLKMSTLTGADTGTYFCARMGYEDGYVGGVYITVGAFNLWGQ GTLVTVSSGQPK (SEQ ID NO:89).

777. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QSLEEYGGDLVKPGASLTLTCTASGLDFSGIYWACWVRQAPGKGLEWVACMDNRVYATWA KGRFTSSKTSSTTVTLQMTSLTAADTATYFCARGGYGGRGLVFNLVWGQGTTLVTVSSGQPK (SEQ ID NO:111).

778. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an

anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLVQSGAEVKKPGASVKVSKASGYTFTDHAIHWRQAPGQRLEWIGYFSPGNQDIKYNE KFKDRATLTADKSASTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:264).

779. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLVQSGAEVKKPGASVKVSKASGYTFTDHAIHWRQAPGQRLEWIGYFSPGNQDIKYSQ KFKGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:265).

780. The chimeric TCR of any one of embodiments 735 to 771, wherein the anti-glyco-cMET variable heavy chain comprises an amino acid of when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLVQSGAEVKKPGASVKVSKASGYTFTDHAIHWRQAPGQRLEWIGYFSPGNADTKYS QKFQGRVTITADKSASTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:266).

781. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAIHWRQAPGQGLEWIGYFSPGNQDIKYNE KFKDRATLTADKSTSTAYMELSSLRSEDVAVYFCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:267).

782. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFTDHAIHWRQAPGQGLEWIGYFSPGNQDIKYNQ KFKGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:268).

783. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of QVQLVQSGAEVKKPGSSVKVSKASGYTFSDHAIHWRQAPGQGLEWIGYFSPGNADINYAQ KFQGRVTITADKSTSTAYMELSSLRSEDVAVYYCKRSLPGDFDYWGQGLTVTVSS (SEQ ID NO:269).

784. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of

EVQLVQSGAEVKKPGESLKISCKASGYTFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNEK
FKDQATLSADKSISTAYLQWSSLKASDTAMYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:270).

785. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of
EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNNGDIKYNE
KFKGQVTISADKSISTAYLQWSSLKASDTAMYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:271).

786. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of
EVQLVQSGAEVKKPGESLKISCKGSGYSFTDHAIHWVRQMPGKGLEWIGYFSPGNADIRYSEK
FQGQVTISADKSISTAYLQWSSLKASDTAMYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:272).

787. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of
QVQLVQSGSELKKPGASVKVSKASGYTFTDHAIHWVRQAPGQGLEWIGYFSPGNNGDIKYNE
KFKDRAVLSADKSVSTAYLQISSLKAEDTAVYFCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:273).

788. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of
QVQLVQSGSELKKPGASVKVSKASGYTFTDHAIHWVRQAPGQGLEWIGYISTGNNGDIKYNQ
KFTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:274).

789. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising the amino acid sequence of
QVQLVQSGSELKKPGASVKVSKASGYTFTDHAIHWVRQAPGQGLEWIGYISTGNANITYAQQ
FTGRAVLSLDKSVSTAYLQISSLKAEDTAVYYCKRSLPGDFDYWGQGTTLTVSS (SEQ ID
NO:275).

790. The chimeric TCR of any one of embodiments 735 to 771, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable heavy chain comprising:

- (a) a complementarity determining region (CDR) H1 comprising the amino acid sequence of GYTFTDHA (SEQ ID NO:133), DHAIH (SEQ ID

NO:139), GYTFTDH (SEQ ID NO:145), GYTFTDHAIH (SEQ ID NO:205), or DH (SEQ ID NO:253);

- (b) a CDR-H2 comprising the amino acid sequence of FSPGNX₁DX₂ (SEQ ID NO:134), YFSPGNX₁DX₂X₃YX₄EKFKX₅ (SEQ ID NO:140), SPGNX₁D (SEQ ID NO:146), or YFSPGNX₁DX₂X₃YX₄EKFKX₅ (SEQ ID NO:206); and
- (c) a CDR-H3 comprising the amino acid sequence of KRSLPGX₆X₇DX₈ (SEQ ID NO:135), or SLPGX₆X₇DX₈ (SEQ ID NO:141).

791. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of NIVMTQSPKSMMSVGERVTLSCKASENVGIYVSWYQQKPEQSPKLLIYGPSNRYTGVPDRFT GSGSATDFTLTISSVQAEDLADYHCGQSYSPFTFGSGTKLEIK (SEQ ID NO:2).

792. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DVQITQSPSYLAASPGETITINCRASKSVSEYLAWYQEKPGKTNKLLIYSGSTLHSGIPSRFSGS GSGTDFTLTITSLAPEDFAMYFCQQHNEYPTFGAGTKLELK (SEQ ID NO:24).

793. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DVQISQSPSYLAASPGETITINCRASKSINNYLVWYQEKPGKTIKPLIYSGSTLQTGTSPSRFSGS GSGTDFSLTISSELEPEDFAMYYCQQHNEYPTFGAGTKLELK (SEQ ID NO:46).

794. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DVVMTQTPASVGA AVGGTVTIKQASQSISNWLAWYQQKPGQPPKLLIYSASYLESVPSRFSGSGSGTEFTLTISDLECADAAATYYCQCTYGSSGDSGSWDFGGGTEVVVKGDPV (SEQ ID NO:68).

795. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DVVMTQTASPVSA AVGGTVTIKQASQTISSYLAWYQQKPGQPPKLLIYATSYLESGVPSRFK GSGSGTQFTLTISGVQCDDAATYYCQCSYSGSGYSWTFGGGTEVVVKGDPV (SEQ ID NO:90).

796. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of

DPVLTQTPPSVSAAVGGTVTIKCQSSQSVYNNNELSWYQQKPGQPPKLLIYDASTLASGVPSR
FKGSGSGTQFTLTISGVQCDDAATYYCQGIYYIGDWYSAFGGGTEVVVKGDPV (SEQ ID
NO:112).

797. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DVQITQSPSFLSASVGDRVTITCRASKSVSEYLAWEKPKGKANKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:276).

798. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DIQLTQSPSFLSASVGDRVTITCRASKSVSEYLAWEKPKGKAPKLLIYSGSTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:277).

799. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of DIQLTQSPSFLSASVGDRVTITCRASKSISEYLAWEKPKGKAPKLLIYASSTLHSGVPSRFSGSGSGTEFTLTISLQPEDFATYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:278).

800. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of EVVITQSPATLSVSPGERATLSCRASKSVSEYLAWEKPGQANRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYFCQQHNEYPTFGQGTKLEIK(SEQ ID NO:279).

801. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of EIVMTQSPATLSVSPGERATLSCRASKSVSEYLAWEKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:280).

802. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of EIVMTQSPATLSVSPGERATLSCRASQSVSEYLAWEKPGQAPRLLIYSGSTLHSGIPARFSGSGSGTEFTLTISLQSEDFAVYYCQQHNEYPTFGQGTKLEIK(SEQ ID NO:281).

803. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of EVVITQSPDFQSVTPKEKVTITCRASKSVSEYLAWEKPDQSNKLLIYSGSTLHSGVPSRFSGSGSGTDFTLTINSLEAEDAATYFCQQHNEYPTFGQGTKLEIK (SEQ ID NO:282).

804. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSVSEYLAWYQQKPDQSPKLLIYSGSTLHSGVPSRFSGSGSGTDFLTINSLEAEDAATYYCQQHNEYPTFGQGKLEIK (SEQ ID NO:283).

805. The chimeric TCR of any one of embodiments 735 to 790 when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising the amino acid sequence of EIVLTQSPDFQSVTPKEKVTITCRASKSISSYLAWYQQKPDQSPKLLIYSGSTLFSGVPSRFSGSGSGTDFLTINSLEAEDAATYYCQQHNEYPTFGQGKLEIK (SEQ ID NO:284).

806. The chimeric TCR of any one of embodiments 735 to 790, when depending directly or indirectly from embodiment 708, wherein the antigen-binding fragment comprises an anti-glyco-cMET variable light chain comprising:

- (a) a CDR-L1 comprising the amino acid sequence of $X_{10}X_{11}X_{12}X_{13}X_{14}Y$ (SEQ ID NO:136), or $X_9ASX_{10}X_{11}X_{12}X_{13}X_{14}YX_{15}X_{16}$ (SEQ ID NO:142),
- (b) a CDR-L2 comprising the amino acid sequence of $X_{17}X_{18}S$ (SEQ ID NO:137), or $X_{17}X_{18}SX_{19}X_{20}X_{21}X_{22}$ (SEQ ID NO:143); and
- (c) a CDR-L3 comprising the amino acid sequence of $X_{23}QX_{24}X_{25}X_{26}YPFT$ (SEQ ID NO:138).

807. The chimeric TCR of any one of embodiments 740, 741, 747 to 749, 753, and 754, when comprising a first and/or a second linker, the first and/or second linkers comprise, individually, a constant domain or fragment thereof from an immunoglobulin or from a T cell receptor subunit.

808. The chimeric TCR of embodiment 807, wherein the first and/or second linkers comprise, individually, a CH1, CH2, CH3, CH4, or CL antibody domain, or a fragment of any one thereof.

809. The chimeric TCR of embodiment 807, wherein the first and/or second linkers comprise, individually, a $C\alpha$, $C\beta$, $C\gamma$, or $C\delta$ TCR domain, or a fragment of any one thereof.

810. The chimeric TCR of embodiment 809, wherein the first polypeptide chain comprises a $C\alpha$ TCR domain or a fragment thereof, and the second polypeptide chain comprises a $C\beta$ TCR domain or a fragment thereon.

811. The chimeric TCR of embodiment 809, wherein the first polypeptide chain comprises a $C\beta$ TCR domain or a fragment thereof, and the second polypeptide chain comprises a $C\alpha$ TCR domain or a fragment thereon.

812. The chimeric TCR of embodiment 809, wherein the first polypeptide chain comprises a $C\gamma$ TCR domain or a fragment thereof, and the second polypeptide chain comprises a $C\delta$ TCR domain or a fragment thereon.

813. The chimeric TCR of embodiment 809, wherein the first polypeptide chain comprises a C δ TCR domain or a fragment thereof, and the second polypeptide chain comprises a C γ TCR domain or a fragment thereon.

814. The chimeric TCR of any one of embodiments 809 to 813, wherein the first TCR constant region domain and the second TCR constant region domain each comprise at least one mutation relative to the wildtype TCR constant region domain.

815. The chimeric TCR of embodiment 814, wherein the C α TCR domain comprises a substitution at an amino acid corresponding to amino acid position 48 of wildtype human C α TCR and the C β TCR domain comprises a substitution at an amino acid corresponding amino acid position 57 of wildtype human C β TCR.

816. The chimeric TCR of embodiment 814 or 815 wherein the C α TCR domain comprises a substitution at an amino acid corresponding to amino acid position 85 of wildtype human C α TCR and the C β TCR domain comprises a substitution at an amino acid corresponding to amino acid position 88 of wildtype human C β TCR.

817. The chimeric TCR of any one of embodiments 735 to 816, wherein the first TCR domain further comprises a first connecting peptide of a TCR subunit, or a fragment thereof, N-terminal to the first TCR transmembrane domain.

818. The chimeric TCR of any one of embodiments 735 to 817, wherein the second TCR domain further comprises a second connecting peptide of a TCR subunit, or a fragment thereof, N-terminal to the second TCR transmembrane domain.

819. The chimeric TCR of embodiment 818, comprising a disulfide bond between a residue in the first connecting peptide and a residue in the second connecting peptide.

820. The chimeric TCR of any one of embodiments 735 to 819, wherein the first TCR domain further comprises a first TCR intracellular domain comprising a TCR intracellular sequence C-terminal to the first transmembrane domain.

821. The chimeric TCR of any one of embodiments 735 to 820, wherein the second TCR domain further comprises a second TCR intracellular domain comprising a TCR intracellular sequence C-terminal to the second transmembrane domain.

822. The chimeric TCR of any one of embodiments 735 to 821, wherein the first polypeptide chain further comprises a first accessory intracellular domain comprising a co-stimulatory intracellular signaling sequence C-terminal to the first transmembrane domain.

823. The chimeric TCR of any one of embodiments 735 to 822, wherein the second polypeptide chain further comprises a second accessory intracellular domain comprising a co-stimulatory intracellular signaling sequence C-terminal to the second transmembrane domain.

824. The chimeric TCR of any one of embodiments 735 to 823, further comprising a cleavable peptide linker, configured to temporarily associate the first polypeptide chain with the second polypeptide chain during and/or shortly after protein translation.

825. The chimeric TCR of embodiment 824, wherein the cleavable peptide linker is a protease cleavable peptide linker.

826. The chimeric TCR of embodiment 824 or 825, wherein the peptide linker comprises the sequence ATNFSLLKQAGDVEENPGP (SEQ ID NO:317).

827. The chimeric TCR of any one of embodiments 735 to 826, wherein the first TCR domain is a TCR α chain or a fragment thereof and the second TCR domain is a TCR β chain or a fragment thereof.

828. The chimeric TCR of any one of embodiments 735 to 826, wherein the first TCR domain is a TCR β chain or a fragment thereof and the second TCR domain is a TCR α chain or a fragment thereof.

829. The chimeric TCR of any one of embodiments 735 to 826, wherein the first TCR domain is a TCR δ chain or a fragment thereof and the second TCR domain is a TCR γ chain or a fragment thereof.

830. The chimeric TCR of any one of embodiments 735 to 826, wherein the first TCR domain is a TCR γ chain or a fragment thereof and the second TCR domain is a TCR δ chain or a fragment thereof.

831. The chimeric TCR of any one of embodiments 735 to 830, comprising, from N- to C-terminus, (i) the anti-glyco-cMET variable heavy chain (VH), (ii) the first TCR domain, (iii) a cleavable peptide linker, (iv) the anti-glyco-cMET variable light chain (VL), and (v) the second TCR domain.

832. The chimeric TCR of any one of embodiments 735 to 830, comprising, from N- to C-terminus, (i) the anti-glyco- cMET variable heavy chain (VH), (ii) the second TCR domain, (iii) a cleavable peptide linker, (iv) the anti-glyco-cMET common light chain (CL), and (v) first second TCR domain.

833. The chimeric TCR of any one of embodiments 735 to 830, comprising, from N- to C-terminus, (i) the anti-glyco-cMET variable light chain (VL), (ii) the first TCR domain, (iii) a cleavable peptide linker, (iv) the anti-glyco-cMET variable heavy chain (VH), and (v) the second TCR domain.

834. The chimeric TCR of any one of embodiments 735 to 830, comprising, from N- to C-terminus, (i) the anti-glyco-cMET variable light chain (VL), (ii) the second TCR domain, (iii) a cleavable peptide linker, (iv) the anti-glyco-cMET variable heavy chain (VH), and (v) the first TCR domain.

835. A nucleic acid comprising a coding region for an anti-glyco-cMET antibody or antigen-binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, or the chimeric TCR of any one of embodiments 735 to 834.

836. The nucleic acid of embodiment 835 in which the coding region is codon-optimized for expression in a human cell.

837. A vector comprising the nucleic acid of embodiment 835 or embodiment 836.

838. The vector of embodiment 837 which is a viral vector.

839. The vector of embodiment 838 wherein the viral vector is a lentiviral vector.

840. A host cell engineered to express the nucleic acid of embodiment 835 or embodiment 836.

841. The host cell of embodiment 840, which is a human T-cell engineered to express the CAR of any one of embodiments 689 to 724.

842. The host cell of embodiment 840, which is a human NK cell engineered to express the CAR of any one of embodiments 689 to 724.

843. The host cell of embodiment 840, which is a human T-cell engineered to express the chimeric TCR of any one of embodiments 735 to 834.

844. A host cell comprising the vector of any one of embodiments 837 to 839.

845. The host cell of embodiment 844 which is a T-cell and wherein the vector encodes the CAR of any one of embodiments 690 to 725.

846. The host cell of embodiment 844 which is a T-cell and wherein the vector encodes the chimeric TCR of any one of embodiments 735 to 834.

847. A pharmaceutical composition comprising (a) the anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, or the host cell of any one of embodiments 840 to 846, and (b) a physiologically suitable buffer, adjuvant, diluent, or combination thereof.

848. A method of treating cancer comprising administering to a subject in need thereof an effective amount of the anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847.

849. The method of embodiment 848, wherein the subject is suffering from lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.

850. The method of embodiment 849, wherein the subject is suffering from lung cancer.
851. The method of embodiment 849, wherein the subject is suffering from breast cancer.
852. The method of embodiment 849, wherein the subject is suffering from pancreatic cancer.
853. The method of embodiment 849, wherein the subject is suffering from ovarian cancer.
854. The method of embodiment 849, wherein the subject is suffering from cholangiocarcinoma.
855. The method of embodiment 849, wherein the subject is suffering from colon cancer.
856. The method of embodiment 849, wherein the subject is suffering from thyroid cancer.
857. The method of embodiment 849, wherein the subject is suffering from liver cancer.
858. The method of embodiment 849, wherein the subject is suffering from gastric carcinoma.
859. A method of detecting cancer in a biological sample, comprising contacting a sample with an anti-glyco-cMET antibody or antigen-binding fragment according to any one of embodiments 1 to 657 and detecting binding of the anti-glyco-cMET antibody or antigen-binding fragment.
860. The method of embodiment 859, further comprising quantitating the binding of the anti-glyco-cMET antibody or antigen-binding fragment.
861. The method of embodiment 859 or embodiment 860, wherein the binding is compared to a normal tissue control as a negative/baseline control and/or to a cancerous tissue control as a positive control.
862. The method of any one of embodiments 859 to 861, wherein the cancer is lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.
863. The method of embodiment 862, wherein the cancer is lung cancer.
864. The method of embodiment 862, wherein the cancer is breast cancer.
865. The method of embodiment 862, wherein the cancer is pancreatic cancer.
866. The method of embodiment 862, wherein the cancer is ovarian cancer.
867. The method of embodiment 862, wherein the cancer is cholangiocarcinoma.
868. The method of embodiment 862, wherein the cancer is colon cancer.
869. The method of embodiment 862, wherein the cancer is thyroid cancer.
870. The method of embodiment 862, wherein the cancer is liver cancer.

871. The method of embodiment 862, wherein the cancer is gastric carcinoma.

872. The method of any one of embodiments 848 to 868, when depending from any one of embodiments 670 to 673, which further comprises administering to the subject genetically modified T-cells engineered to express a CAR comprising a NKG2D receptor capable of specifically binding the MIC protein domain.

873. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use as a medicament.

874. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of cancer, optionally wherein the cancer is lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.

875. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of lung cancer.

876. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of

embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of breast cancer.

877. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of pancreatic cancer.

878. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of ovarian cancer.

879. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of cholangiocarcinoma.

880. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of colon cancer.

881. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric

TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of thyroid cancer.

882. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of liver cancer.

883. The anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for use in the treatment of gastric carcinoma.

884. Use of the anti-glyco-cMET antibody or antigen binding fragment of any of embodiments 1 to 657, the bispecific antibody of any one of embodiments 658 to 663, the fusion protein of any one of embodiments 664 to 688, the CAR of any one of embodiments 689 to 724, the antibody-drug conjugate of any one of embodiments 725 to 734, the chimeric TCR of any one of embodiments 735 to 834, the nucleic acid of embodiment 835 or embodiment 836, the vector of any one of embodiments 837 to 839, the host cell of any one of embodiments 840 to 846, or the pharmaceutical composition of embodiment 847 for the manufacture of a medicament for the treatment of cancer, optionally wherein the cancer is lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.

885. The use according to embodiment 884, wherein the cancer is lung cancer.

886. The use according to embodiment 884, wherein the cancer is breast cancer.

887. The use according to embodiment 884, wherein the cancer is pancreatic cancer.

888. The use according to embodiment 884, wherein the cancer is ovarian cancer.

889. The use according to embodiment 884, wherein the cancer is cholangiocarcinoma.

890. The use according to embodiment 884, wherein the cancer is colon cancer.

891. The use according to embodiment 884, wherein the cancer is thyroid cancer.
892. The use according to embodiment 884, wherein the cancer is liver cancer.
893. The use according to embodiment 884, wherein the cancer is gastric carcinoma.
894. A peptide of 13-30 amino acids in length comprising a cMET peptide comprising PTKSFISGGSTITGVGKLN (SEQ ID NO:286), or a fragment thereof comprising amino acids corresponding to amino acids 9 and 10 of PTKSFISGGSTITGVGKLN (SEQ ID NO:286).
895. The peptide of embodiment 894 which is 15-25 amino acids in length.
896. The peptide of embodiment 894 which is 18-25 amino acids in length.
897. The peptide of any one of embodiments 894 to 896, wherein the fragment of the cMET peptide comprises amino acids 7-11, 7-12, 7-13, 7-14, 7-15, 7-16, 7-17, 7-18, 7-19, or 7-20 of SEQ ID NO:286.
898. The peptide of any one of embodiments 894 to 897, wherein the fragment of the cMET peptide comprises amino acids 6-11, 6-12, 6-13, 6-14, 6-15, 6-16, 6-17, 6-18, 6-19, or 6-20 of SEQ ID NO:286.
899. The peptide of any one of embodiments 894 to 898, wherein the fragment of the cMET peptide comprises amino acids 5-11, 5-12, 5-13, 5-14, 5-15, 5-16, 5-17, 5-18, 5-19, or 5-20 of SEQ ID NO:286.
900. The peptide of any one of embodiments 894 to 899, wherein the fragment of the cMET peptide comprises amino acids 4-11, 4-12, 4-13, 4-14, 4-15, 4-16, 4-17, 4-18, 4-19, or 4-20 of SEQ ID NO:286.
901. The peptide of any one of embodiments 894 to 900, wherein the fragment of the cMET peptide comprises amino acids 3-11, 3-12, 3-13, 3-14, 3-15, 3-16, 3-17, 3-18, 3-19, or 3-20 of SEQ ID NO:286.
902. The peptide of any one of embodiments 894 to 901, wherein the fragment of the cMET peptide comprises amino acids 2-11, 2-12, 2-13, 2-14, 2-15, 2-16, 2-17, 2-18, 2-19, or 2-20 of SEQ ID NO:286.
903. The peptide of any one of embodiments 894 to 902, wherein the fragment of the cMET peptide comprises amino acids 1-11, 1-12, 1-13, 1-14, 1-15, 1-16, 1-17, 1-18, 1-19, or 1-20 of SEQ ID NO:286.
904. The peptide of any one of embodiments 894 to 903 which comprises PTKSFISGGSTITGVGKLN (SEQ ID NO:286).
905. The peptide of any one of embodiments 894 to 904 which consists of PTKSFISGGSTITGVGKLN (SEQ ID NO:286).
906. The peptide of any one of embodiments 894 to 905 which is O-glycosylated at the threonine corresponding to position 9 of PTKSFISGGSTITGVGKLN (SEQ ID NO:286), and/or the threonine corresponding to position 10 of PTKSFISGGSTITGVGKLN (SEQ ID NO:286).

907. The peptide of embodiment 906, wherein the O-glycosylation comprises or consists of GalNAc.

908. A peptide of 13-30 amino acids in length comprising a cMET peptide PTKSFISGG**ST**ITGVGKLN (SEQ ID NO:285) that has been O-glycosylated on the serine and threonine residues shown with bold and underlined text, or a fragment thereof comprising amino acids corresponding to amino acids 9 and 10 of PTKSFISGG**ST**ITGVGKLN (SEQ ID NO:285).

909. The peptide of embodiment 908 which is 15-25 amino acids in length.

910. The peptide of embodiment 908 which is 18-25 amino acids in length.

911. The peptide of any one of embodiments 908 to 910, wherein the fragment of the cMET peptide comprises amino acids 7-11, 7-12, 7-13, 7-14, 7-15, 7-16, 7-17, 7-18, 7-19, or 7-20 of SEQ ID NO:285.

912. The peptide of any one of embodiments 908 to 911, wherein the fragment of the cMET peptide comprises amino acids 6-11, 6-12, 6-13, 6-14, 6-15, 6-16, 6-17, 6-18, 6-19, or 6-20 of SEQ ID NO:285.

913. The peptide of any one of embodiments 908 to 912, wherein the fragment of the cMET peptide comprises amino acids 5-11, 5-12, 5-13, 5-14, 5-15, 5-16, 5-17, 5-18, 5-19, or 5-20 of SEQ ID NO:285.

914. The peptide of any one of embodiments 908 to 913, wherein the fragment of the cMET peptide comprises amino acids 4-11, 4-12, 4-13, 4-14, 4-15, 4-16, 4-17, 4-18, 4-19, or 4-20 of SEQ ID NO:285.

915. The peptide of any one of embodiments 908 to 914, wherein the fragment of the cMET peptide comprises amino acids 3-11, 3-12, 3-13, 3-14, 3-15, 3-16, 3-17, 3-18, 3-19, or 3-20 of SEQ ID NO:285.

916. The peptide of any one of embodiments 908 to 915, wherein the fragment of the cMET peptide comprises amino acids 2-11, 2-12, 2-13, 2-14, 2-15, 2-16, 2-17, 2-18, 2-19, or 2-20 of SEQ ID NO:285.

917. The peptide of any one of embodiments 908 to 916, wherein the fragment of the cMET peptide comprises amino acids 1-11, 1-12, 1-13, 1-14, 1-15, 1-16, 1-17, 1-18, 1-19, or 1-20 of SEQ ID NO:285.

918. The peptide of any one of embodiments 908 to 917 which comprises PTKSFISGG**ST**ITGVGKLN (SEQ ID NO:285).

919. The peptide of any one of embodiments 908 to 918 which consists of PTKSFISGG**ST**ITGVGKLN (SEQ ID NO:285).

920. The peptide of any one of embodiments 908 to 919, wherein the O-glycosylation comprises or consists of GalNAc.

921. A composition comprising the peptide of any one of embodiments 894 to 920 and adjuvant.

922. The composition of embodiment 921, wherein the adjuvant comprises a Freund's adjuvant and/or an aluminum salt (e.g., aluminum hydroxide).

923. A method of generating antibodies against a tumor-associated form of cMET, comprising administering to an animal:

- (a) the peptide of any one of embodiments 906 to 920; or
- (b) the composition of embodiment 921 or 922 wherein the composition

comprises the peptide of any one of embodiments 906 to 920.

924. The method of embodiment 923, further comprising collecting antibodies from the animal following the administering step.

925. A method of eliciting an immune response against a tumor-associated form of cMET, comprising administering to a subject:

- (a) the peptide of any one of embodiments 906 to 920; or
- (b) the composition of embodiment 921 or 922 wherein the composition

comprises the peptide of any one of embodiments 906 to 920.

926. The method of any one of embodiments 923 to 925, wherein the animal is a mouse or a rabbit.

927. The anti-glyco-cMET antibody or antigen binding fragment, bispecific antibody, fusion protein, CAR, antibody-drug conjugate, the chimeric TCR, pharmaceutical composition method or use as described in any one of the preceding embodiments, wherein the determination of competing is made using an antibody competition assay, optionally wherein the assay is an assay described in Section 5.1.

928. The anti-glyco-cMET antibody or antigen binding fragment, bispecific antibody, fusion protein, CAR, antibody-drug conjugate, the chimeric TCR, pharmaceutical composition method or use of embodiment 927, wherein competing is present if the anti-glyco-cMET antibody or anti-glyco-LAMP1 antibody fragment decreases binding of a reference antibody by at least about 20% 30%, 40%, 50%, 60%, 70%, 80%, 90%, or 95% when tested at a reference antibody concentration that is 80% of maximal binding under the specific assay conditions used and a test antibody concentration that is 10-fold higher than the reference antibody concentration.

[0428] All publications, patents, patent applications and other documents cited in this application are hereby incorporated by reference in their entireties for all purposes to the same extent as if each individual publication, patent, patent application or other document were individually indicated to be incorporated by reference for all purposes. In the event that there is an inconsistency between the teachings of one or more of the references incorporated herein and the present disclosure, the teachings of the present specification are intended.

What is claimed is:

1. An anti-glyco-cMET antibody or antigen binding fragment that specifically binds to a cMET peptide PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285) that has been glycosylated with GalNAc on the serine and threonine residues shown with bold and underlined text (“the cMET glycopeptide”).

2. The anti-glyco-cMET antibody or antigen binding fragment of claim 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence and a light chain variable (VL) sequence of:

- (a) SEQ ID NO:1 and SEQ ID NO:2, respectively;
- (b) SEQ ID NO:23 and SEQ ID NO:24, respectively;
- (c) SEQ ID NO:45 and SEQ ID NO:46, respectively;
- (d) SEQ ID NO:67 and SEQ ID NO:68, respectively;
- (e) SEQ ID NO:89 and SEQ ID NO:90, respectively; or
- (f) SEQ ID NO:111 and SEQ ID NO:112, respectively;

for binding to the cMET glycopeptide.

3. The anti-glyco-cMET antibody or antigen binding fragment of claim 1, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of any one of SEQ ID NOS:264-275 and a light chain variable (VL) sequence of any one of SEQ ID NOS:276-284 for binding to the cMET glycopeptide.

4. The anti-glyco-cMET antibody or antigen binding fragment of any one of claims 1 to 3, which specifically binds to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

5. The anti-glyco-cMET antibody or antigen binding fragment of claim 4, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence and a light chain variable (VL) sequence of:

- (a) SEQ ID NO:1 and SEQ ID NO:2, respectively;
- (b) SEQ ID NO:23 and SEQ ID NO:24, respectively; or
- (c) SEQ ID NO:45 and SEQ ID NO:46, respectively,

for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

6. The anti-glyco-cMET antibody or antigen binding fragment of claim 4, wherein the anti-glyco-cMET antibody or antigen binding fragment competes with an antibody or antigen binding fragment comprising a heavy chain variable (VH) sequence of any one of SEQ ID NOS:264-275 and a light chain variable (VL) sequence of any one of SEQ ID NOS:276-284 for binding to COSMC knock-out T47D cells or COSMC knock-out A549 cells.

7. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen binding fragment according to any one of claims 1 to 6, comprising:

(a) a complementarity determining region (CDR) H1 comprising the amino acid sequence of SEQ ID NO:133, SEQ ID NO:139, SEQ ID NO:145, SEQ ID NO:205, or SEQ ID NO:253;

(b) a CDR-H2 comprising the amino acid sequence of SEQ ID NO:134, SEQ ID NO:140, SEQ ID NO:146, SEQ ID NO:206, or SEQ ID NO:254;

(c) a CDR-H3 comprising the amino acid sequence of SEQ ID NO:135, SEQ ID NO:141, SEQ ID NO:147, SEQ ID NO:207, or SEQ ID NO:255;

(d) a CDR-L1 comprising the amino acid sequence of SEQ ID NO:136, SEQ ID NO:142, SEQ ID NO:148, SEQ ID NO:208, or SEQ ID NO:256;

(e) a CDR-L2 comprising the amino acid sequence of SEQ ID NO:137, SEQ ID NO:143, SEQ ID NO:149, SEQ ID NO:209, or SEQ ID NO:257; and

(f) a CDR-L3 comprising the amino acid sequence of SEQ ID NO:138, SEQ ID NO:144, SEQ ID NO:150, SEQ ID NO:210, or SEQ ID NO:258.

8. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 7, which comprises:

(a) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:3-5, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:6-8, respectively;

(b) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:9-11, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:12-14, respectively; or

(c) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:15-17, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:18-20, respectively.

9. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 7, which comprises:

(a) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:25-27, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:28-30, respectively;

(b) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:31-33, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:34-36, respectively; or

(c) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:37-39, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:40-42, respectively.

10. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 7, which comprises:

(a) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:47-49, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:50-52, respectively;

(b) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:53-55, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:56-58, respectively; or

(c) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:59-61, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:62-64, respectively.

11. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 6, comprising:

(a) a complementarity determining region (CDR) H1 comprising the amino acid sequence of SEQ ID NO:151, SEQ ID NO:157, SEQ ID NO:163, SEQ ID NO:211, or SEQ ID NO:259;

(b) a CDR-H2 comprising the amino acid sequence of SEQ ID NO:152, SEQ ID NO:158, SEQ ID NO:164, SEQ ID NO:212, or SEQ ID NO:260;

(c) a CDR-H3 comprising the amino acid sequence of SEQ ID NO:153, SEQ ID NO:159, SEQ ID NO:165, SEQ ID NO:213, or SEQ ID NO:261;

(d) a CDR-L1 comprising the amino acid sequence of SEQ ID NO:154, SEQ ID NO:160, SEQ ID NO:166, SEQ ID NO:214, or SEQ ID NO:262;

(e) a CDR-L2 comprising the amino acid sequence of SEQ ID NO:155, SEQ ID NO:161, SEQ ID NO:167, SEQ ID NO:215, or SEQ ID NO:263; and

(f) a CDR-L3 comprising the amino acid sequence of SEQ ID NO:156, SEQ ID NO:162, SEQ ID NO:168, SEQ ID NO:216, or SEQ ID NO:342.

12. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 6, which comprises:

(a) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:69-71, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:72-74, respectively;

(b) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:75-77, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:78-80, respectively; or

(c) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:81-83, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:84-86, respectively.

13. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 6, which comprises:

(a) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:91-93, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:94-96, respectively;

(b) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:97-99, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:100-102, respectively; or

(c) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:103-105, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:106-108, respectively.

14. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 6, which comprises:

(a) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:113-115, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:116-118, respectively;

(b) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:119-121, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:122-124, respectively; or

(c) a VH comprising CDR-H1, CDR-H2, and CDR-H3 having the amino sequences of SEQ ID NOS:125-127, respectively, and a VL comprising CDR-L1, CDR-L2, and CDR-L3 having the amino acid sequences of SEQ ID NOS:128-130, respectively.

15. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 14, which is a chimeric or humanized antibody or antigen-binding fragment of a chimeric or humanized antibody.

16. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 15, which comprises:

(a) a VH comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:1 and a VL comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:2;

(b) a VH comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:23 and a VL comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:24;

(c) a VH comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:45 and a VL comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:46;

(d) a VH comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:67 and a VL comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:68;

(e) a VH comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:89 and a VL comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:90; or

(f) a VH comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:111 and a VL comprising an amino acid sequence having at least 95% sequence identity to SEQ ID NO:112.

17. An anti-glyco-cMET antibody or antigen-binding fragment, which is optionally an anti-cMET antibody or antigen-binding fragment according to any one of claims 1 to 16, that competes with a reference antibody or antigen binding fragment comprising:

(a) a heavy chain variable (VH) sequence of SEQ ID NO:1 and a light chain variable (VL) sequence of SEQ ID NO:2;

(b) a heavy chain variable (VH) sequence of SEQ ID NO:23 and a light chain variable (VL) sequence of SEQ ID NO:24;

(c) a heavy chain variable (VH) sequence of SEQ ID NO:45 and a light chain variable (VL) sequence of SEQ ID NO:46;

(d) a heavy chain variable (VH) sequence of SEQ ID NO:67 and a light chain variable (VL) sequence of SEQ ID NO:68;

(e) a heavy chain variable (VH) sequence of SEQ ID NO:89 and a light chain variable (VL) sequence of SEQ ID NO:90;

(f) a heavy chain variable (VH) sequence of SEQ ID NO:111 and a light chain variable (VL) sequence of SEQ ID NO:112; or

(g) a humanized heavy chain variable (VH) sequence of any one of SEQ ID NOS:264-275 and a humanized light chain variable (VL) sequence of SEQ ID NOS:276-284,

for binding to a cMET peptide PTKSFISGGSTITGVGKLN (SEQ ID NO:285) that has been glycosylated with GalNAc on the serine and threonine residues shown with bold and underlined text (“the cMET glycopeptide”), the anti-glyco-cMET antibody or antigen-binding fragment comprising:

(h) a VH sequence with first, second and third CDR means within the VH sequence; and

(i) a VL sequence with fourth, fifth and sixth CDR means within the VL sequence,

wherein the first, second, third, fourth, fifth, and sixth CDR means cooperate to effect binding of the anti-glyco-cMET antibody or antigen-binding fragment to the cMET glycopeptide.

18. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 17, which preferentially binds to a glyco-cMET epitope that is overexpressed on cancer cells as compared to normal cells.

19. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 18, which specifically binds to a cMET peptide PTKSFISGGSTITGVGKLN (SEQ ID NO:285) that has been glycosylated with STn on the serine and threonine residues shown with bold and underlined text.

20. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 1 to 18, which does not specifically bind to a cMET peptide PTKSFISGGSTITGVGKLN (SEQ ID NO:285) that has been glycosylated with STn on the serine and threonine residues shown with bold and underlined text.

21. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 20, which binds to the cMET glycopeptide with a binding affinity (KD) of 1 nM to 200 nM as measured by surface plasmon resonance or bio-layer interferometry.

22. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 21, which does not specifically bind to the unglycosylated cMET peptide PTKSFISGGSTITGVGKLN (SEQ ID NO:286) (the “unglycosylated cMET peptide”).

23. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 22, which does not specifically bind to the MUC1 tandem repeat (VTSAPDTRPAPGSTAPPAHG)₃ (SEQ ID NO:288) that has been glycosylated in vitro using purified recombinant human glycosyltransferases GalNAc-T1, GalNAc-T2, and GalNAc-T4 (“the first MUC1 glycopeptide”).

24. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 23, which does not specifically bind to the MUC1 peptide TAPPAHGV**TS**APD**TR**PAPG**ST**APPAHGVT (SEQ ID NO:289) that has been glycosylated in vitro with GalNAc on the serine and threonine residues shown with bold and underlined text (the “second MUC1 glycopeptide”).

25. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 24, which does not specifically bind to the CD44v6 peptide GYRQ**TP**KEDSH**ST**TGTAAA (SEQ ID NO:345) that has been glycosylated in vitro with GalNAc on the threonine and serine residues shown with bold and underlined text (the “CD44v6 glycopeptide”).

26. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 25, which does not specifically bind to the MUC4 peptide CTIPSTAMHTR**STA**APIILP (SEQ ID NO:291) that has been glycosylated in vitro with GalNAc on the serine and threonine residues shown with bold and underlined text (the “MUC4 glycopeptide”).

27. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 26, which does not specifically bind to the LAMP1 peptide CEQDRP**SPT**TAPPAPPSPSP (SEQ ID NO:292) that has been glycosylated in vitro with GalNAc on the serine and threonine residues shown with bold and underlined text (the “LAMP1 glycopeptide”).

28. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 27, which is multivalent.

29. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 28, which is an antigen-binding fragment.

30. The anti-glyco-cMET antibody or antigen-binding fragment of claim 29, wherein the antigen-binding fragment is in the form of a single-chain variable fragment (scFv).

31. The anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 28, which is in the form of a multispecific antibody.

32. The anti-glyco-cMET antibody or antigen-binding fragment of claim 31, wherein the multispecific antibody is a bispecific antibody that binds to a second epitope that is different from the first epitope.

33. The anti-glyco-cMET antibody or antigen-binding fragment of claim 32, wherein the bispecific antibody is a bottle opener, mAb-Fv, mAb-scFv, central-scFv, one-armed central-scFv, or dual scFv format bispecific antibody.

34. The anti-glyco-cMET antibody or antigen-binding fragment of claim 32, wherein the bispecific antibody is a bispecific domain-exchanged antibody (e.g., a CrossMab), a Fab-arm exchange antibody, a bispecific T-cell engager (BiTE), or a dual-affinity retargeting molecule (DART).

35. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 32 to 34, wherein the second epitope is a cMET epitope.

36. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 32 to 34, wherein the second epitope is a cMET epitope that is overexpressed on cancer cells as compared to normal cells.

37. The anti-glyco-cMET antibody or antigen-binding fragment of any one of claims 32 to 34, wherein the second epitope is a T-cell epitope.

38. The anti-glyco-cMET antibody or antigen-binding fragment of claim 37, wherein the T-cell epitope comprises a CD3 epitope, a CD8 epitope, a CD16 epitope, a CD25 epitope, a CD28 epitope, or an NKG2D epitope.

39. A fusion protein comprising the amino acid sequence of the anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 38, operably linked to at least a second amino acid sequence.

40. A chimeric antigen receptor (CAR) comprising one or more antigen-binding fragments according to claim 29 or claim 30.

41. A chimeric antigen receptor (CAR), whose amino acid sequence comprises the amino acid sequence of SEQ ID NO:348, SEQ ID NO:339, SEQ ID NO:340, or SEQ ID NO:341.

42. An antibody-drug conjugate comprising the anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 38 or the fusion protein of claim 39 conjugated to a cytotoxic agent.

43. A chimeric T cell receptor (TCR) comprising
- (a) an antigen-binding fragment according to claim 29 or claim 30;
 - (b) a first polypeptide chain comprising a first TCR domain comprising a first TCR transmembrane domain from a first TCR subunit; and
 - (c) a second polypeptide chain comprising a second TCR domain comprising a second TCR transmembrane domain from a second TCR subunit.
44. A nucleic acid comprising a coding region for an anti-glyco-cMET antibody or antigen-binding fragment of any of claims 1 to 38, the fusion protein of claim 39, the CAR of claim 40 or claim 41, or the chimeric TCR of claim 43.
45. A vector comprising the nucleic acid of claim 44.
46. A host cell engineered to express the nucleic acid of claim 44 or comprising the vector of claim 45.
47. A pharmaceutical composition comprising (a) the anti-glyco-cMET antibody or antigen binding fragment of any of claims 1 to 38, the fusion protein of claim 39, the CAR of claim 40 or claim 41, the antibody-drug conjugate of claim 42, the chimeric TCR of claim 43, the nucleic acid of claim 44, the vector of claim 45, or the host cell of claim 46, and (b) a physiologically suitable buffer, adjuvant, diluent, or combination thereof.
48. A method of treating cancer comprising administering to a subject in need thereof an effective amount of the anti-glyco-cMET antibody or antigen binding fragment of any of claims 1 to 38, the fusion protein of claim 39, the CAR of claim 40 or claim 41, the antibody-drug conjugate of claim 42, the chimeric TCR of claim 43, the nucleic acid of claim 44, the vector of claim 45, the host cell of claim 46, or the pharmaceutical composition of claim 47.
49. The method of claim 48, wherein the subject is suffering from lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.
50. A method of detecting cancer in a biological sample, comprising contacting a sample with an anti-glyco-cMET antibody or antigen-binding fragment according to any one of claims 1 to 38 and detecting binding of the anti-glyco-cMET antibody or antigen-binding fragment.
51. The method of claim 50, wherein the cancer is lung cancer, breast cancer, pancreatic cancer, ovarian cancer, cholangiocarcinoma, colon cancer, thyroid cancer, liver cancer, or gastric carcinoma.

52. A peptide of 13-30 amino acids in length comprising a cMET peptide comprising PTKSFISGGSTITGVGKNLN (SEQ ID NO:286), or a fragment thereof comprising amino acids corresponding to amino acids 9 and 10 of PTKSFISGGSTITGVGKNLN (SEQ ID NO:286).

53. A peptide of 13-30 amino acids in length comprising a cMET peptide PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285) that has been O-glycosylated on the serine and threonine residues shown with bold and underlined text, or a fragment thereof comprising amino acids corresponding to amino acids 9 and 10 of PTKSFISGG**ST**ITGVGKNLN (SEQ ID NO:285).

54. A composition comprising the peptide of claim 52 or claim 53 and an adjuvant.

55. A method of generating antibodies against a tumor-associated form of cMET, comprising administering to an animal:

(a) the peptide of claim 53; or

(b) The composition of claim 54, wherein the composition comprises the peptide of claim 53.

56. A method of eliciting an immune response against a tumor-associated form of cMET, comprising administering to a subject:

(a) the peptide of claim 53; or

(b) the composition of claim 54, wherein the composition comprises the peptide of claim 53.

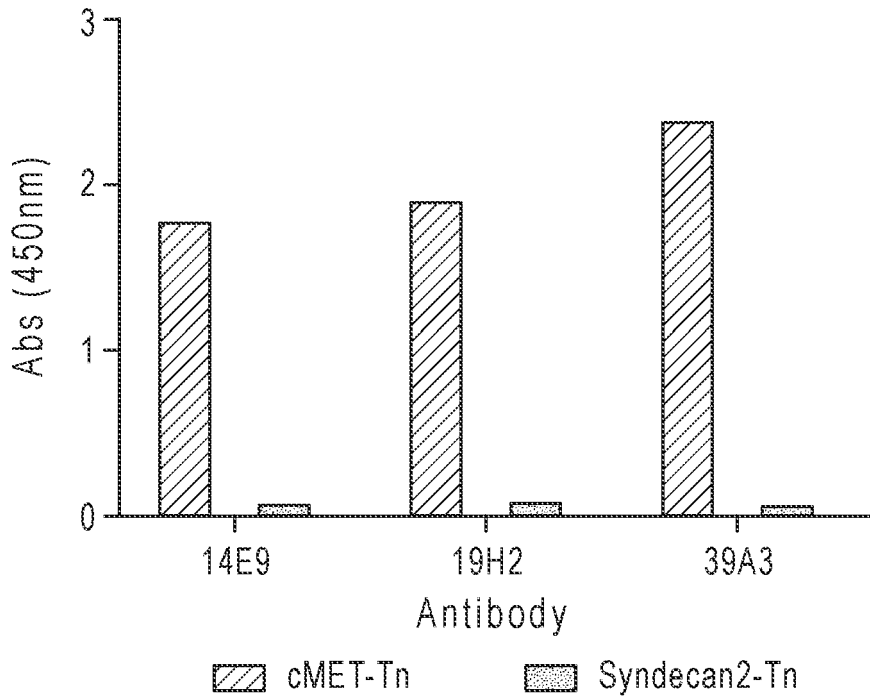


FIG. 1

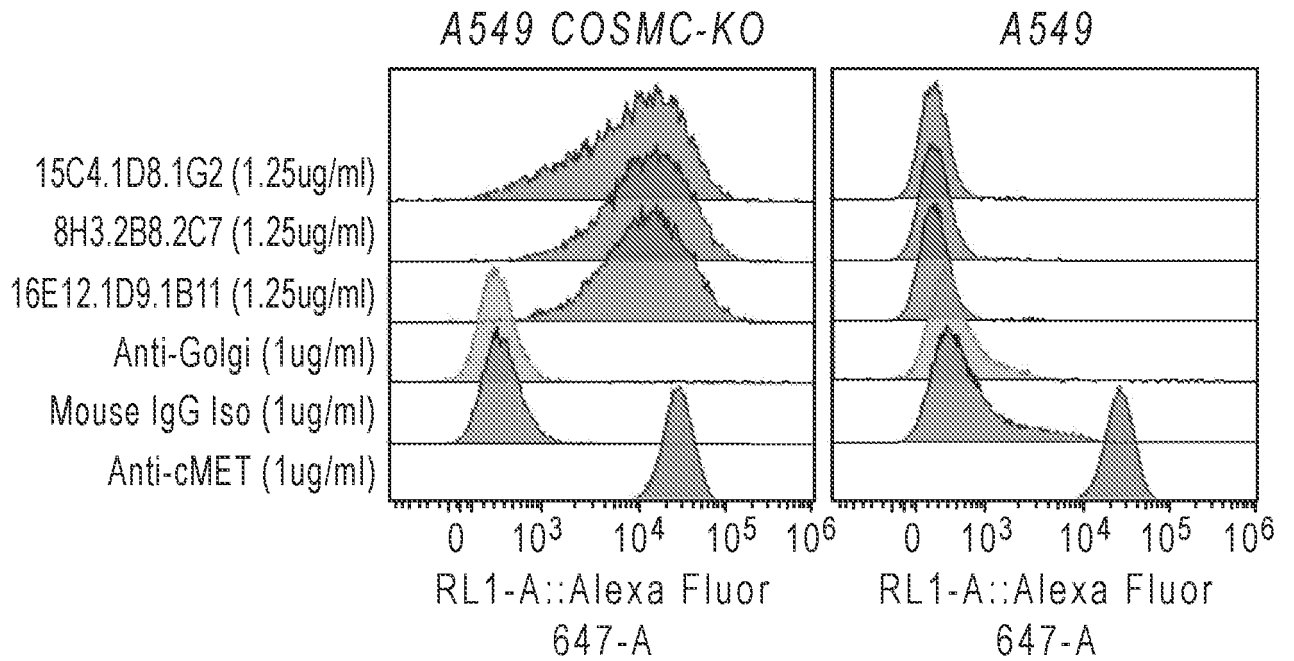


FIG. 2A

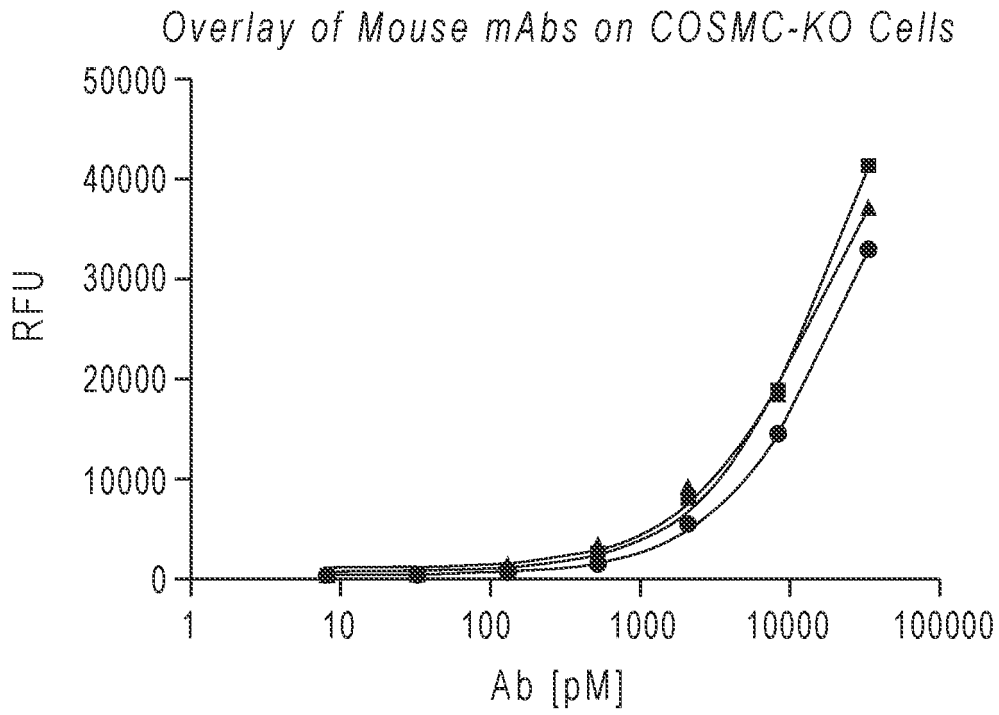


FIG. 2B-1

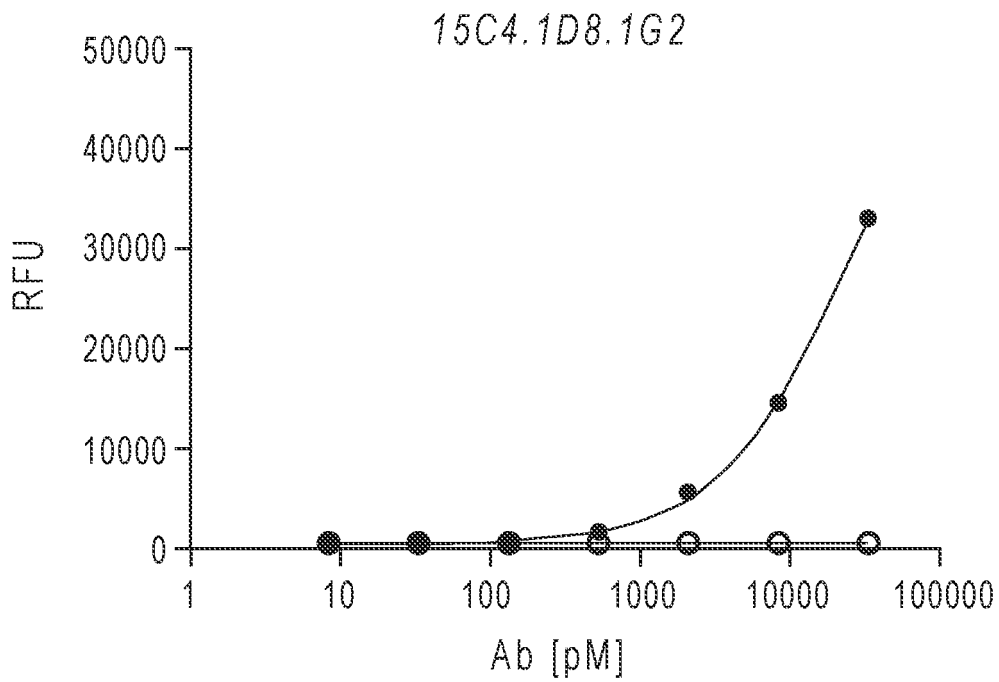


FIG. 2B-2

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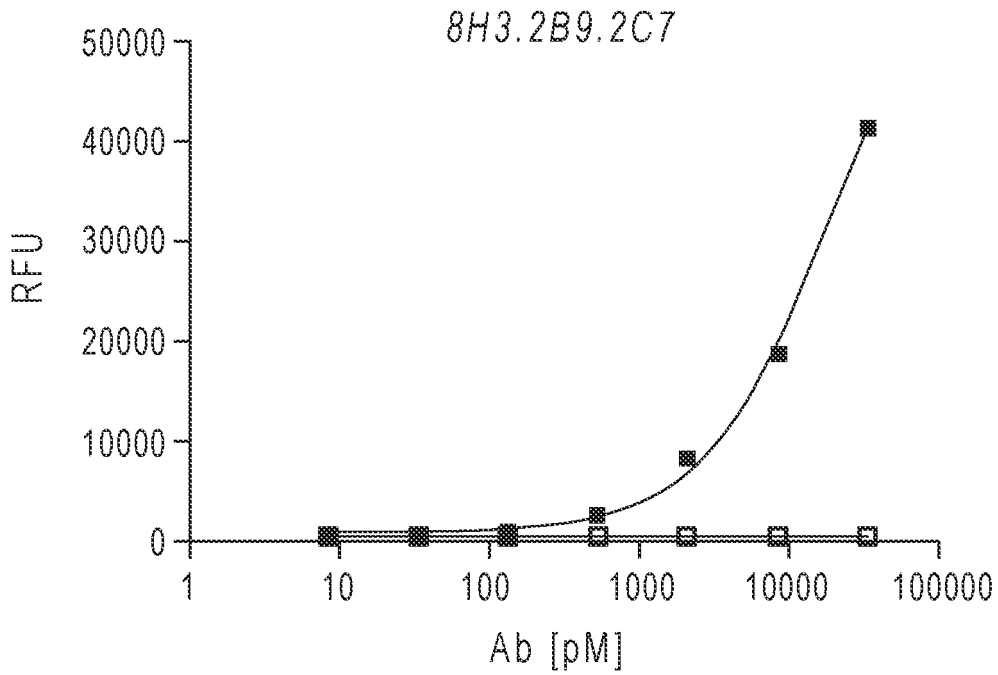


FIG. 2B-3

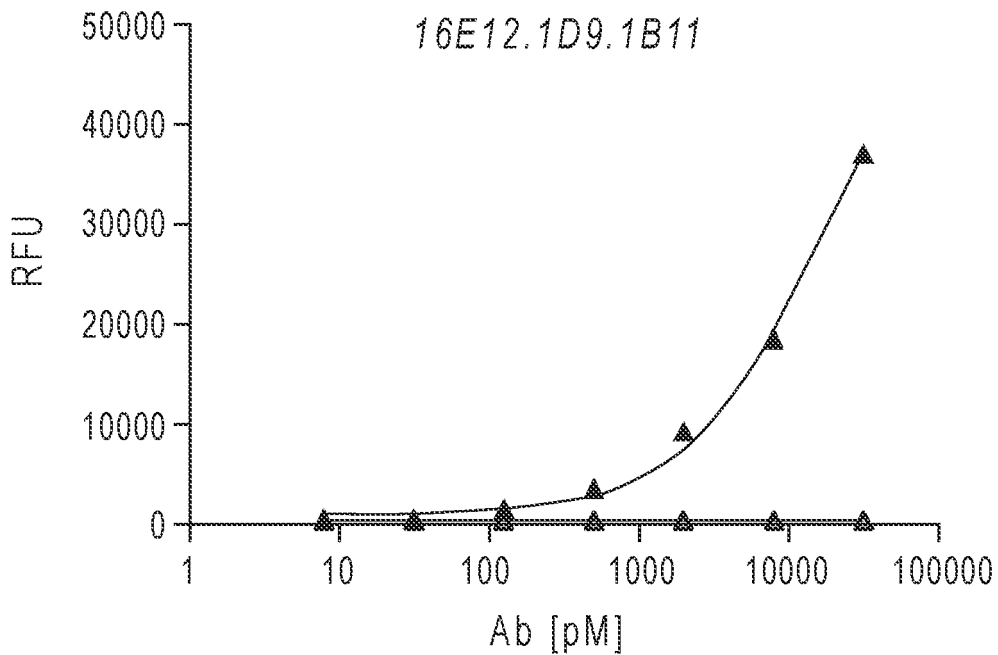


FIG. 2B-4

FIG. 2B-5

Antibody	A549	
	COSMC-KO	A549
15C4.1D8.1G2	●	⊙
8H3.2B9.2C7	■	⊠
16E12.1D9.1B11	▲	⊡

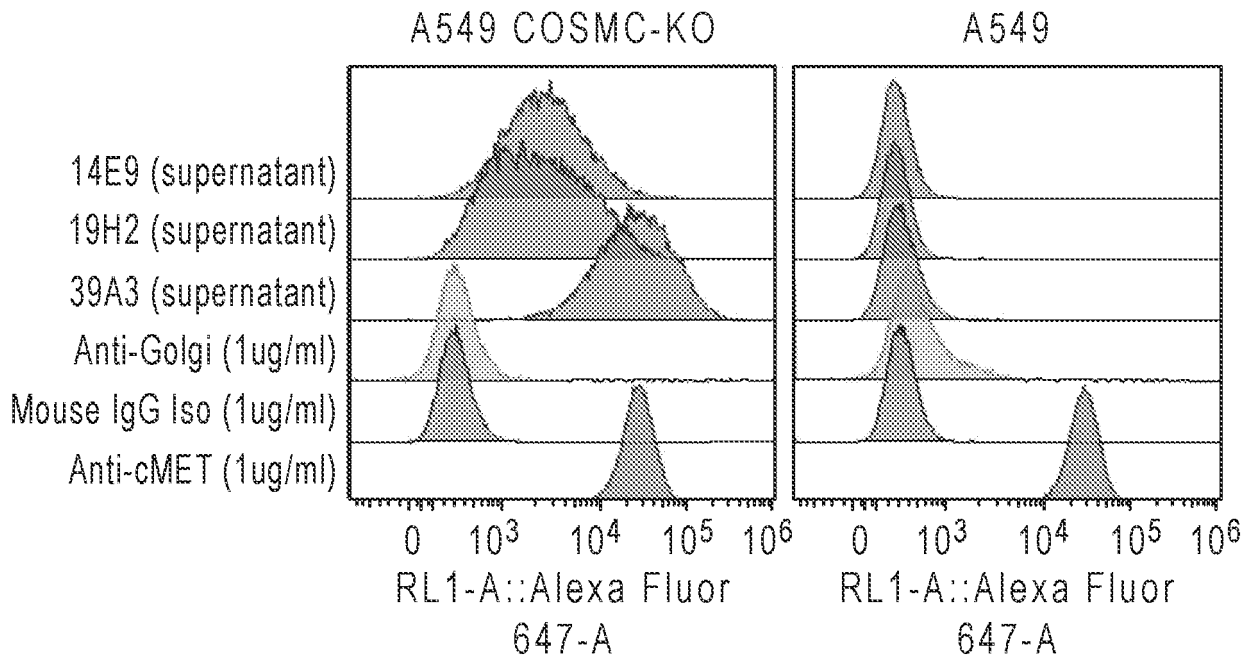


FIG. 3A

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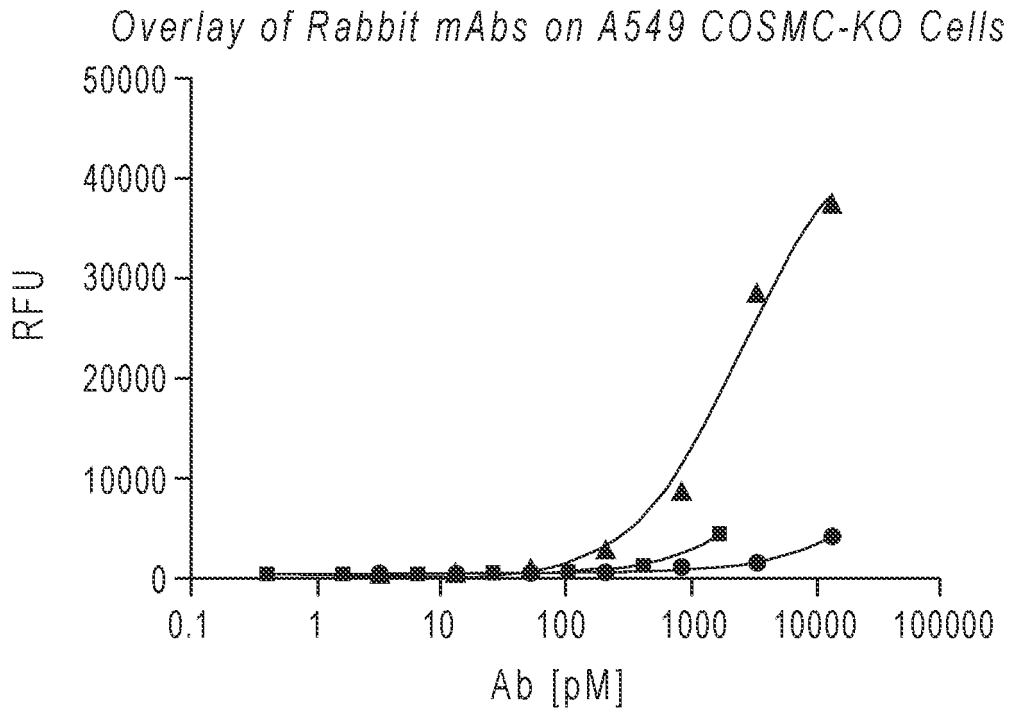


FIG. 3B-1

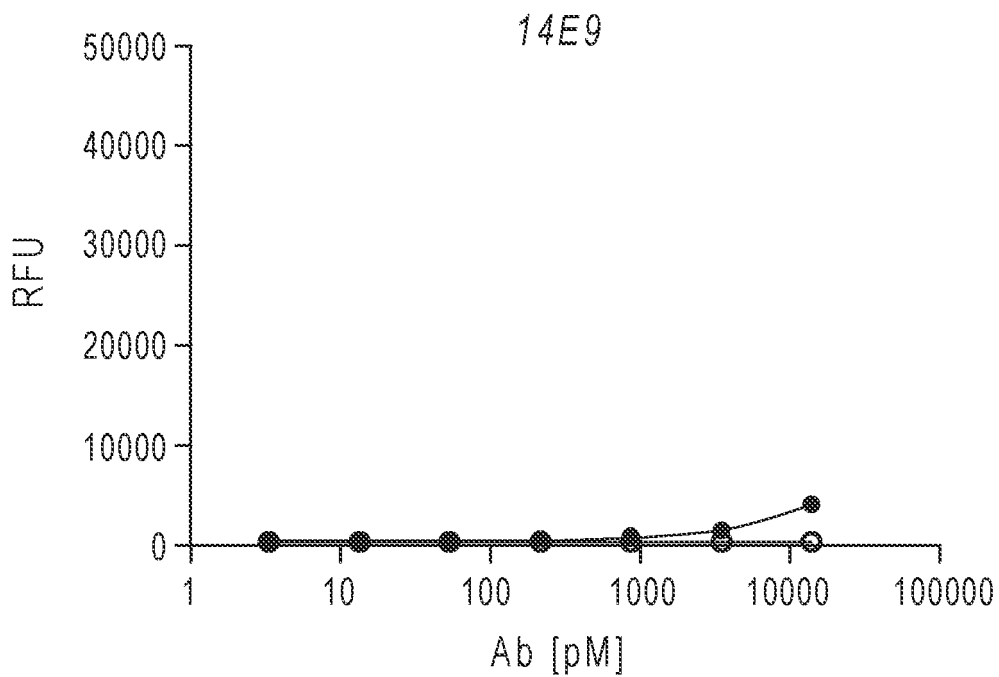


FIG. 3B-2

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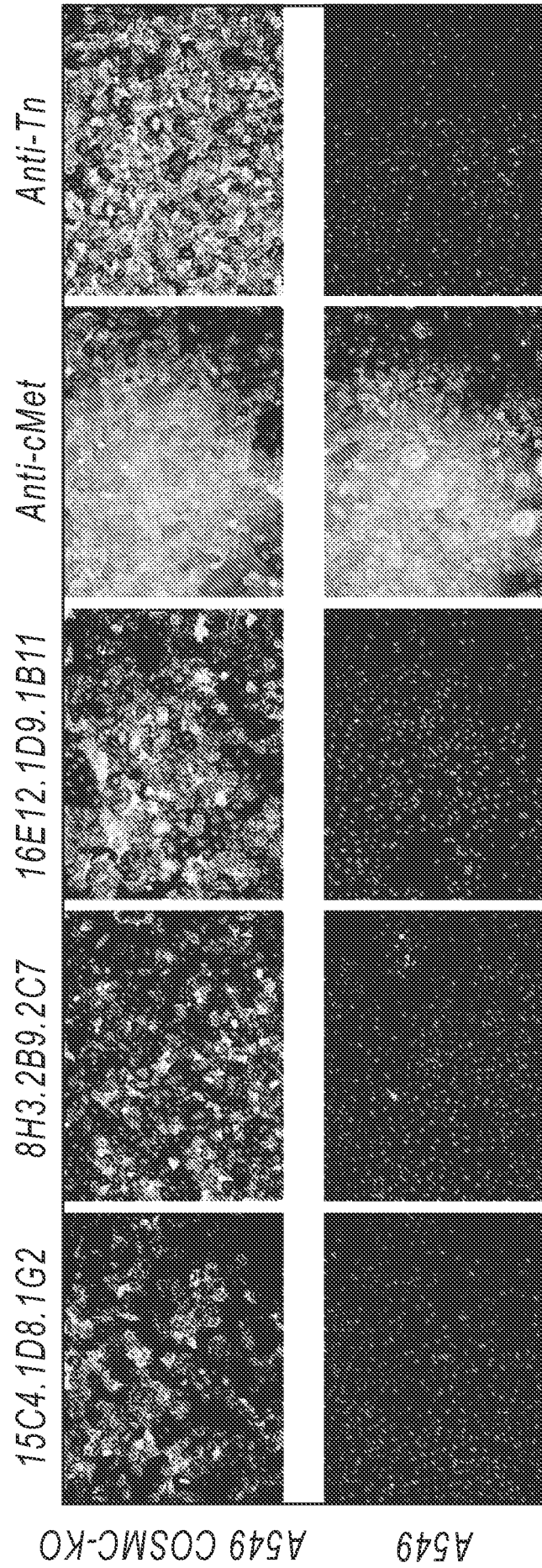


FIG. 4A

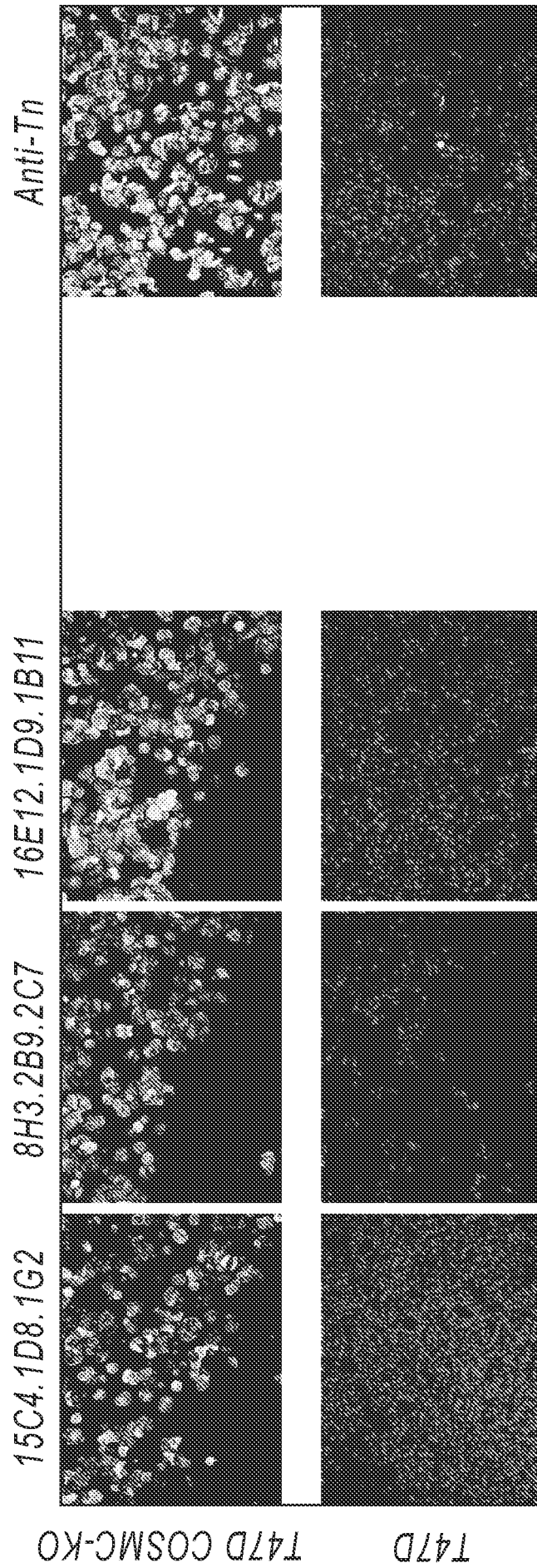


FIG. 4B

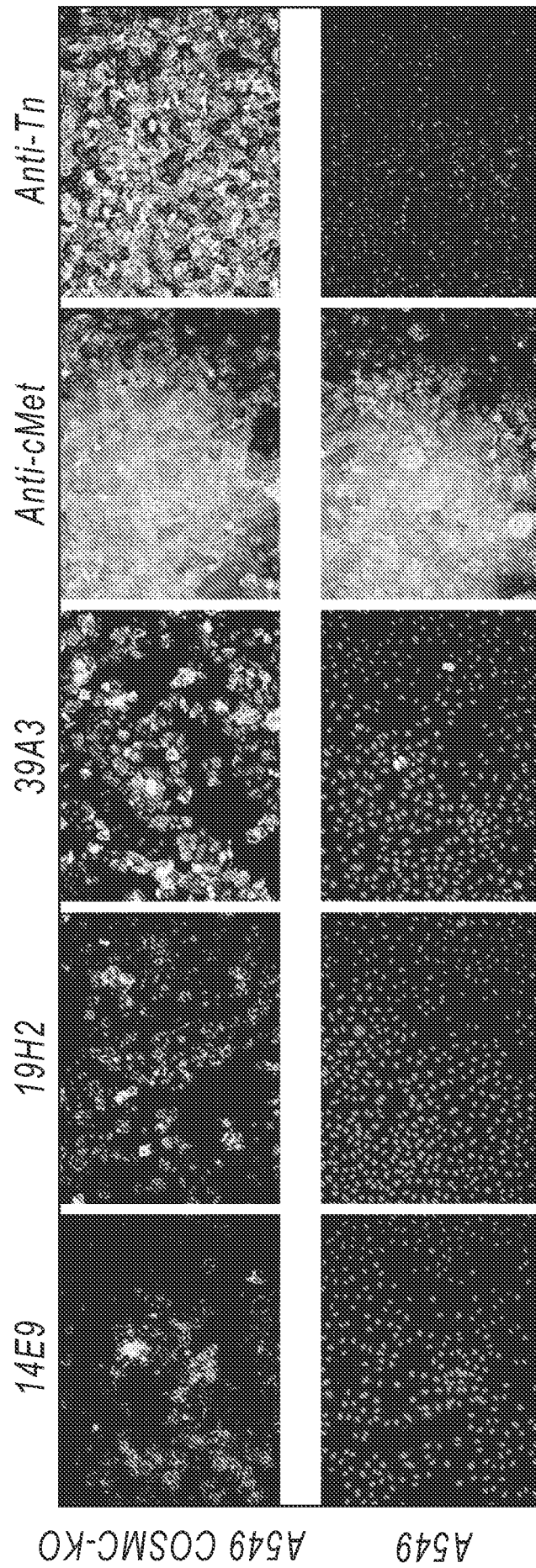
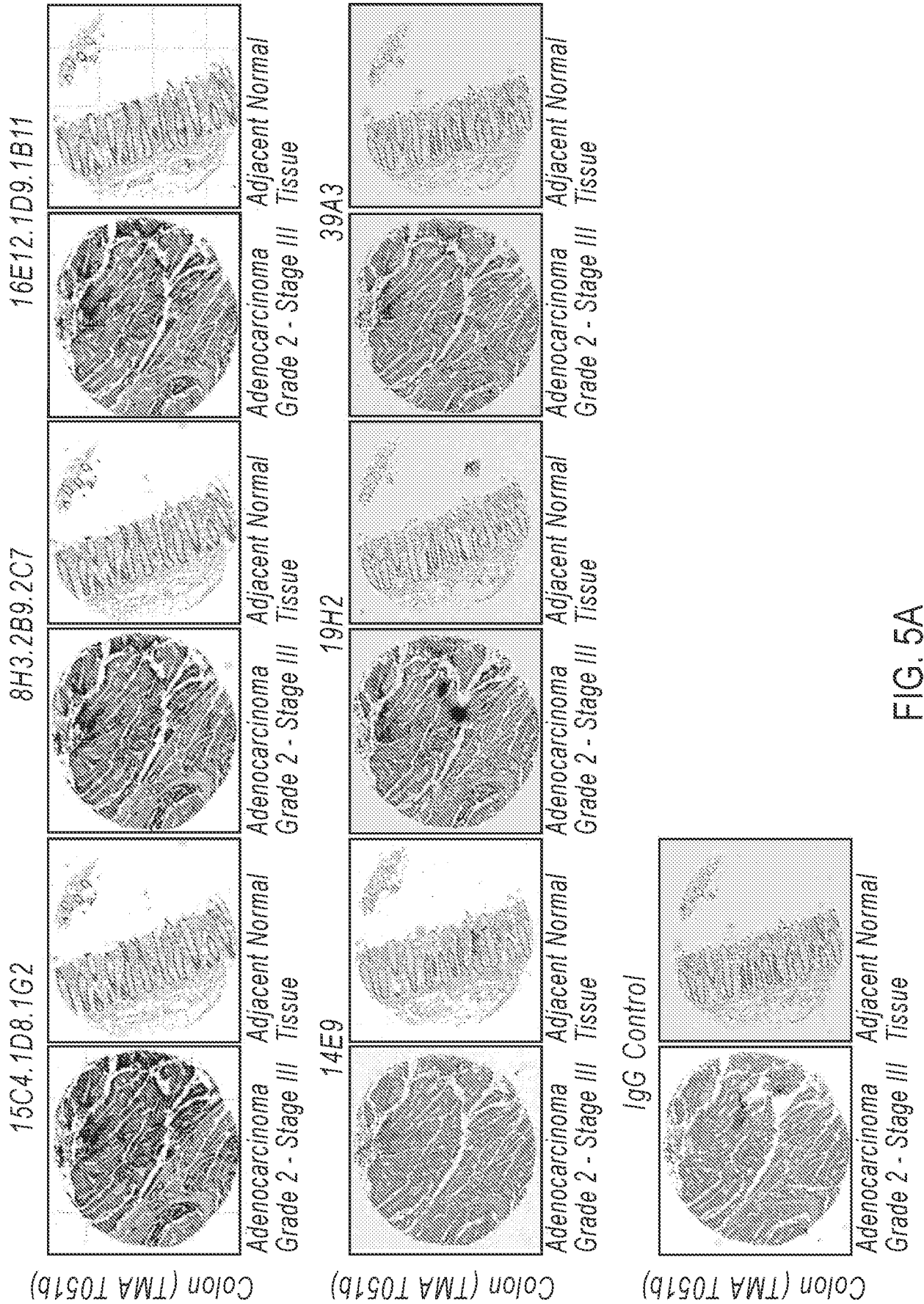


FIG. 4C



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Colon (TMA T051b)

Antibody	Cases	Positive (Surface)	Negative
15C4.1D8.1G2	6	6 (100%)	0 (0%)
8H3.2B9.2C7	6	6 (100%)	0 (0%)
16E12.1D9.1B11	6	6 (100%)	0 (0%)

Colon (TMA051b)

Antibody	Cases	Positive (Surface)	Negative
14E9	6	3 (50%)	3 (50%)
19H2	6	3 (50%)	3 (50%)
39A3	6	3 (50%)	3 (50%)

FIG. 5B

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Human Cancer Tissue

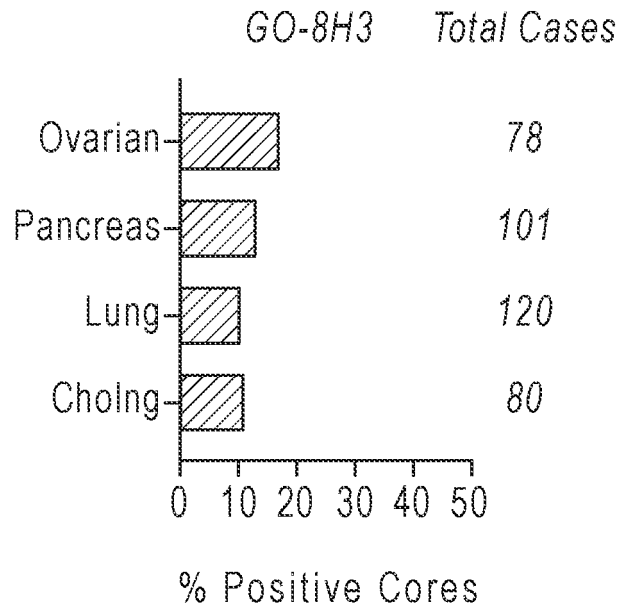
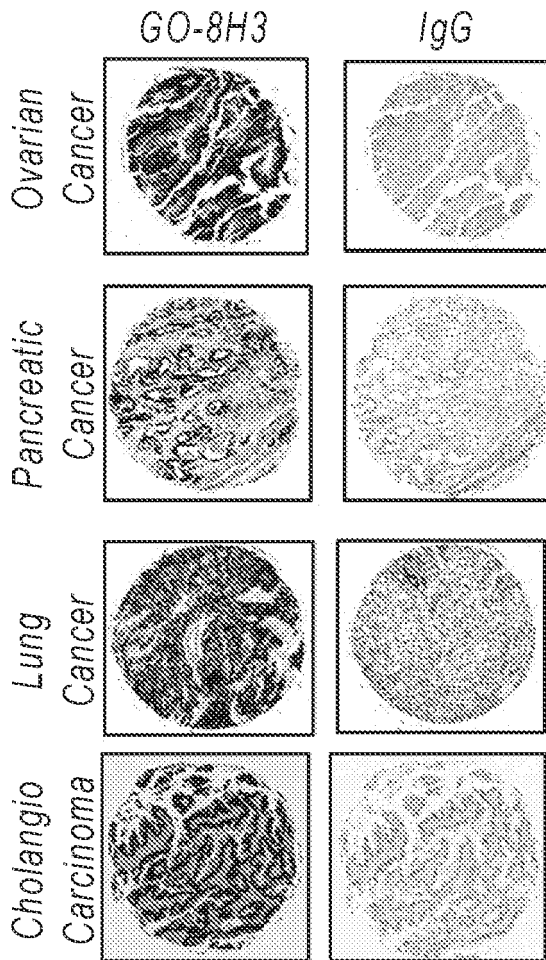


FIG. 6A-2

FIG. 6A-1

Human Normal Tissue

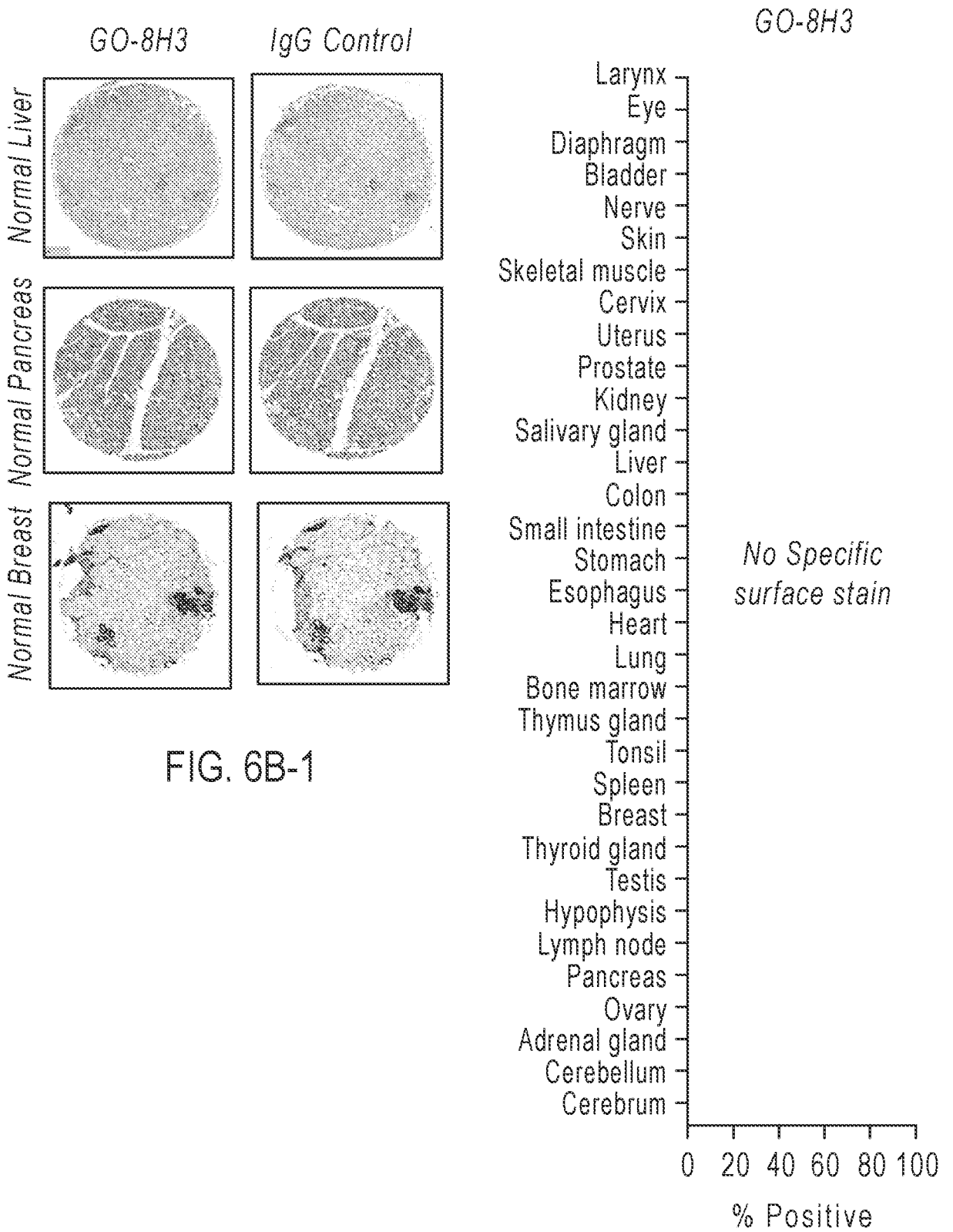


FIG. 6B-1

FIG. 6B-2

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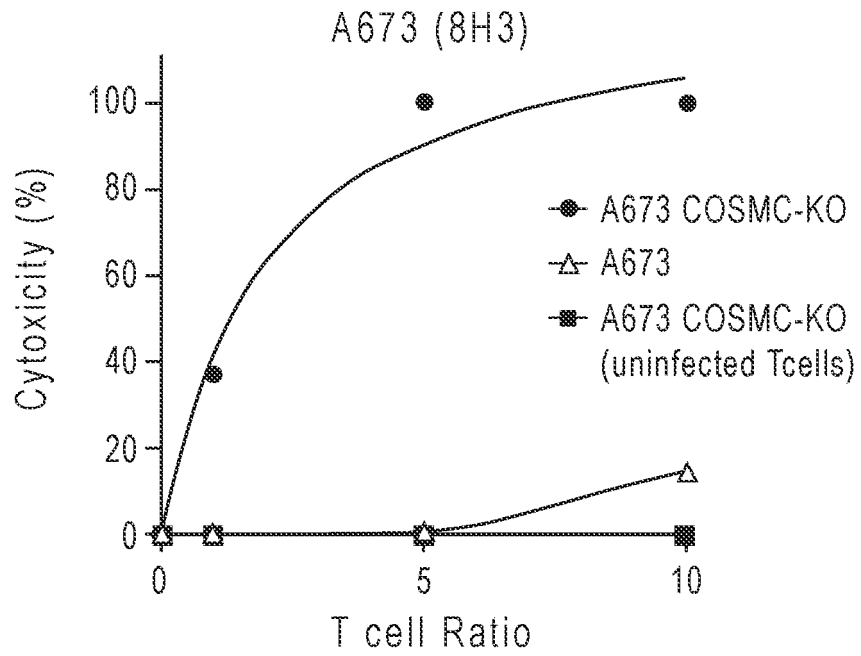


FIG. 7A

KT50 (Time to kill 50% of cells)

Target Cell	T Cell Ratio	8H3 (KT50)	T Cells (KT50)
A673 COSMC-KO	5:1	5.73 hrs	N/A
	10:1	4.98 hrs	N/A

FIG. 7B

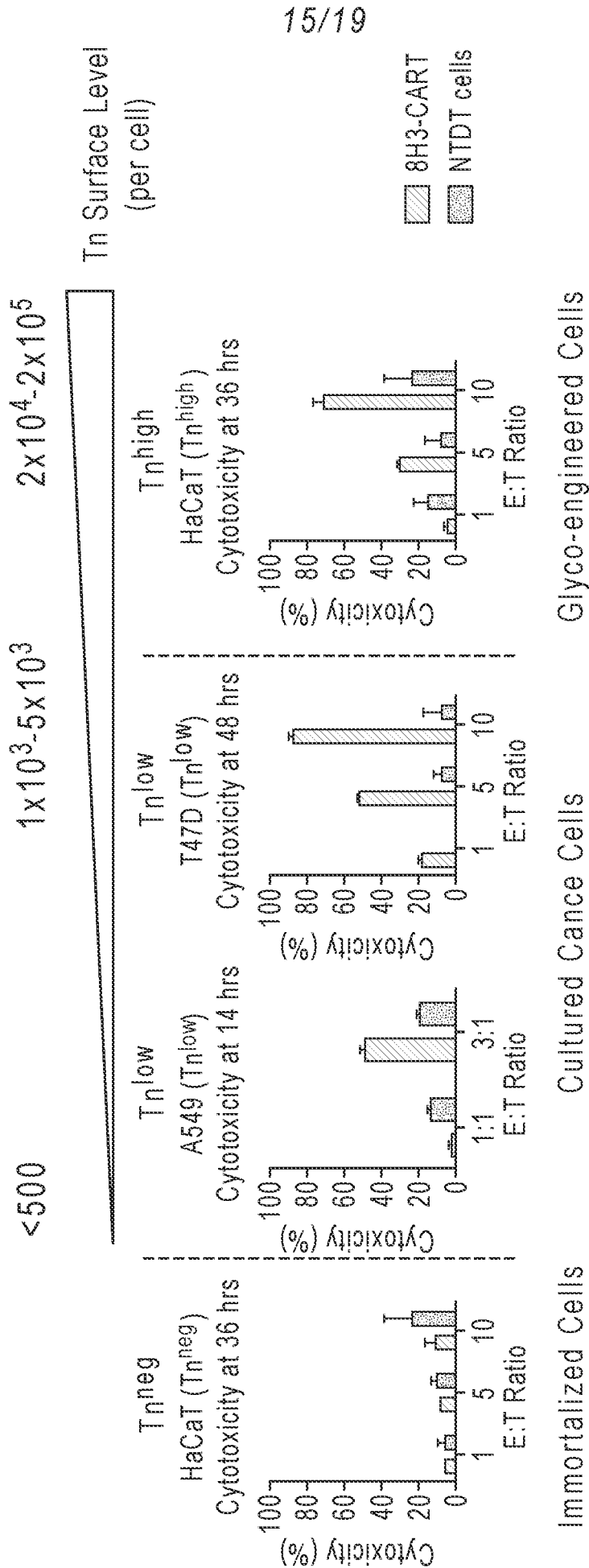
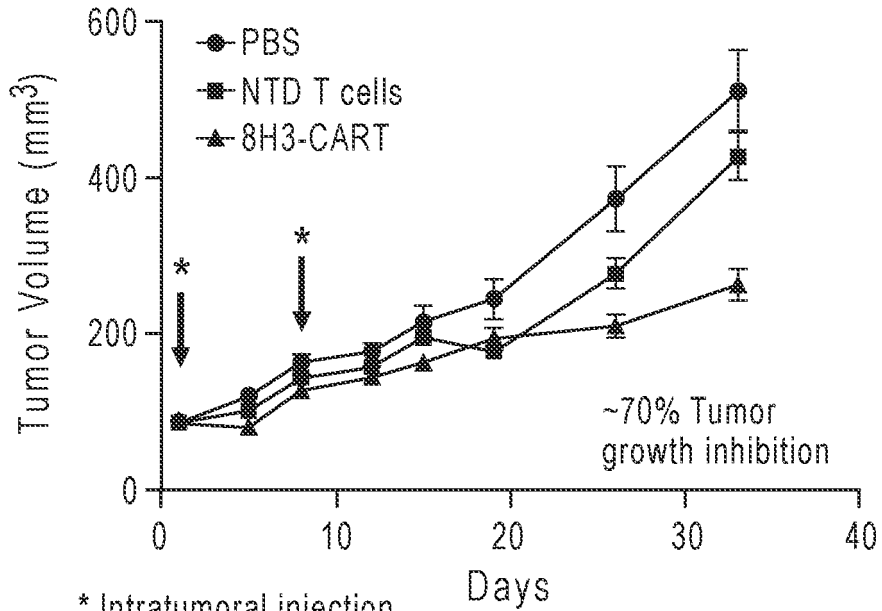


FIG. 7C

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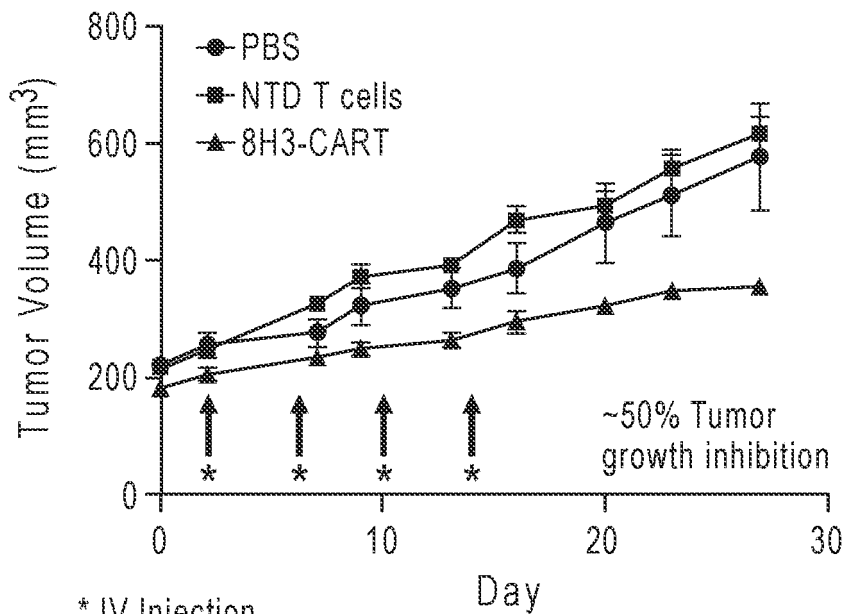
Tumor Growth Curve of A549 CDx (2000 Tn surface binding sites)



* Intratumoral injection
Error bars=SEM
N=5

FIG. 8A

Tumor Growth Curve of Lung PDx (~6000 Tn surface binding sites)



* IV Injection
Error bars=SEM
N=5

FIG. 8B

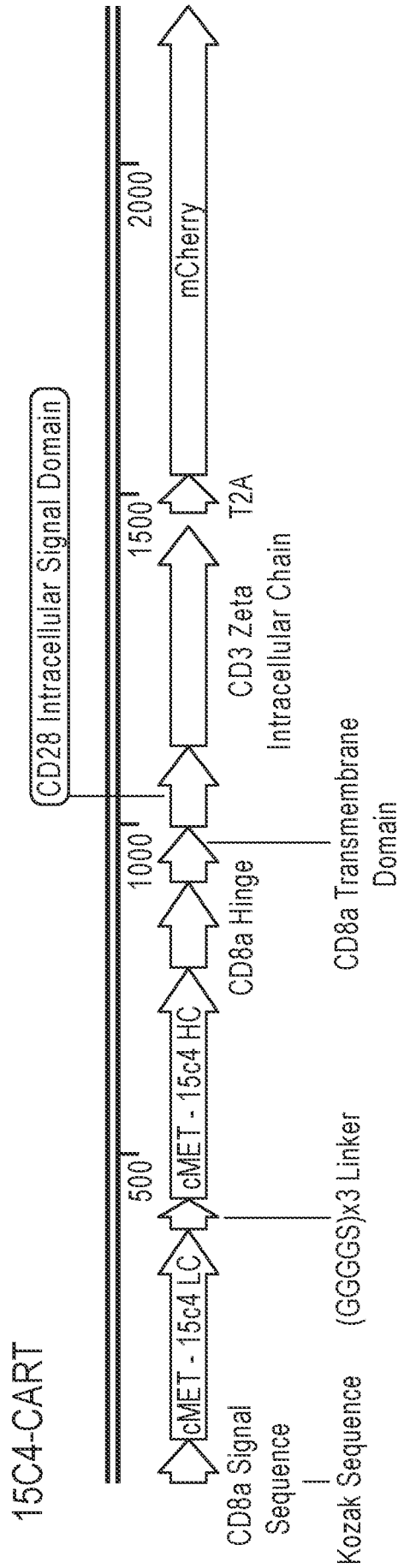


FIG. 9A

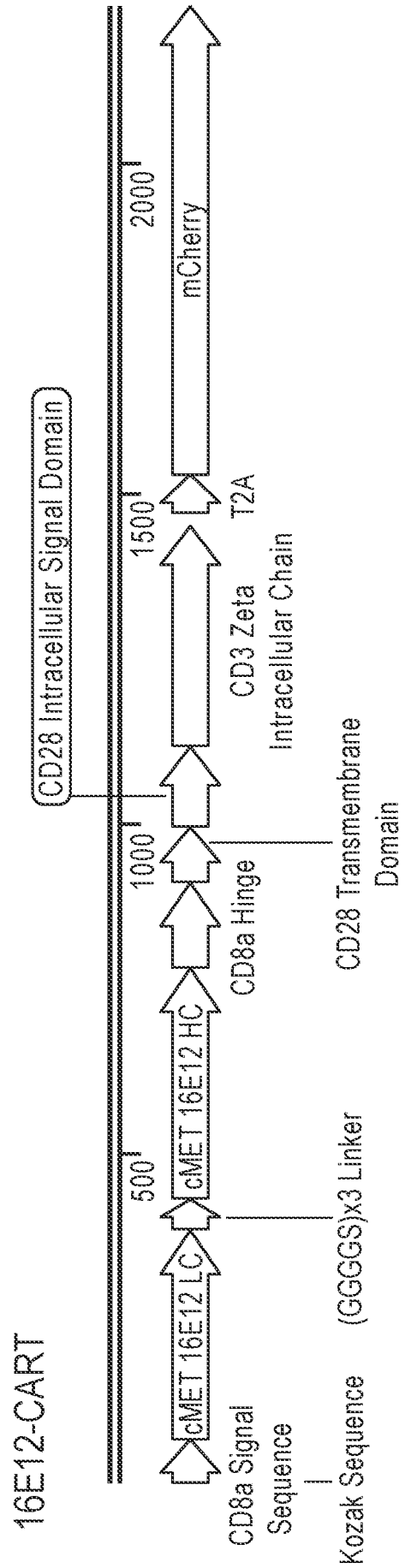


FIG. 9B

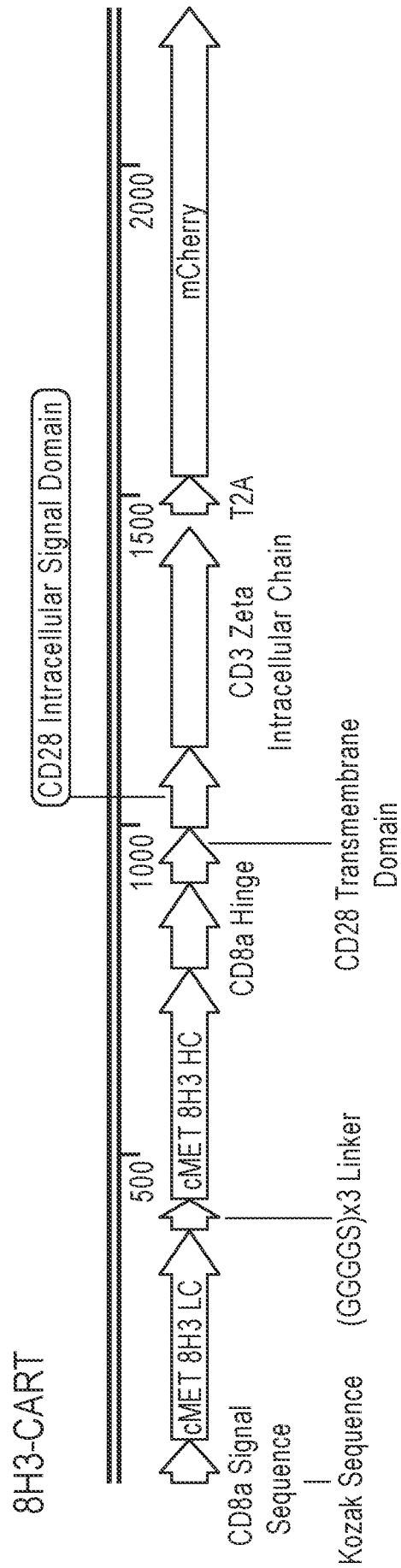


FIG. 9C

Cytotoxicity of hu8H3-CART on A673 cells (2:1 ratio)

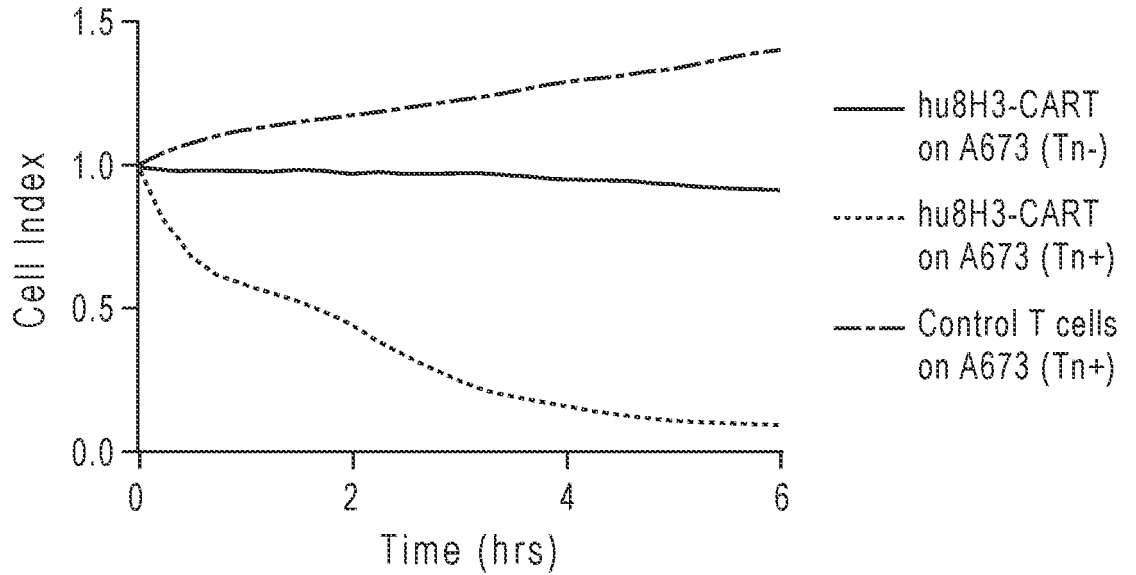


FIG. 10