



US012383631B2

(12) **United States Patent**  
**Pattabiraman et al.**

(10) **Patent No.:** US 12,383,631 B2  
(45) **Date of Patent:** \*Aug. 12, 2025

(54) **IMMUNE ANTIGEN SPECIFIC IL-18  
IMMUNOCYTOKINES AND USES THEREOF**

(71) Applicant: **Bright Peak Therapeutics AG**, Basel (CH)

(72) Inventors: **Vijaya Raghavan Pattabiraman**, Volketswil (CH); **Bertolt Kreft**, Kleinmachow (DE); **Jean-Philippe Carralot**, Saint-Louis (FR); **Arnaud Goepfert**, Sierentz (FR); **Kea Martin**, Istein (DE); **Andrew Chi**, San Diego, CA (US)

(73) Assignee: **Bright Peak Therapeutics AG**, Basel (CH)

(\*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 150 days.

This patent is subject to a terminal disclaimer.

(21) Appl. No.: **18/113,413**

(22) Filed: **Feb. 23, 2023**

(65) **Prior Publication Data**

US 2023/0355795 A1 Nov. 9, 2023

**Related U.S. Application Data**

(60) Provisional application No. 63/479,529, filed on Jan. 11, 2023, provisional application No. 63/313,222, filed on Feb. 23, 2022.

(51) **Int. Cl.**  
*A61K 47/68* (2017.01)

(52) **U.S. Cl.**  
*CPC .....* *A61K 47/6849* (2017.08); *A61K 47/6813* (2017.08)

(58) **Field of Classification Search**  
*CPC .....* *A61K 47/6849; A61K 47/6813*  
See application file for complete search history.

(56) **References Cited**

**U.S. PATENT DOCUMENTS**

7,253,260 B2	8/2007	Janson et al.
7,311,902 B2	12/2007	Bam et al.
7,442,526 B2	10/2008	Johanson et al.
7,524,488 B2	4/2009	Dinarello et al.
7,595,039 B2	9/2009	Dede et al.
7,667,076 B2	2/2010	Bode et al.
7,736,639 B2 *	6/2010	Bam .....
		A61P 35/02
		514/19.3
7,875,709 B2	1/2011	Dinarello et al.
8,216,805 B2	7/2012	Carter et al.
8,679,471 B2	3/2014	Carroll et al.
9,764,038 B2	9/2017	Dennler et al.
9,862,779 B2	1/2018	Beck et al.
10,081,684 B2	9/2018	Ploegh et al.
10,227,383 B2 *	3/2019	Ito .....
10,266,502 B2	4/2019	Louis van Delft et al.
10,434,180 B2	10/2019	Bregeon et al.
10,676,516 B2	6/2020	Viney et al.

10,815,303 B2	10/2020	Yue et al.
10,864,277 B2	12/2020	Grawunder et al.
11,053,293 B2	7/2021	Krupnick et al.
11,168,111 B2	11/2021	Davis et al.
11,219,672 B2	1/2022	Okamura et al.
11,291,721 B2	4/2022	Loew et al.
11,413,331 B2	8/2022	Deak et al.
11,421,022 B2	8/2022	Fenn et al.
12,173,062 B2	12/2024	Kreft et al.
2002/0169291 A1	11/2002	Dinarello et al.
2005/0008615 A1	1/2005	Bam et al.
2009/0286855 A1	11/2009	Dinarello et al.
2010/0291088 A1	11/2010	Ghayur et al.
2016/0107999 A1	4/2016	Debets et al.
2018/0200338 A1	7/2018	Umana et al.
2019/0023760 A1	1/2019	Bode et al.
2019/0062395 A1	2/2019	Merchant et al.
2019/0070262 A1	3/2019	Ring et al.
2019/0194641 A1	6/2019	Spycher et al.
2019/0204330 A1	7/2019	Kelly et al.
2019/0338032 A1	11/2019	Campbell et al.
2019/0352405 A1	11/2019	Fang et al.
2020/0019165 A1	1/2020	Lewandowski et al.
2020/0190165 A1	6/2020	Yamada et al.
2020/0308242 A1 *	10/2020	Lowe .....
		A61P 35/00

(Continued)

**FOREIGN PATENT DOCUMENTS**

EP	2970486 B1	1/2016
WO	WO-9958572 A1	11/1999

(Continued)

**OTHER PUBLICATIONS**

Kim, SH, Azam T, Yoon DY et al. 2001. Site-specific mutations in the mature form of human IL-18 with enhanced biological activity and decreased neutralization by IL-18 binding protein. Proceedings of the National Academy of Sciences of the US 98:3304\_3309 DOI 10.1073/pnas.051634098. (Year: 2001).\*  
Yamamoto Y, Kato Z, Matsukuma E, Li A, Omoya K, Hashimoto K, Ohnishi H, Kondo N. 2004. Generation of highly stable IL-18 based on a ligand receptor complex structure. Biochemical and Biophysical Research Communications 317:181\_186 DOI 10.1016/j.bbrc.2004.03.024. (Year: 2004).\*

(Continued)

*Primary Examiner* — Daniel E Kolker

*Assistant Examiner* — Julia A Rossi

(74) *Attorney, Agent, or Firm* — WILSON SONSINI GOODRICH & ROSATI

(57) **ABSTRACT**

The present disclosure relates to modified immunocytokine compositions comprising antibodies or antigen binding fragments specific for immune cell antigens and IL-18 polypeptides. Also provided herein are methods of treatment with and methods of manufacture of immunocytokine compositions.

**20 Claims, 39 Drawing Sheets**

**Specification includes a Sequence Listing.**

(56)	References Cited		
U.S. PATENT DOCUMENTS			
2020/0317787 A1	10/2020	Li et al.	WO WO-2024102693 A3 6/2024
2020/0330557 A1	10/2020	Hadden, II et al.	WO WO-2024102587 A3 7/2024
2021/0128743 A1	5/2021	Spycher et al.	WO WO-2024153183 A1 7/2024
2021/0139541 A1	5/2021	Matsuda et al.	WO WO-2024158838 A2 8/2024
2021/0139549 A1	5/2021	Yamada et al.	WO WO-2024158840 A2 8/2024
2021/0252157 A1	8/2021	Pattabiraman et al.	WO WO-2024158838 A3 10/2024
2021/0300972 A1	9/2021	Yamada et al.	WO WO-2024158840 A3 10/2024
2021/0388049 A1	12/2021	Li et al.	
2022/0023395 A1	1/2022	Bishai et al.	
2022/0041713 A1	2/2022	Viney et al.	
2022/0056091 A1	2/2022	Pattabiraman et al.	
2022/0112258 A1	4/2022	Cui et al.	
2022/0133904 A1	5/2022	Schibli et al.	
2023/0071889 A1	3/2023	Poirier et al.	
2023/0181754 A1	6/2023	Pattabiraman et al.	
2023/0201364 A1	6/2023	Pattabiraman et al.	
2023/0201365 A1	6/2023	Kreft et al.	
2023/0250181 A1	8/2023	Pattabiraman et al.	
2023/0357342 A1	11/2023	Pattabiraman et al.	
2024/0116997 A1	4/2024	Pattabiraman et al.	
2024/0132563 A1	4/2024	Pattabiraman et al.	
2024/0158537 A1	5/2024	Pattabiraman et al.	
2024/0417436 A1	12/2024	Pattabiraman et al.	
2024/0424127 A1	12/2024	Pattabiraman et al.	
FOREIGN PATENT DOCUMENTS			
WO WO-02085923 A2	10/2002		Swencki-Underwood, B., Cunningham, M., Heavner, G., Blasie, C., McCarthy, S., Dougherty, T., Brigham-Burke, M., Gunn, G., Goletz, T., & Snyder, L. (2006). Engineering human IL-18 with increased bioactivity and bioavailability. <i>Cytokine</i> . 34. 114-24. 10.1016/j.cyto.2006.04.004 (Year: 2006).*
WO WO-2004091517 A2	10/2004		Saetang, J., Puseenam, A., Roongsawang, N., Voravuthikunchai, S.P., Sangkhathat, S., & Tipmanee, V. Immunologic function and molecular insight of recombinant Interleukin-18. <i>PLoS One</i> 11(8): e0160321. <a href="https://doi.org/10.1371/journal.pone.0160321">https://doi.org/10.1371/journal.pone.0160321</a> (Year: 2016).*
WO WO-2004096238 A1	11/2004		Krumm, B., Meng, X., Xiang, Y. et al. Identification of small molecule inhibitors of Interleukin-18. <i>Sci Rep</i> 7, 483 (2017). <a href="https://doi.org/10.1038/s41598-017-00532-x">https://doi.org/10.1038/s41598-017-00532-x</a> (Year: 2017).*
WO WO-2005003294 A2	1/2005		Zheng, Xiaohu, et al., "The use of supercytokines, immunocytokines, engager cytokines, and other synthetic cytokines in immunotherapy." <i>Cellular &amp; molecular immunology</i> 19.2 (2022): 192-209.
WO WO-2005019415 A2	3/2005		Holder, Patrick G., et al., "Engineering interferons and interleukins for cancer immunotherapy." <i>Advanced drug delivery reviews</i> 182 (2022): 114112.
WO WO-2006069246 A2	6/2006		Matsuda, Yutaka, et al., "Chemical Site-Specific Conjugation Platform to Improve the Pharmacokinetics and Therapeutic Index of Antibody-Drug Conjugates." <i>Molecular Pharmaceutics</i> 18.11 (2021): 4058-4066.
WO WO-2007079130 A2	7/2007		Biopharma Peg, "PEGs Conjugate a Versatile Approach of Anti-body Drug," retrieved from <a href="https://www.biochempeg.com/article/85.html">https://www.biochempeg.com/article/85.html</a> , Dec. 12, 2019.
WO WO-2014036492 A1	3/2014		Kang, Min Sun, et al., "Recent developments in chemical conjugation strategies targeting native amino acids in proteins and their applications in antibody-drug conjugates." <i>Chemical Science</i> 12.41 (2021): 13613-13647.
WO WO-2015054658 A1	4/2015		Carralot, Jean, et al. "Using Site-Specific Chemical Conjugation to Generate Superior Half-Life Extended or PD1-Targeted Formats of a Potent IL-18 Variant Resistant to IL-18 Binding Protein," <i>Journal for ImmunoTherapy of Cancer</i> . vol. 10. British Med Assoc House, Tavistock Square, London WC1H 9Jr, England, 2022.
WO WO-2016115168 A1	7/2016		Creative Biolabs: "Anti-PD-L1 Antibody-Cytokine Fusion Protein, IgG (84G09)-IL18 CAT#:ACFP-CQ3407", May 10, 2023 (May 10, 2023), XP093045580, Retrieved from the Internet: URL: <a href="https://www.creativebiolabs.net/anti-pd-l1-immunocytokine-igg-84g09-1118-99433.html">https://www.creativebiolabs.net/anti-pd-l1-immunocytokine-igg-84g09-1118-99433.html</a> .
WO WO-2016115168 A1	7/2016		Kim, Soo-Hyun M., et al. "Site-specific mutations in the mature form of human IL-18 with enhanced biological activity and decreased neutralization by IL-18 binding protein." <i>Proceedings of the National Academy of Sciences</i> 98.6 (2001): 3304-3309.
WO WO-2017201352 A1	11/2017		International Search Report and Written Opinion for PCT Application PCT/IB2023/051688, mailed Jun. 15, 2023, 21 pages. (English).
WO WO-2018199337 A1	11/2018		International Search Report and Written Opinion for PCT Application PCT/IB2023/051690, mailed Jun. 20, 2023, 22 pages. (English).
WO WO-2019051015 A1	3/2019		Harmand, Thibault J., et al. "Protein chemical synthesis by $\alpha$ -ketoacid-hydroxylamine ligation," <i>Nature protocols</i> vol. 11, No. 6, May 26, 2016: 1130-1147.
WO WO-2019057772 A1	3/2019		Hein, Christopher D., et al. "Click chemistry, a powerful tool for pharmaceutical sciences," <i>Pharmaceutical Research</i> , vol. 25, No. 10, May 29, 2008: 2216-2230.
WO WO-2019185792 A1	10/2019		Kim, Soo-Hyun, et al., "Structural requirements of six naturally occurring isoforms of the IL-18 binding protein to inhibit IL-18," <i>Proceedings of the National Academy of Sciences</i> , vol. 97, No. 3, Feb. 1, 2000: 1190-1195.
WO WO-2019240287 A1	12/2019		Kim, Soo-Hyun, et al., "Identification of Amino Acid Residues Critical for Biological Activity in Human Interleukin-18," <i>The Journal of Biological Chemistry</i> vol. 277, No. 13, Mar. 29, 2002: 10998-11003.
WO WO-2019240288 A1	12/2019		
WO WO-2020090979 A1	5/2020		
WO WO-2020127369 A1	6/2020		
WO WO-2020163715 A1	8/2020		
WO WO-2020188061 A1	9/2020		
WO WO-2021122866 A1	6/2021		
WO WO-2021133839 A1	7/2021		
WO WO-2022038417 A2	2/2022		
WO WO-2022094473 A1	5/2022		
WO WO-2022172944 A1	8/2022		
WO WO-2022214653 A1	10/2022		
WO WO-2023281479 A1	1/2023		
WO WO-2023281482 A1	1/2023		
WO WO-2023281484 A1	1/2023		
WO WO-2023010021 A1	2/2023		
WO WO-2023013763 A1	2/2023		
WO WO-2023056193 A2	4/2023		
WO WO-2023114829 A1	6/2023		
WO WO-2023118497 A1	6/2023		
WO WO-2023161854 A1	8/2023		
WO WO-2023161856 A1	8/2023		
WO WO-2023225602 A1	11/2023		
WO WO-2024031046 A2	2/2024		
WO WO-2024044780 A1	2/2024		
WO WO-2024051728 A1	3/2024		
WO WO-2024086191 A2	4/2024		
WO WO-2024031046 A3	5/2024		
WO WO-2024086191 A3	5/2024		
WO WO-2024102587 A2	5/2024		
WO WO-2024102693 A2	5/2024		
WO WO-2024051728 A9	6/2024		

(56)

**References Cited****OTHER PUBLICATIONS**

- Krumm, Brian, et al., "Structural basis for antagonism of human interleukin 18 by poxvirus interleukin 18-binding protein," *Proceedings of the National Academy of Sciences* 105.52 (2008): 20711-20715.
- Thirumurugan, Prakasam, et al., "Click chemistry for drug development and diverse chemical-biology applications," *Chemical reviews* 113.7 (2013): 4905-4979.
- Meng, Xiangzhi, et al., "Variola virus IL-18 binding protein interacts with three human IL-18 residues that are part of a binding site for human IL-18 receptor alpha subunit," *Virology* 358.1 (2007): 211-220.
- Swencki-Underwood, Bethany, et al., "Engineering human IL-18 with increased bioactivity and bioavailability," *Cytokine* 34.1-2 (2006): 114-124.
- Tsutsumi, Naotaka, et al., "The structural basis for receptor recognition of human interleukin-18," *Nature communications* 5.1 (2014): 1-13.
- PCT/IB2021/000574 Written Opinion and International Search Report mailed Mar. 15, 2022, (10 Pages).
- Al-Lazikani, Bissan et al. Standard Conformations for the Canonical Structures of Immunoglobulins. *Journal of Molecular Biology* 273(4):927-948 (1997).
- CAS Accession No. 1380723-44-3. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913b1e16ad7c155548c573/> substanceDetails Accessed Jan. 22, 2025. 36 pages.
- CAS Accession No. 1428935-60-7. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913a1016ad7c155548b4a2/> substanceDetails. Accessed Jan. 22, 2025. 32 pages.
- CAS Accession No. 1537032-82-8. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913614116ad7c1555488a31/> substanceDetails. Accessed Jan. 22, 2025. 28 pages.
- CAS Accession No. 1918149-01-5. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913dc216ad7c155548e42f/> substanceDetails. Accessed Jan. 22, 2025. 5 pages.
- CAS Accession No. 2102192-68-5. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913ceb16ad7c155548dc8a/> substanceDetails. Accessed Jan. 22, 2025. 6 pages.
- CAS Accession No. 2118349-31-6. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913e8816ad7c155548ec22/> substanceDetails. Accessed Jan. 22, 2025. 5 pages.
- CAS Accession No. 2216751-26-5. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913f7c16ad7c155548f65d/> substanceDetails. Accessed Jan. 22, 2025. 5 pages.
- CAS Accession No. 2256084-03-2. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/67913bfe16ad7c155548d22a/> substanceDetails. Accessed Jan. 22, 2025. 3 pages.
- CAS Accession No. 857064-38-1. Retrieved from: <https://scifinder-n.cas.org/searchDetail substance/6791409616ad7c1555490514/> substanceDetails. Accessed Jan. 22, 2025. 4 pages.
- Chemical Abstracts Service. CAS Registry: 1415800-37-1. Bromoacetamido-PEG5-azide: pp. 1-9. STN Entry Date Sep. 5, 2014. Retrieved Nov. 12, 2024. Retrieved from: <https://pubchem.ncbi.nlm.nih.gov/compound/77078486>.
- Chothia, Cyrus, and Arthur M. Lesk. Canonical Structures for the Hypervariable Regions of Immunoglobulins. *Journal of Molecular Biology* 196(4):901-917 (1987).
- Co-pending U.S. Appl. No. 18/813,104, inventors Pattabiraman; Vijaya Raghavan et al., filed Aug. 23, 2024.
- Co-pending U.S. Appl. No. 18/813,286, inventors Pattabiraman; Vijaya Raghavan et al., filed Aug. 23, 2024.
- Co-pending U.S. Appl. No. 18/813,358, inventors Kreft; Bertolt et al., filed Aug. 23, 2024.
- Correspondence between C numberings (<https://imgt.org/IMGTScientificChart/Numbering/HuIGHGnber.html>).
- EBI Accession No. AXT 13707. Coding: AXT13707.1. *Bacillus Velezensis PucR Family Transcriptional Regulator*. EMBL-EBI: pp. 1-3. Record created Dec. 2017. Retrieved Nov. 12, 2024. Retrieved from: <https://www.ebi.ac.uk/ena/browser/view/AXT13707>.
- EBI Accession No. AXT 13710. Coding: AXT13710.1. *Bacillus Velezensis Hypothetical Protein*. EMBL-EBI: pp. 1-3. Record created Dec. 2017. Retrieved Nov. 12, 2024. Retrieved from: <https://www.ebi.ac.uk/ena/browser/view/AXT13710>.
- Fujii, Tomohiro et al. AJICAP second generation: improved chemical site-specific conjugation technology for antibody-drug conjugate production. *Bioconjugate Chemistry* 34(4):728-738 (2023).
- Kabat, Elvin A. et al. Sequences of Proteins of Immunological Interest, Fifth Edition. Public Health Service, National Institutes of Health 1:647-669 (1991).
- Liu, Hongyan et al. Fc Engineering for Developing Therapeutic Bispecific Antibodies and Novel Scaffolds. *Frontiers in Immunology* 8(38):1-15 (2017).
- PCT/IB2021/000574 International Preliminary Report on Patentability dated Mar. 2, 2023.
- PCT/IB2021/000574 Invitation to Pay Additional Fees dated Jan. 25, 2022.
- PCT/IB2023/051688 International Preliminary Report on Patentability dated Sep. 6, 2024.
- PCT/IB2023/051690 International Preliminary Report on Patentability dated Sep. 6, 2024.
- Remington's Pharmaceutical Sciences, 17th edition (Mack Publishing Company, Easton, Pa, pp. 1418 (1985).
- Saunders, Kevin O. Conceptual Approaches to Modulating Antibody Effector Functions and Circulation Half-Life. *Frontiers in Immunology* 10:1296, 1-20 (2019).
- U.S. Appl. No. 17/406,847 Office Action dated Feb. 1, 2024.
- U.S. Appl. No. 17/406,847 Office Action dated Nov. 6, 2024.
- Yamada, Kei et al. AJICAP: Affinity Peptide Mediated Regiodivergent Functionalization of Native Antibodies. *Angewandte Chemie* 58(17):5592-5597 (2019).
- Co-pending U.S. Appl. No. 18/813,101, inventors Vijaya; Raghavan Pattabiraman et al., filed Aug. 23, 2024.
- Kim, E.J. et al. Efficient induction of T helper type 1-mediated immune responses in antigen-primed mice by anti-CD3 single-chain Fv/interleukin-18 fusion DNA. *Immunology* 111(1):27-34 (2004).
- PCT/IB2024/058220 International Search Report and Written opinion dated Mar. 21, 2025.
- PCT/IB2024/058220 Invitation to Pay Additional Fees dated Jan. 24, 2025.
- PCT/IB2024/058222 International Search Report and Written Opinion dated Jan. 20, 2025.
- PCT/IB2024/058236 International Search Report and Written Opinion dated Jan. 20, 2025.
- U.S. Appl. No. 17/406,847 Non-Final Office Action dated Apr. 24, 2025.

\* cited by examiner

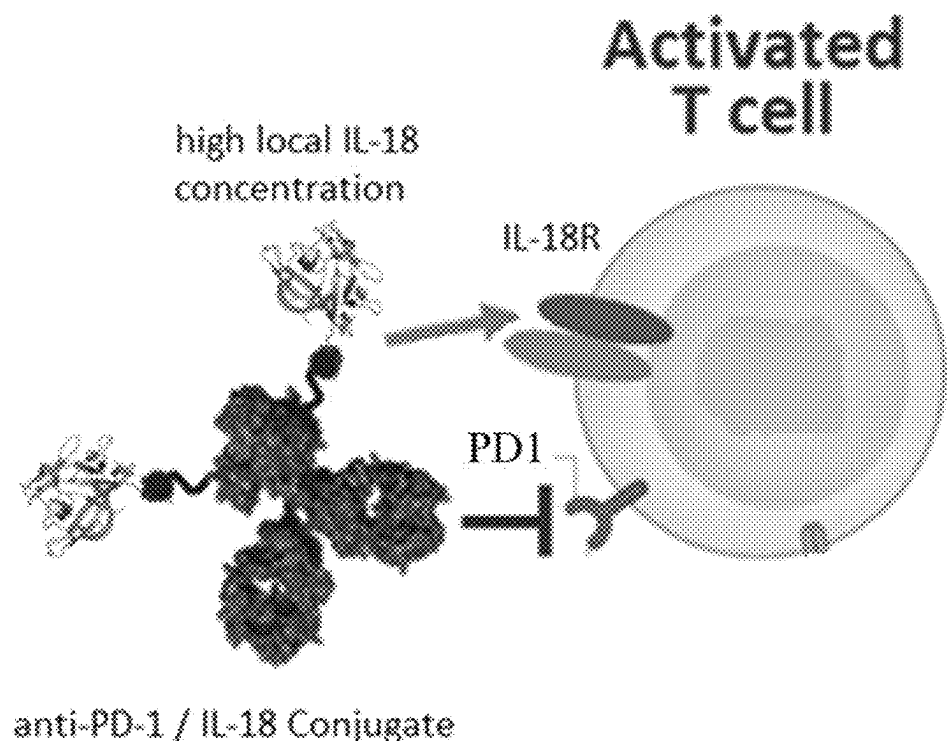


FIGURE 1A

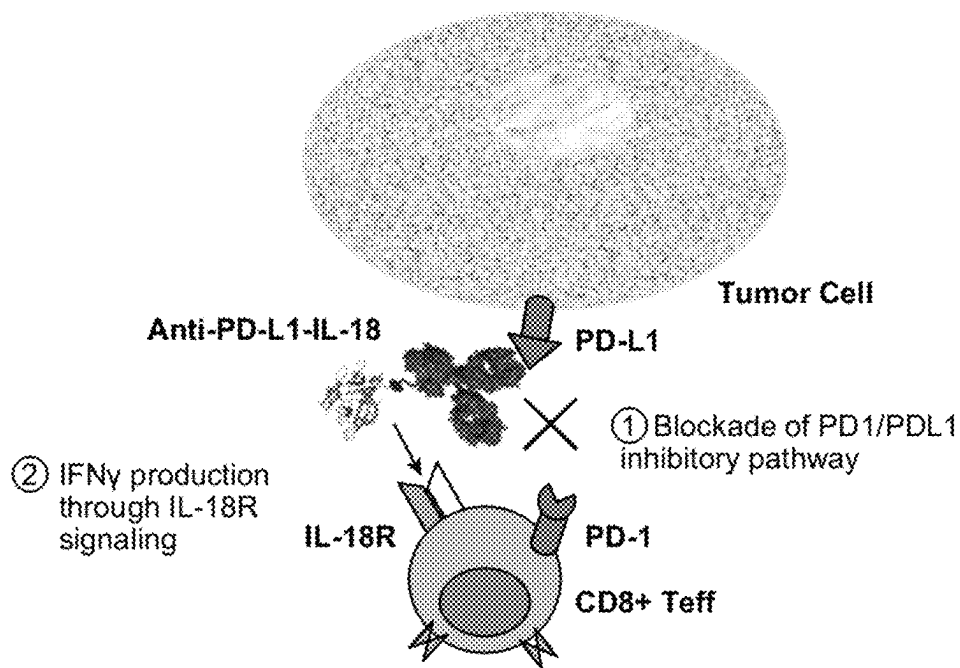
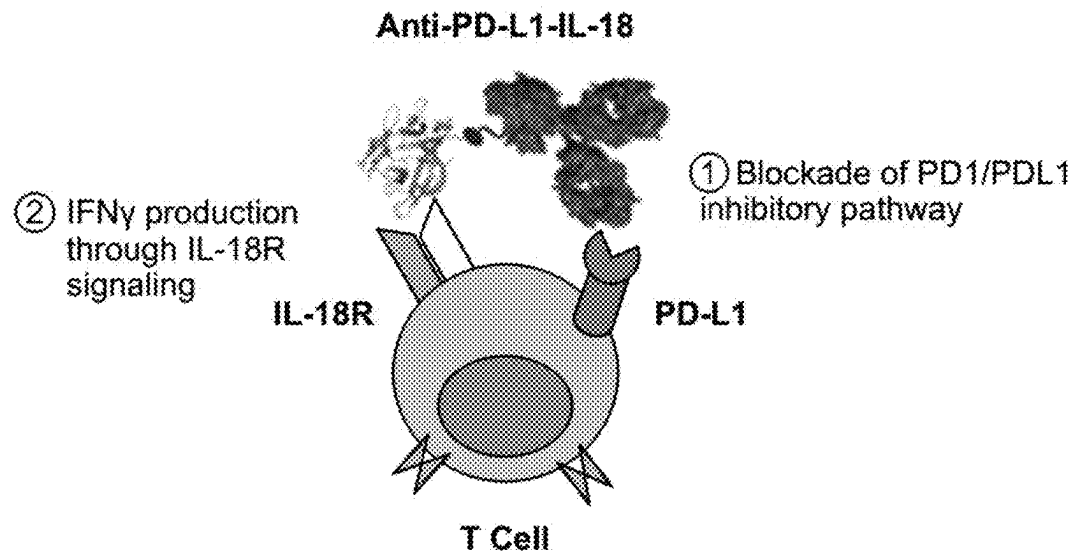
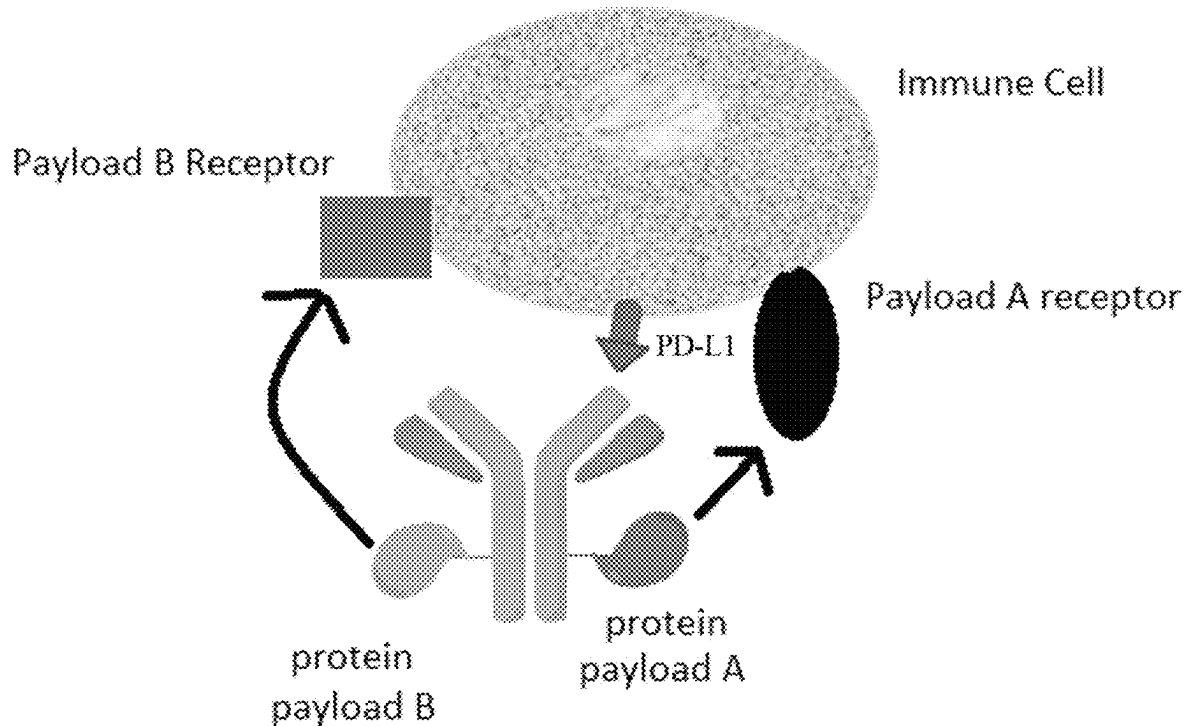


FIGURE 1B

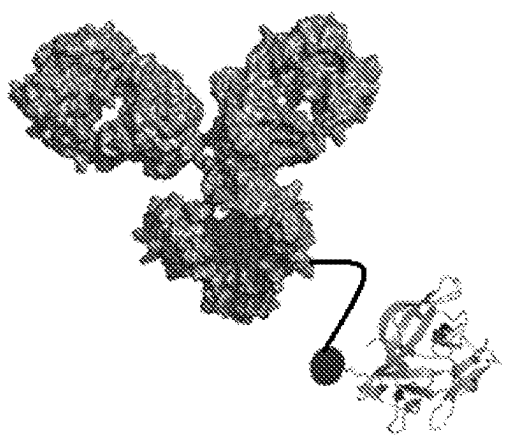


**FIGURE 1C**

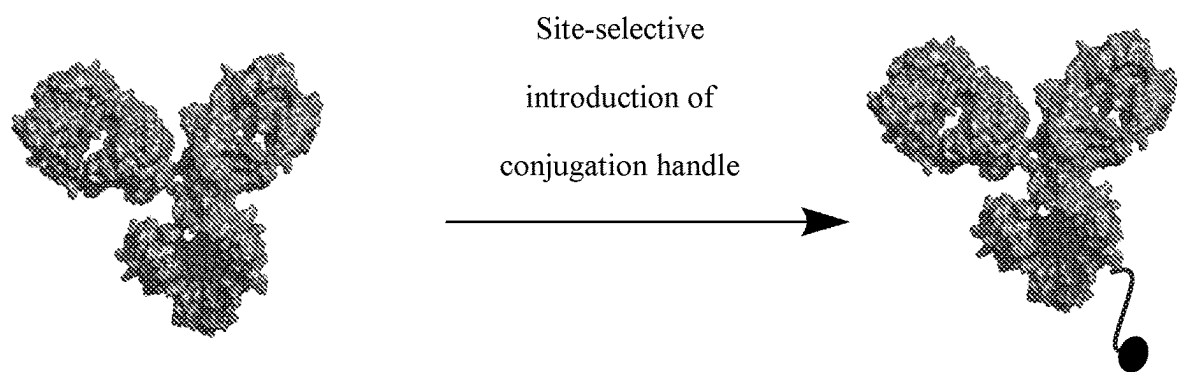
Simultaneous activation of 2 receptors



**FIGURE 1D**



**FIGURE 2**



**FIGURE 3**

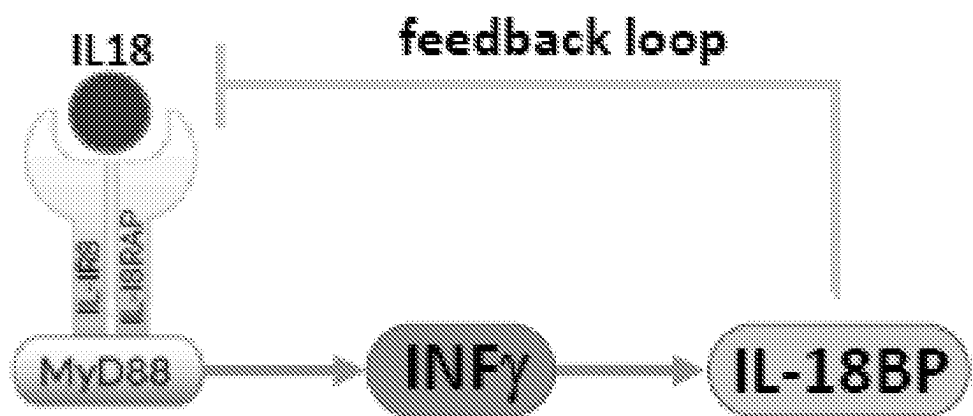


FIGURE 4

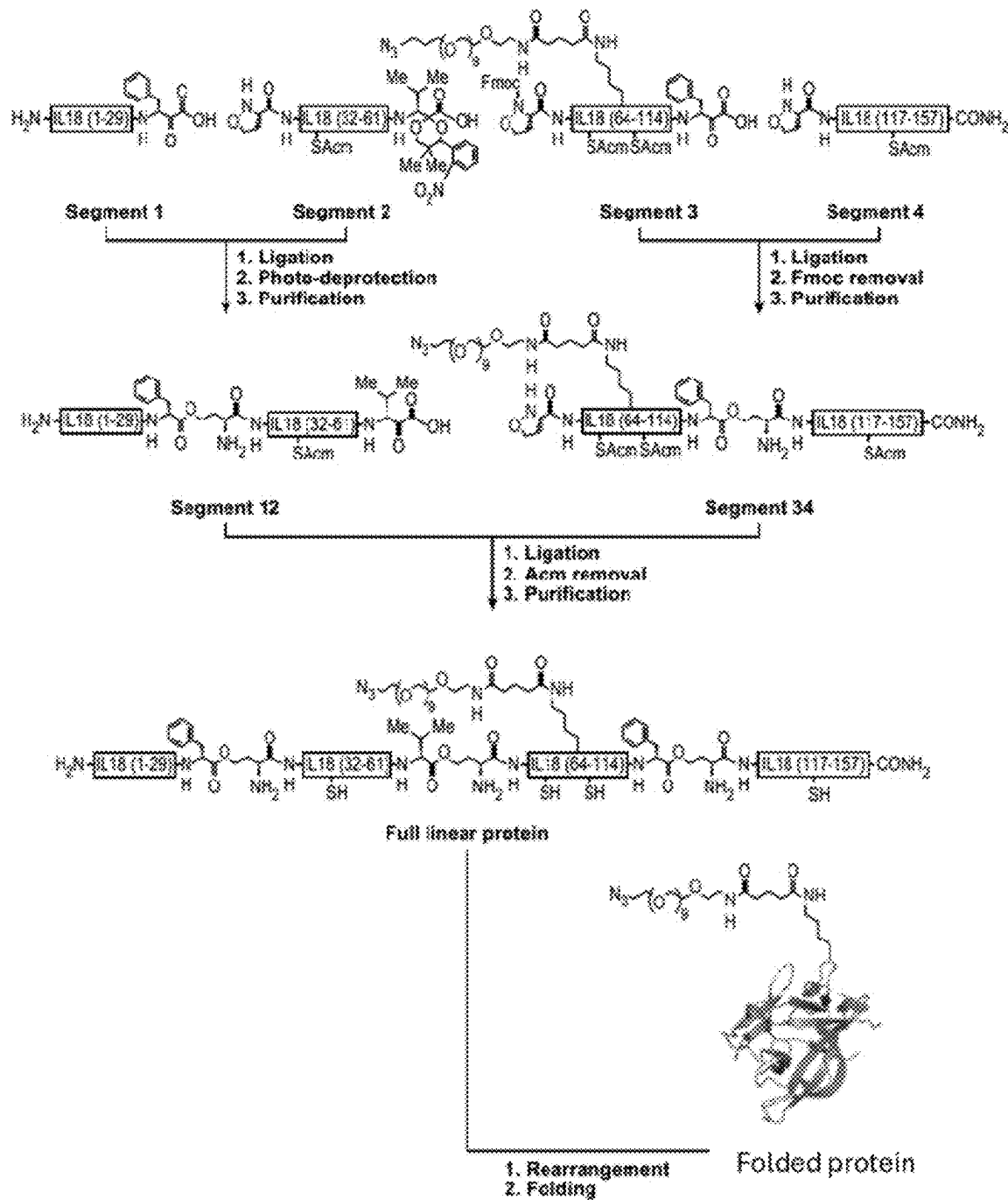
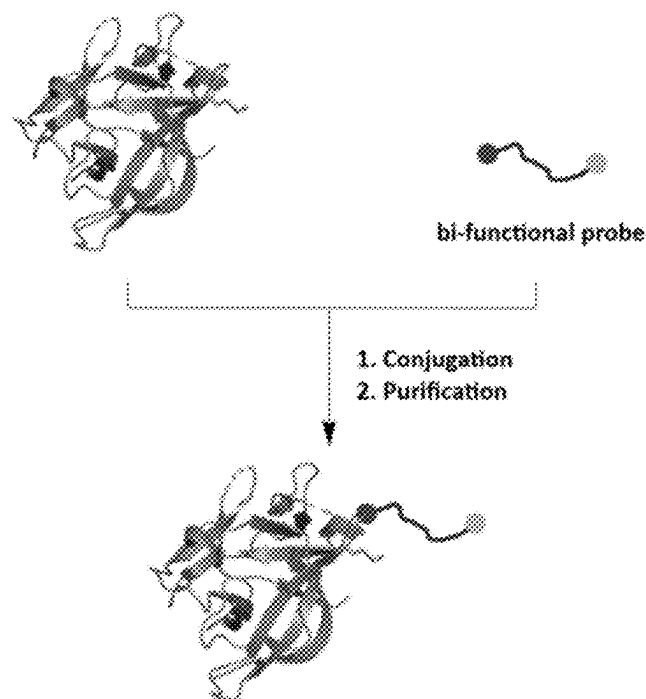
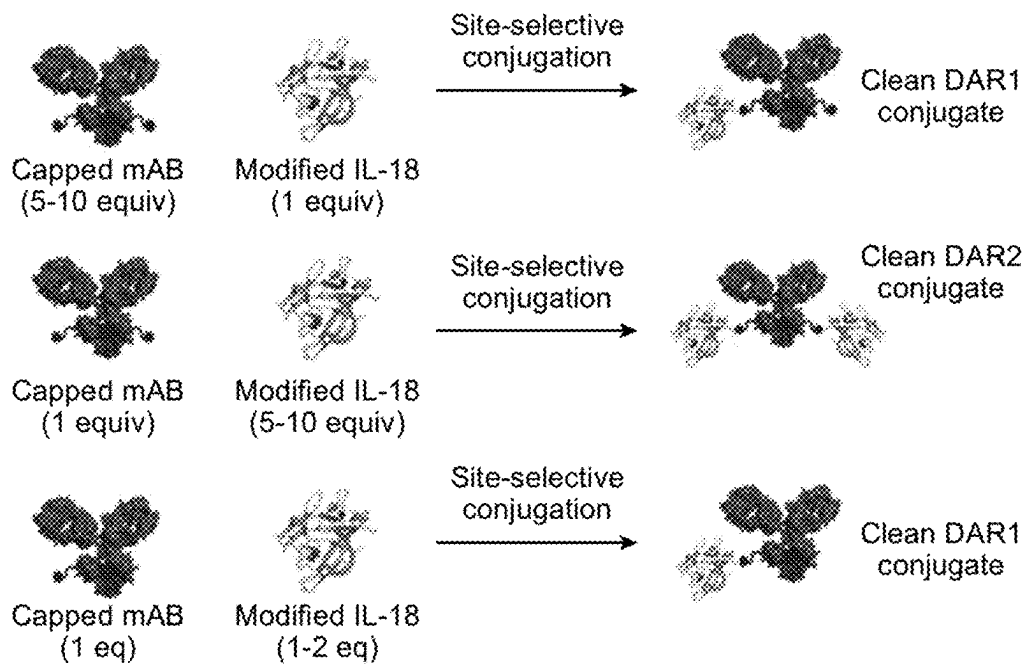


FIGURE 5



**FIGURE 6**



**FIGURE 7**

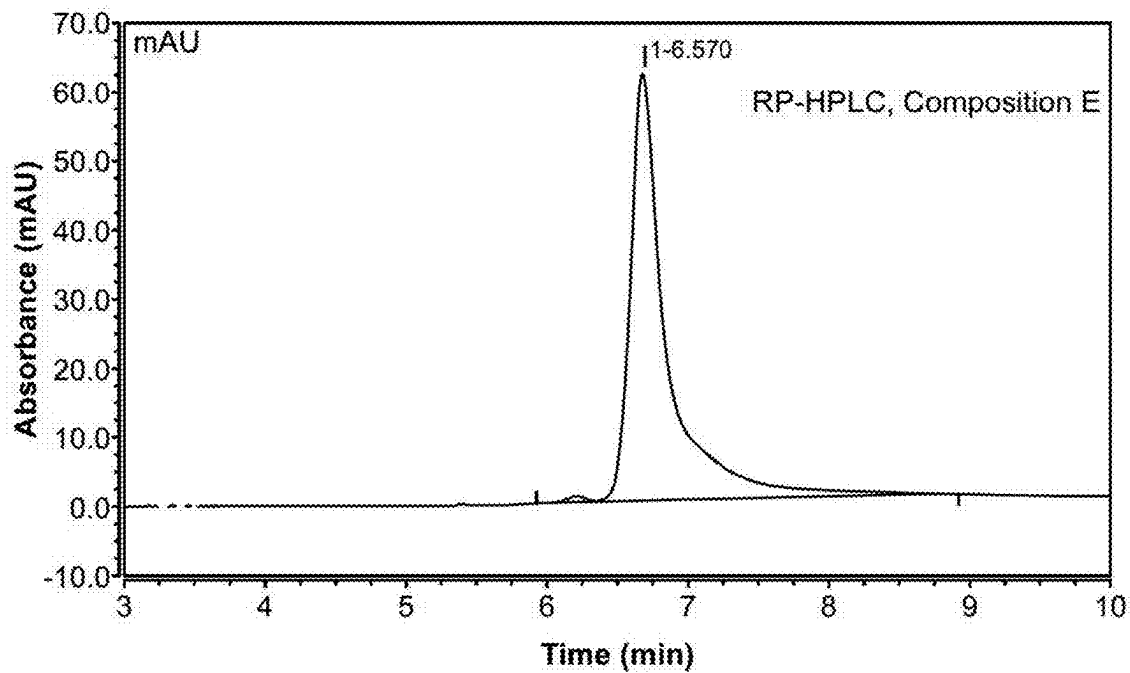


FIGURE 8

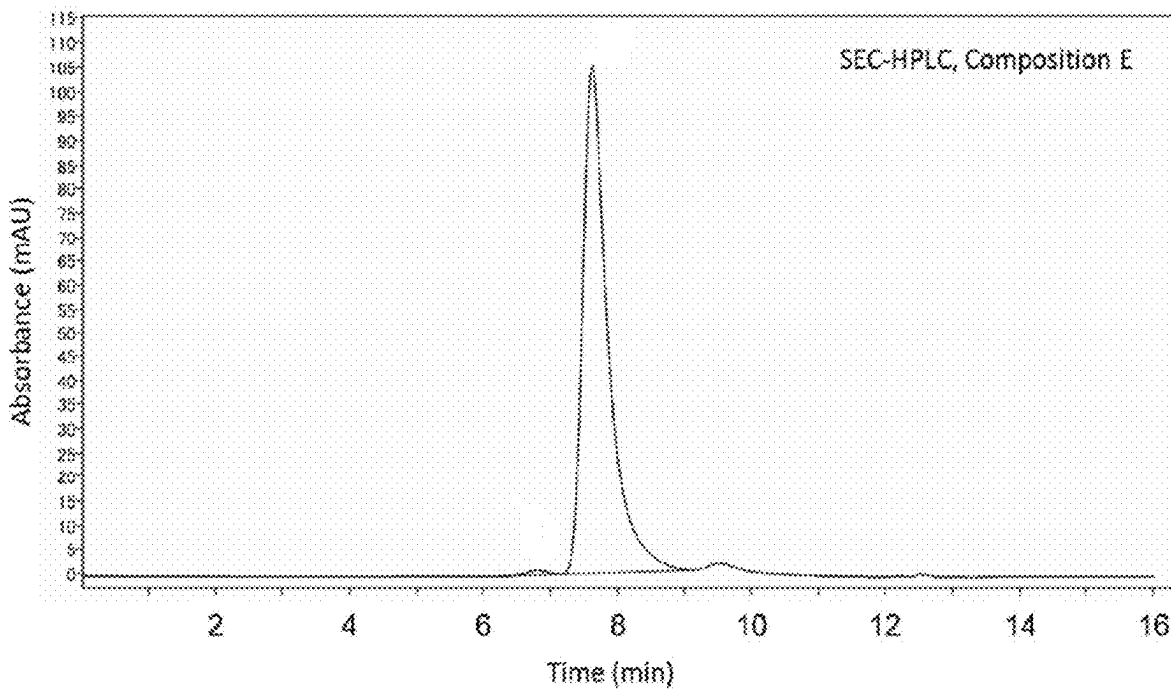


FIGURE 9

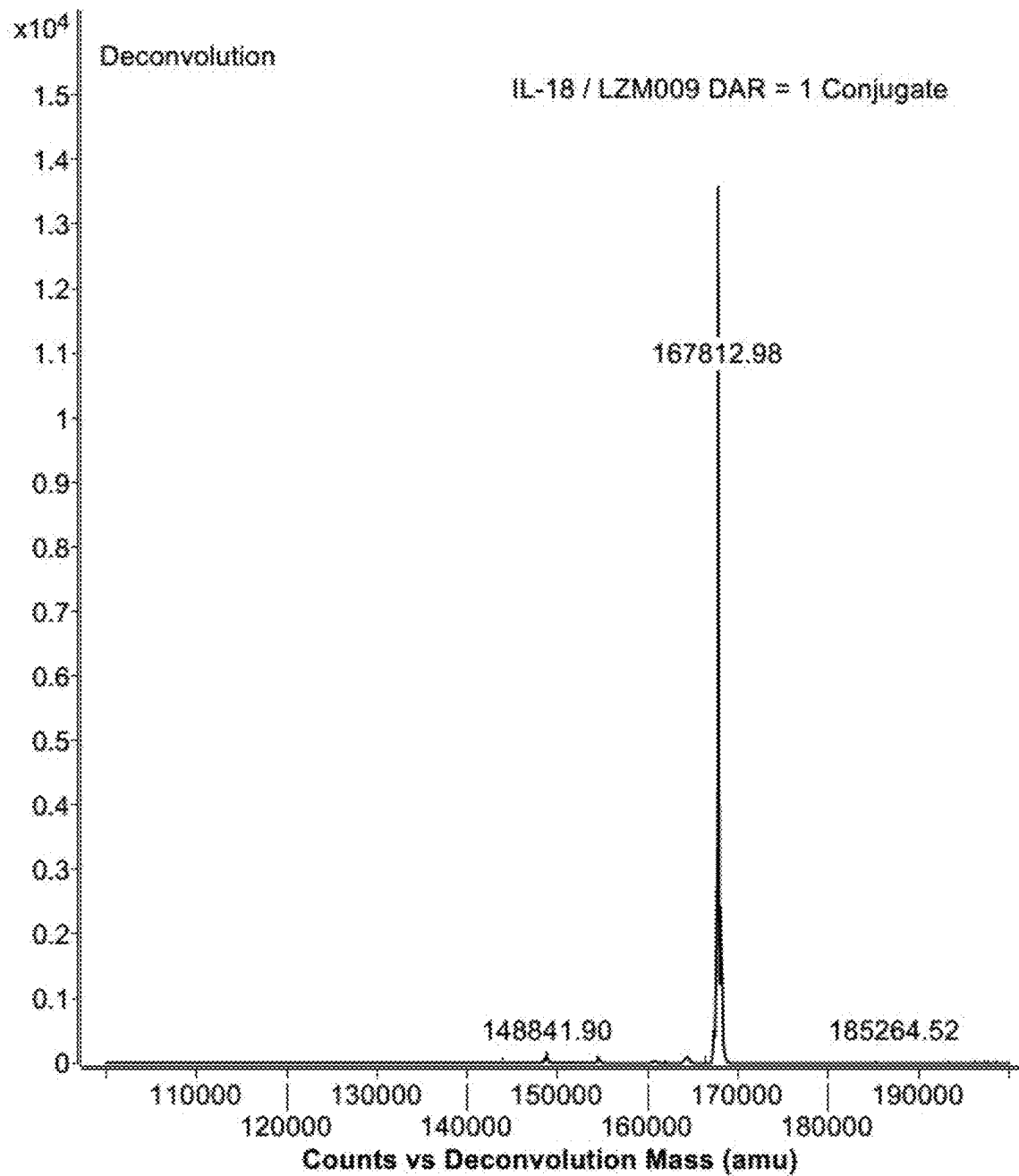


FIGURE 10

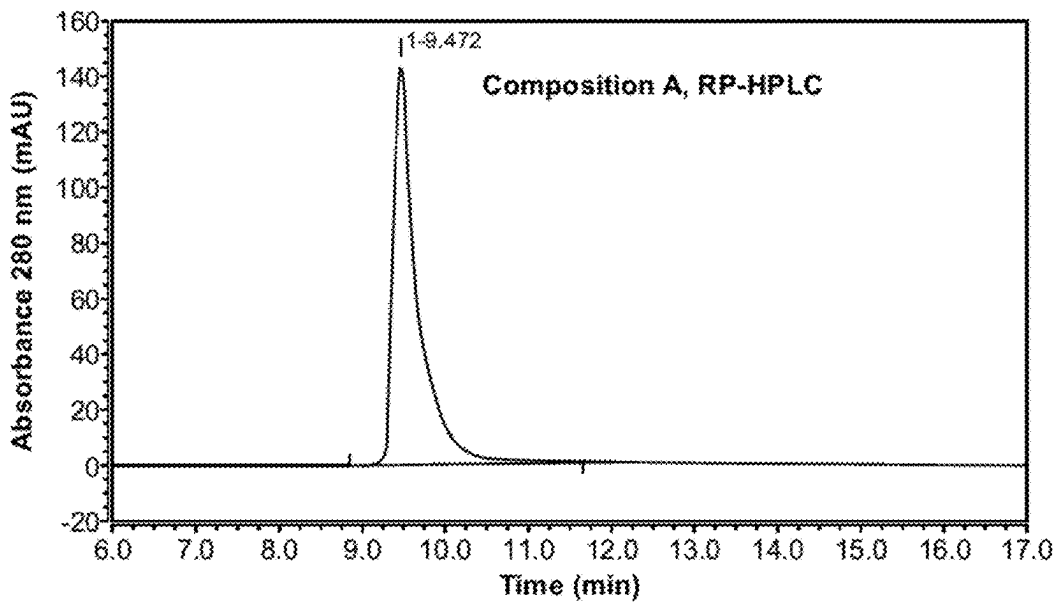


FIGURE 11

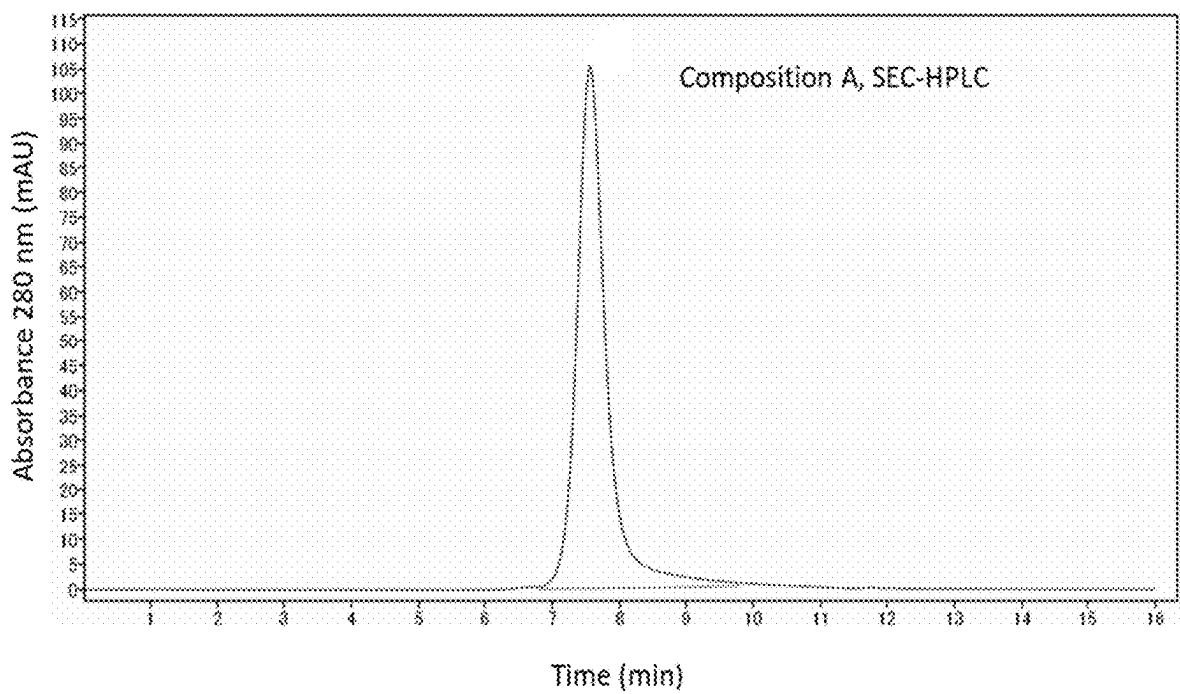


FIGURE 12

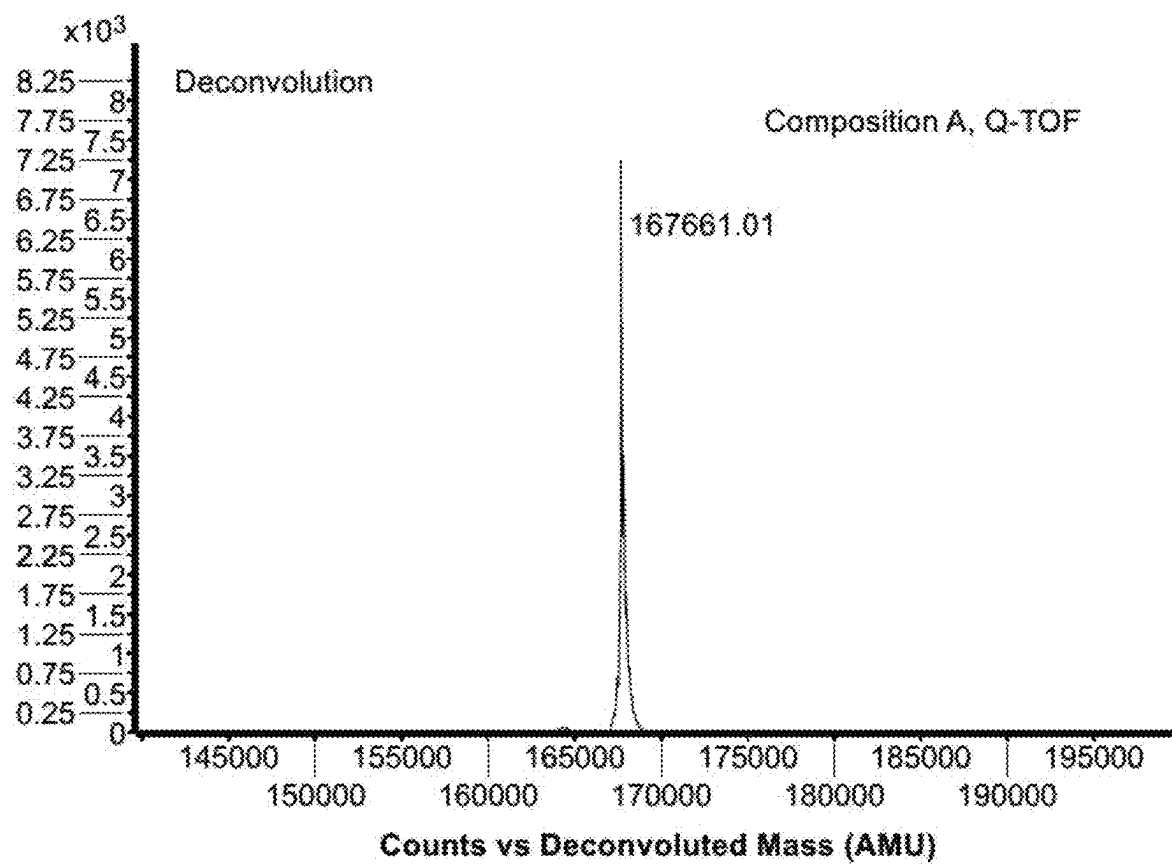


FIGURE 13

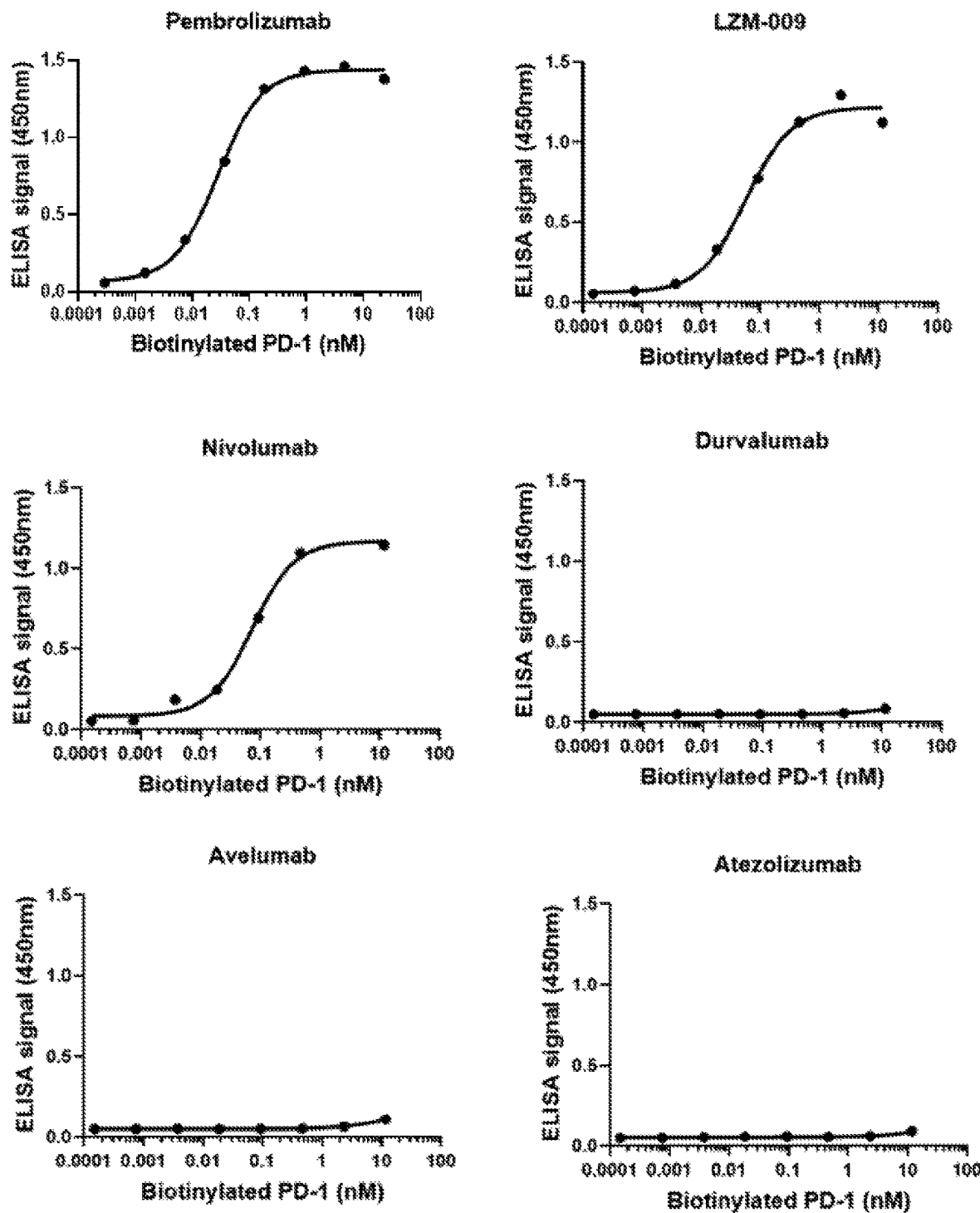


FIGURE 14A

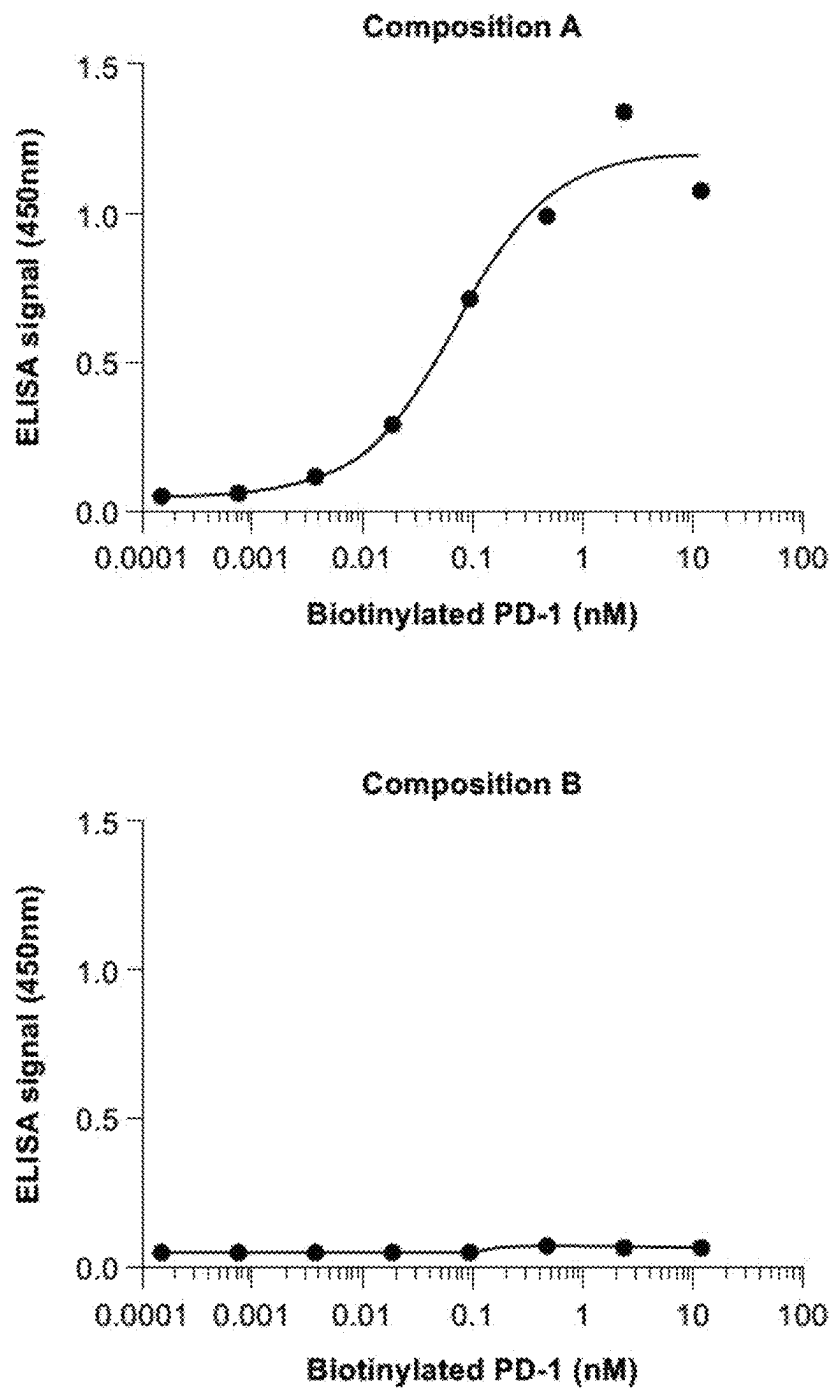


FIGURE 14B

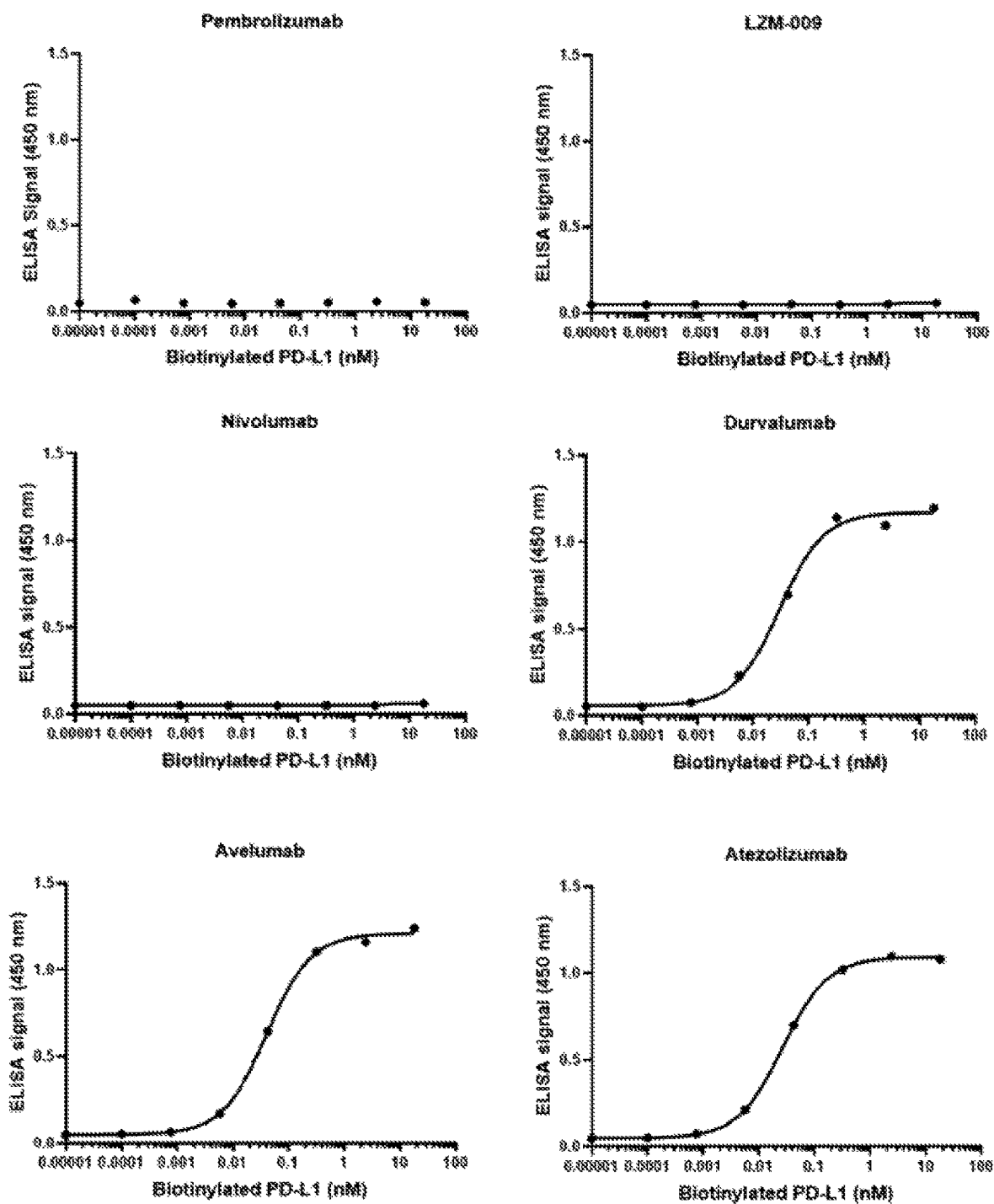
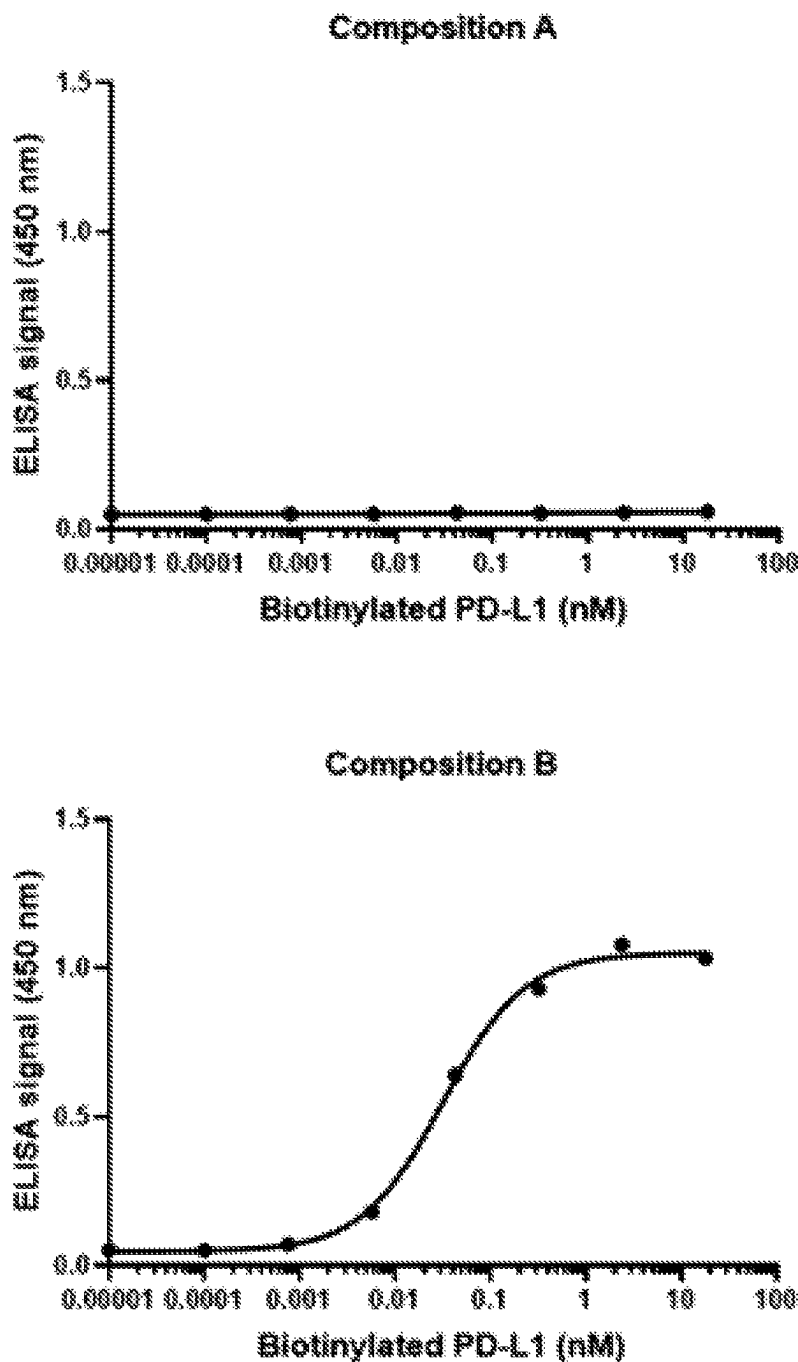


FIGURE 15A

**FIGURE 15B**

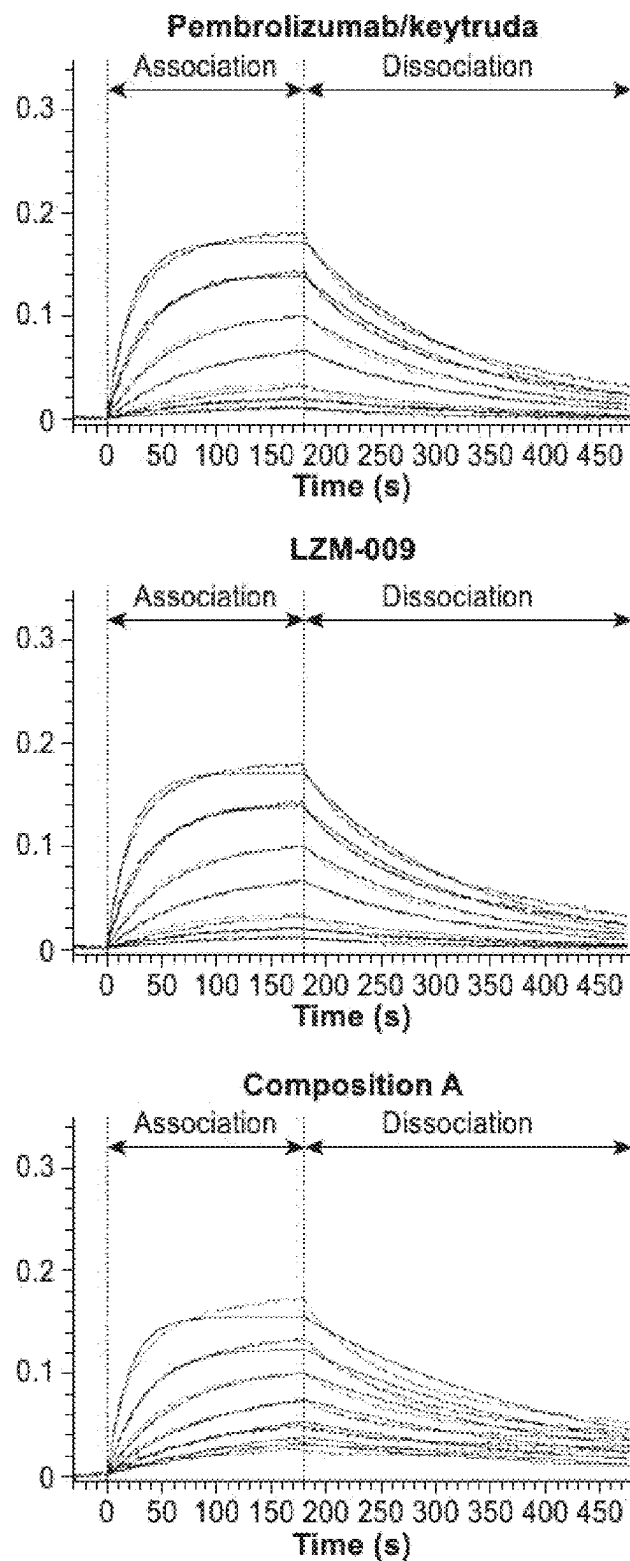


FIGURE 16

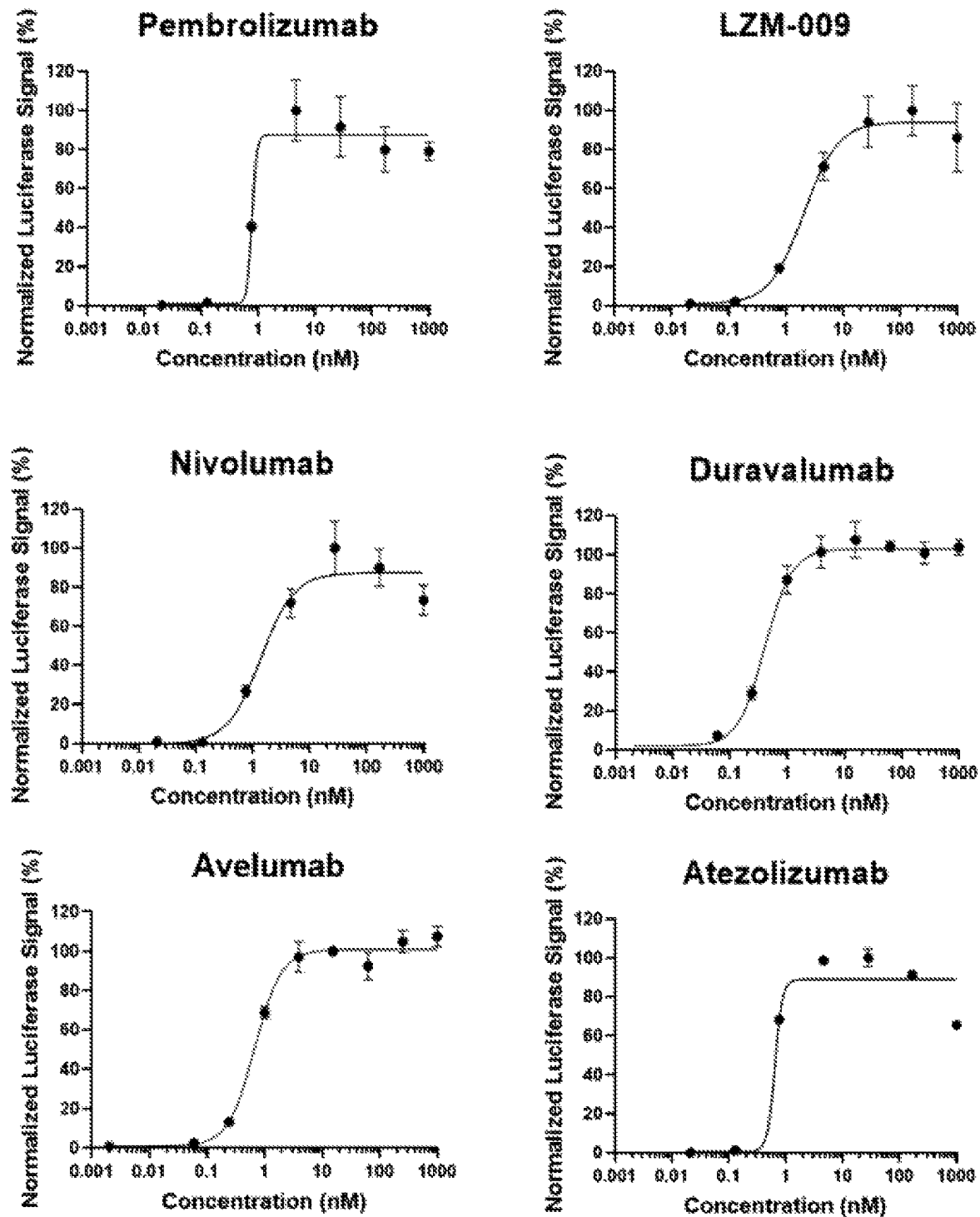


FIGURE 17A

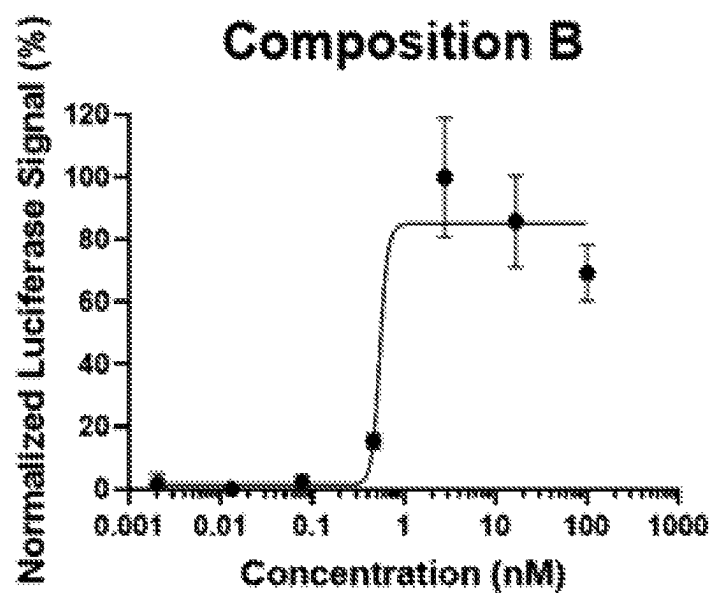
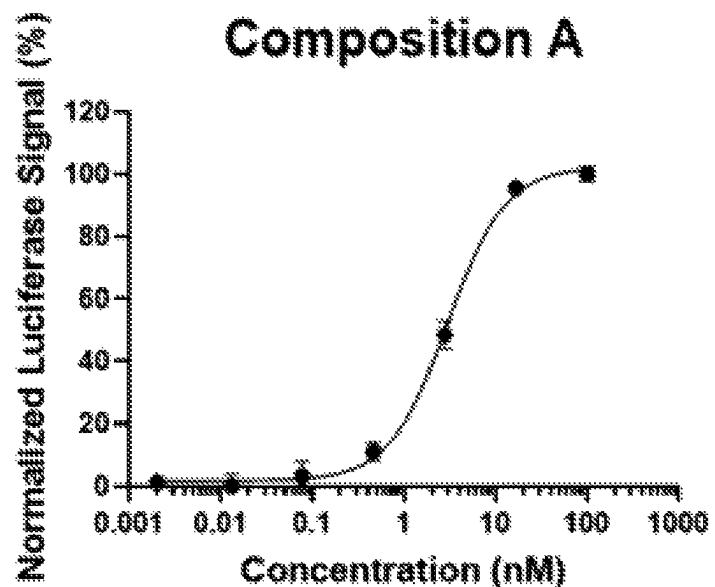


FIGURE 17B

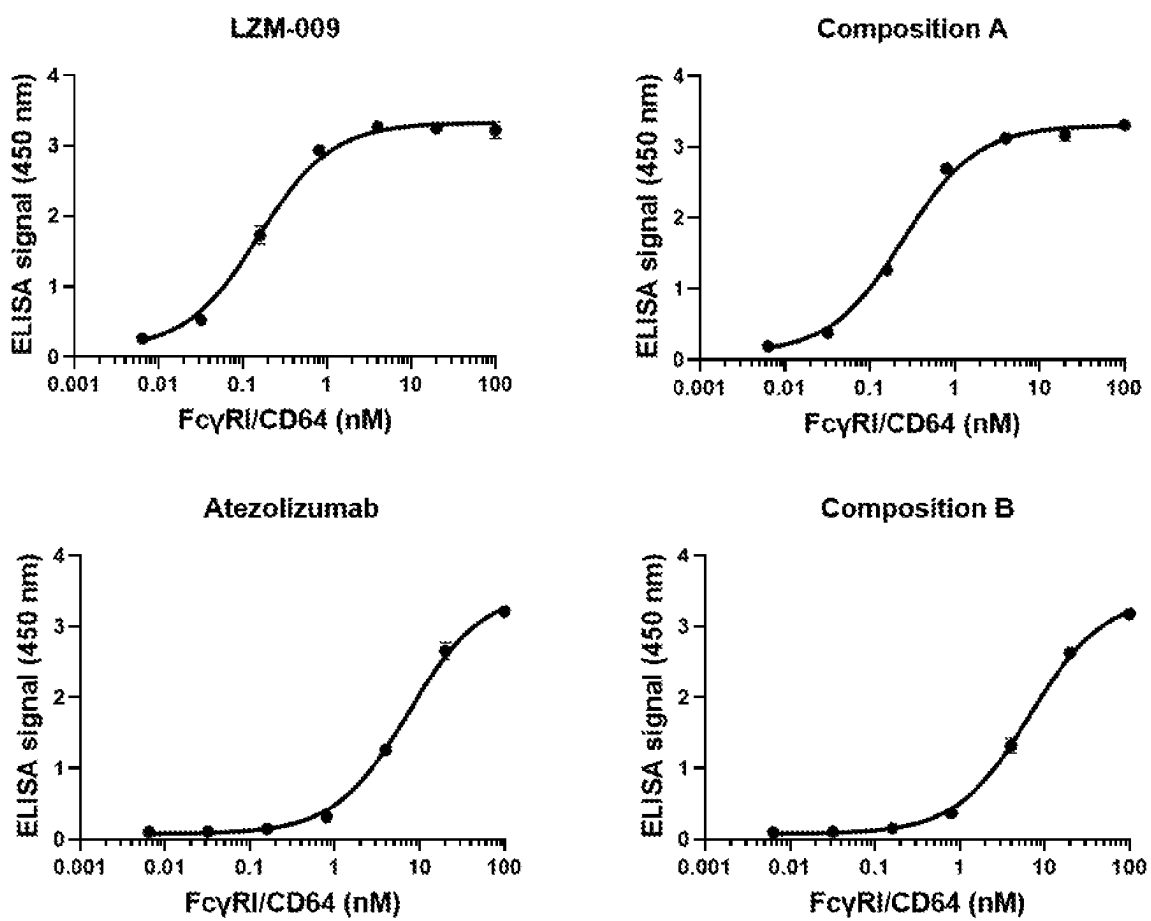


FIGURE 18A

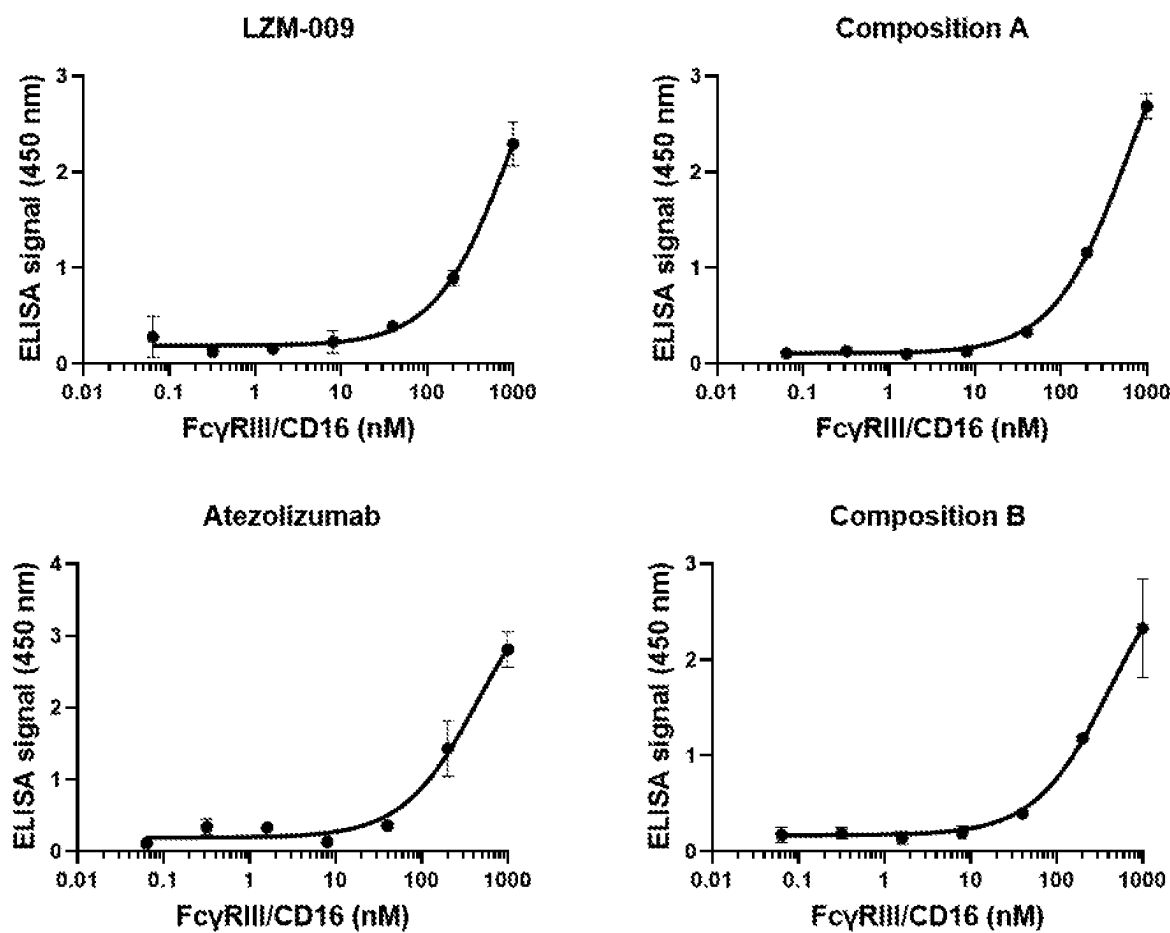


FIGURE 18B

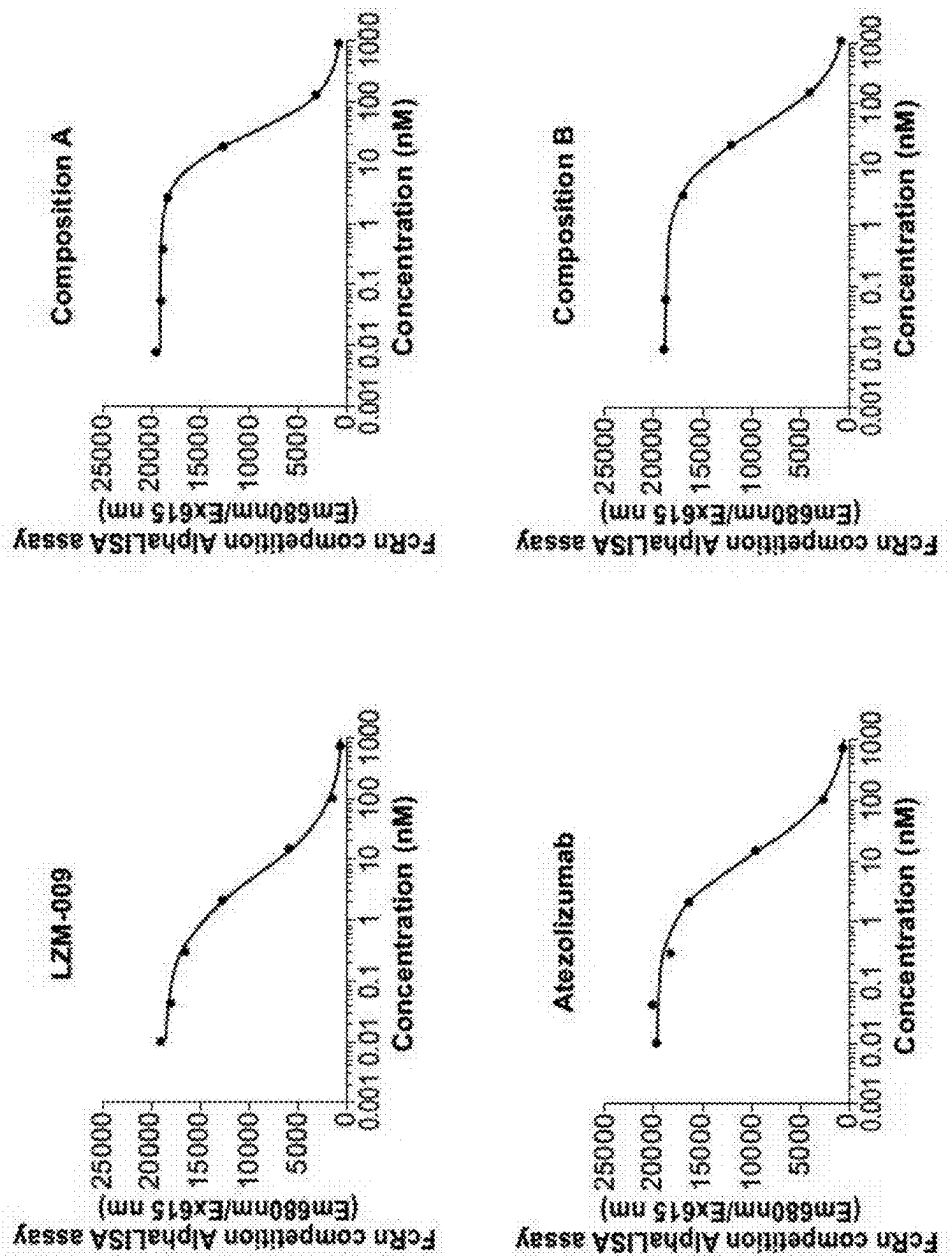
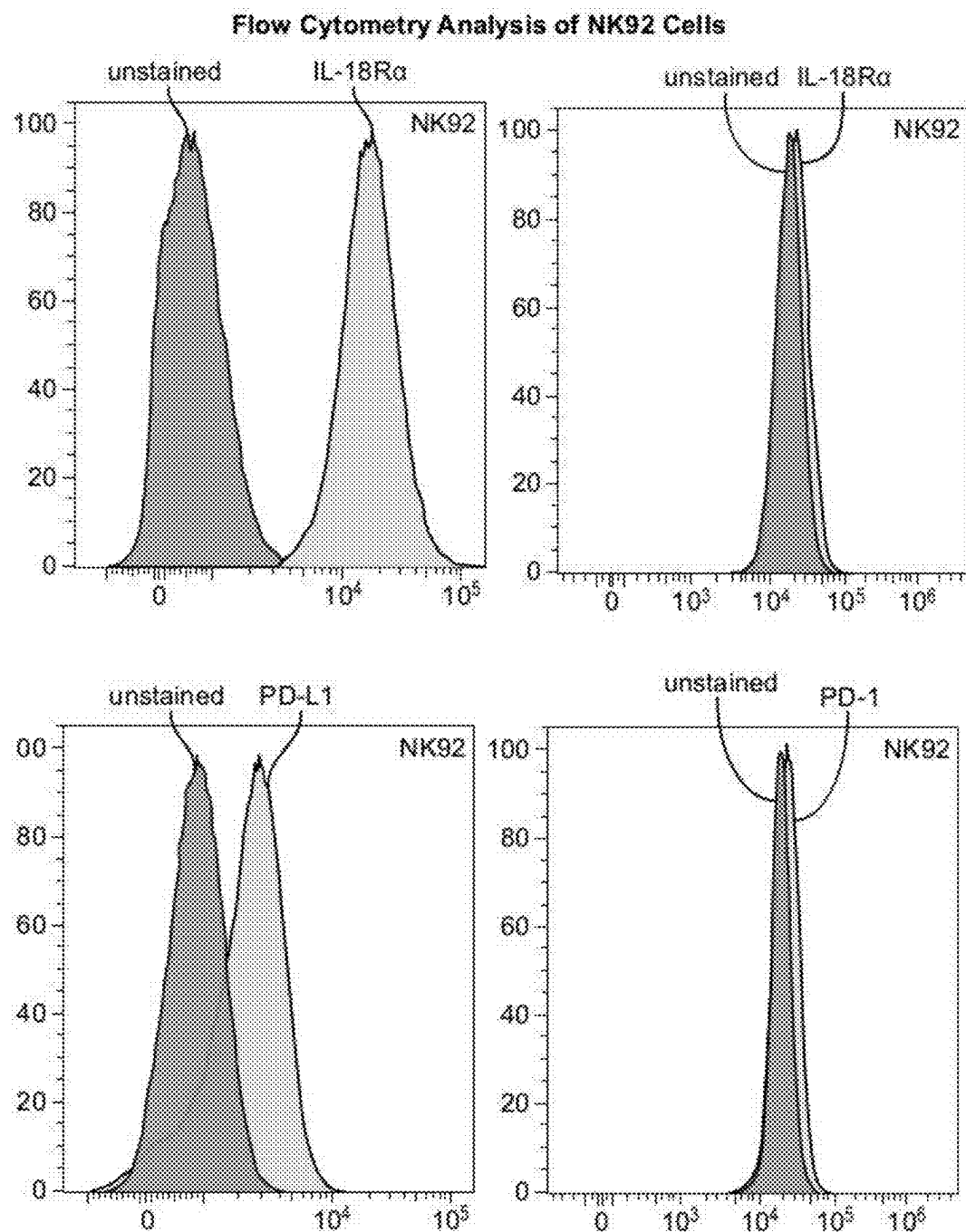


FIGURE 19

**FIGURE 20**

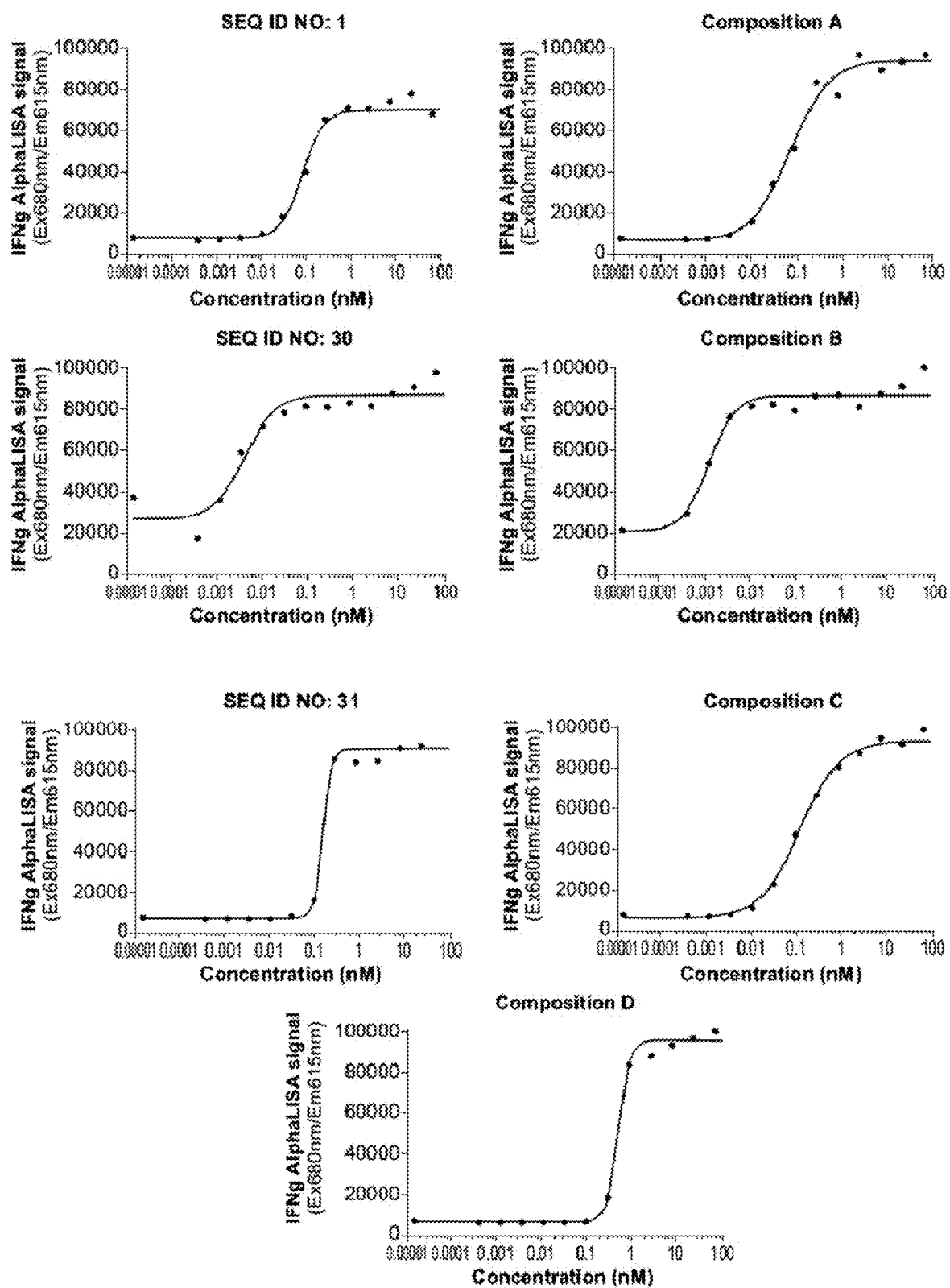
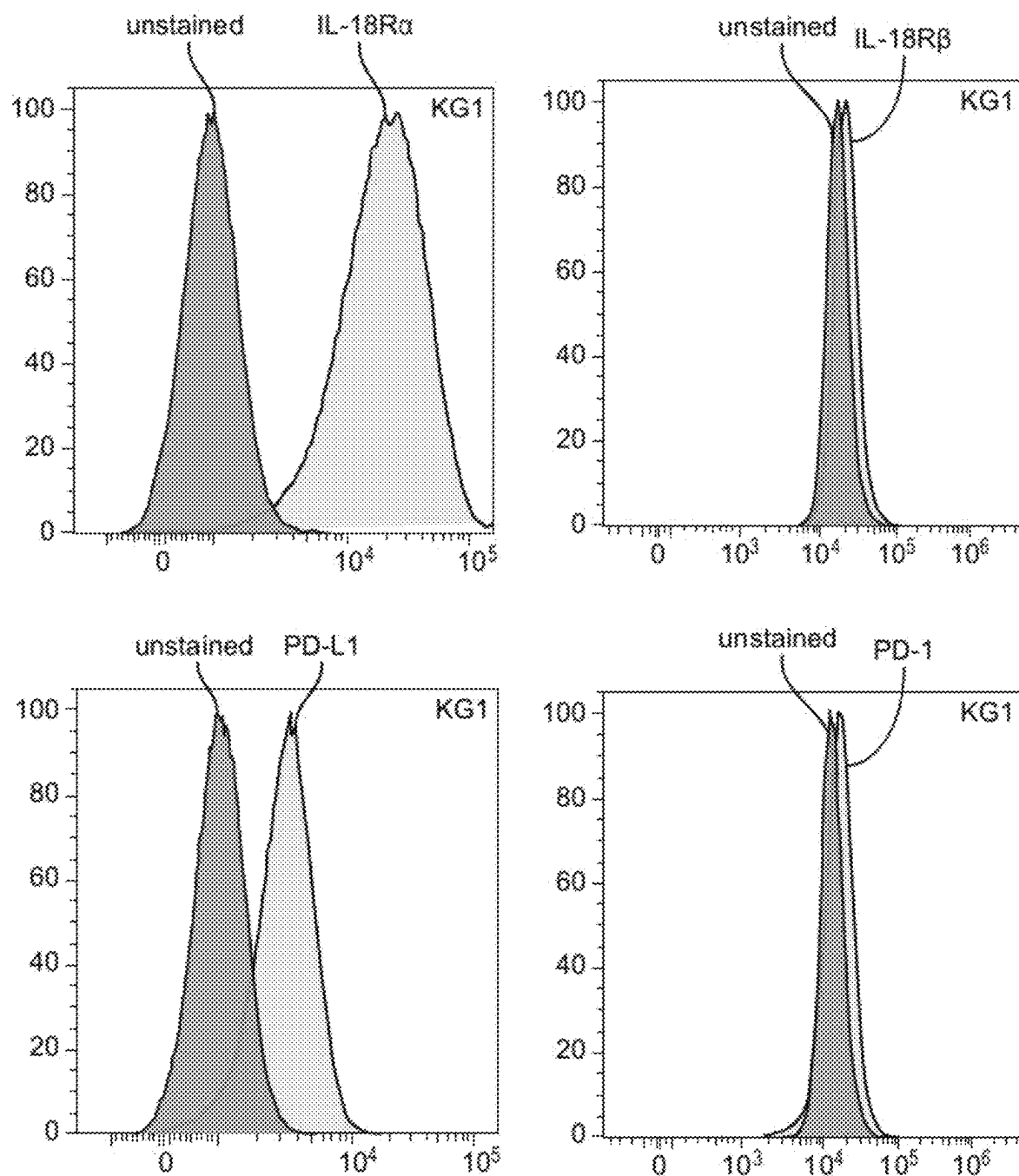


FIGURE 21

**Flow Cytometry Analysis of KG1 Cells****FIGURE 22**

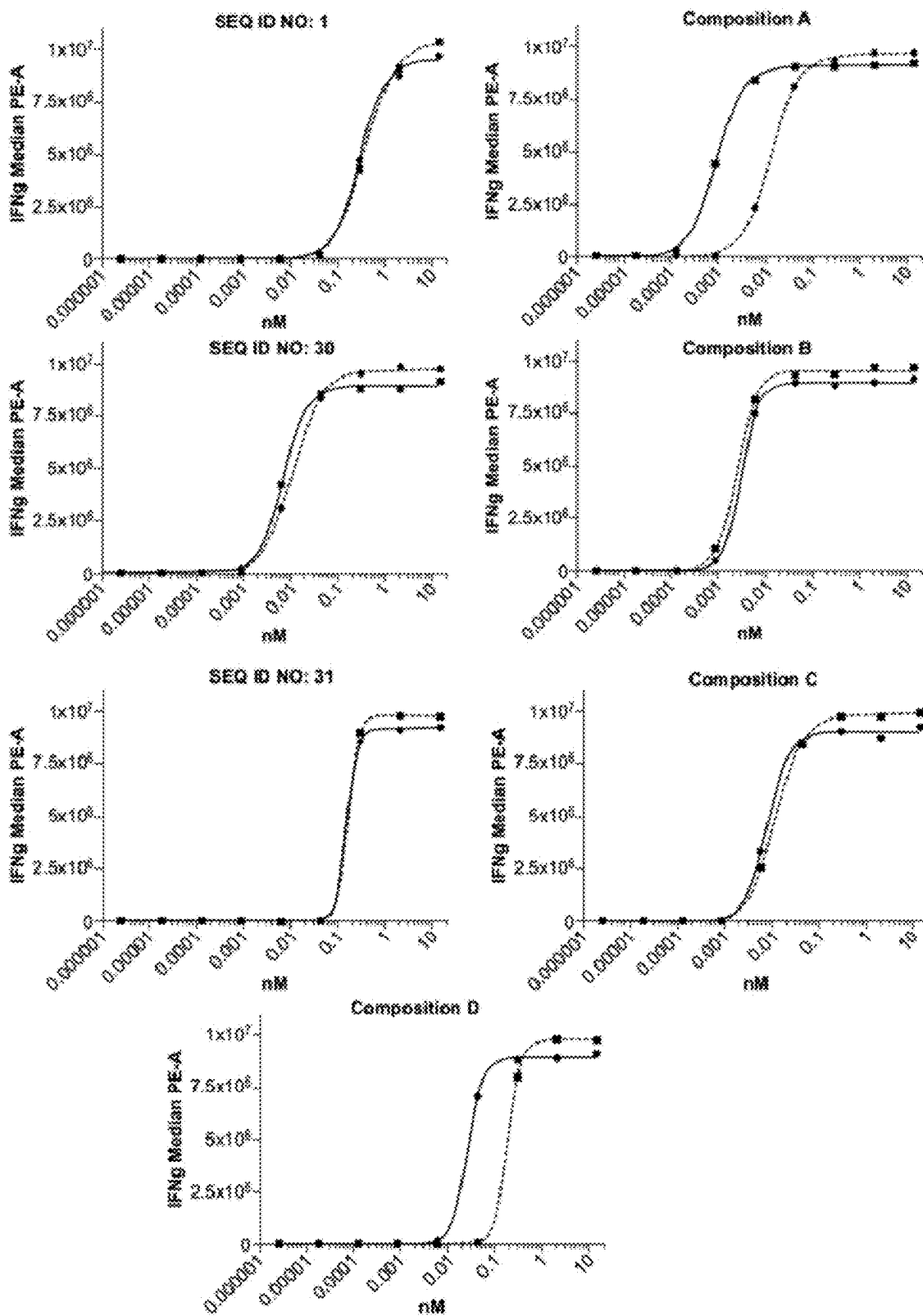


FIGURE 23

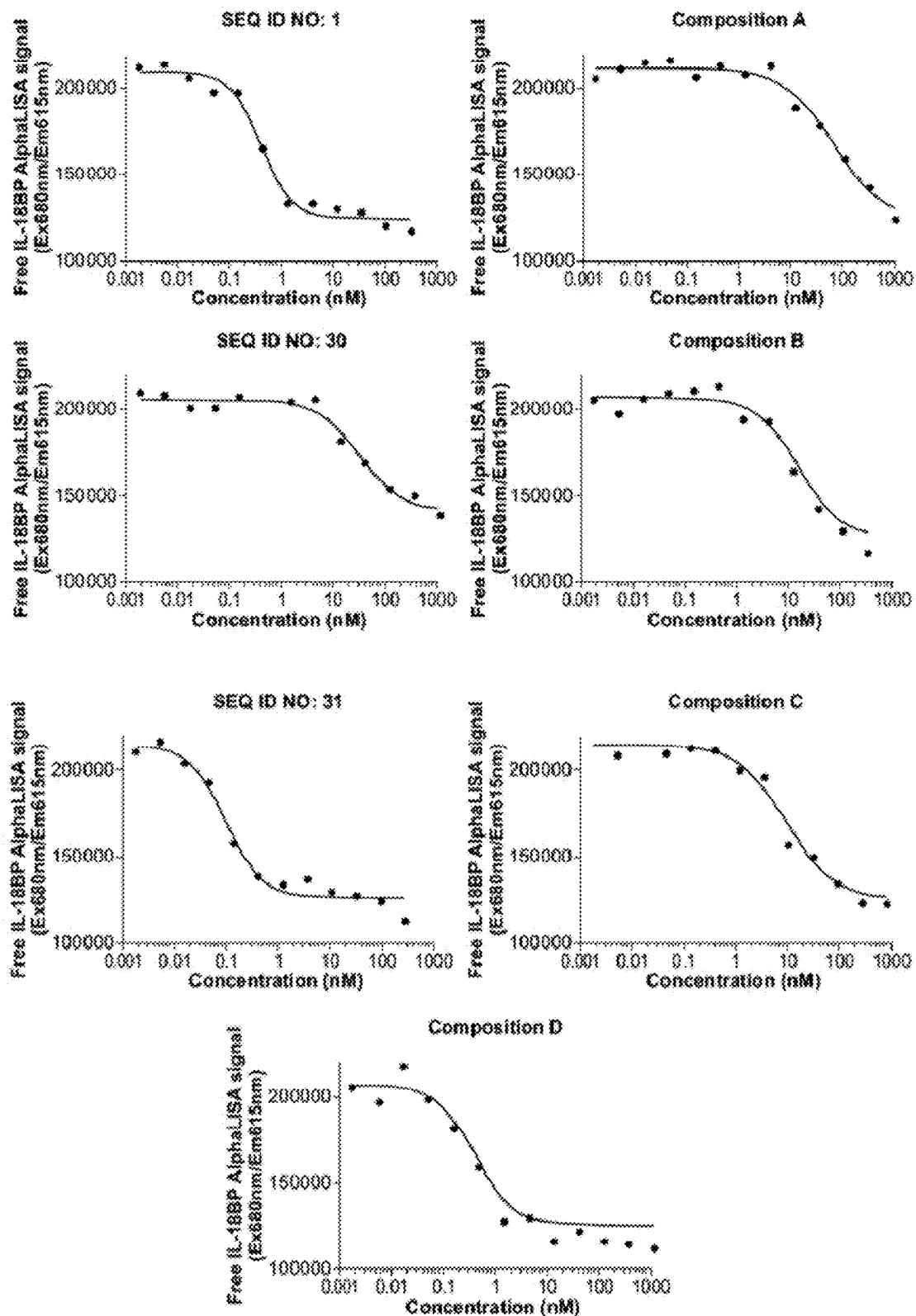


FIGURE 24

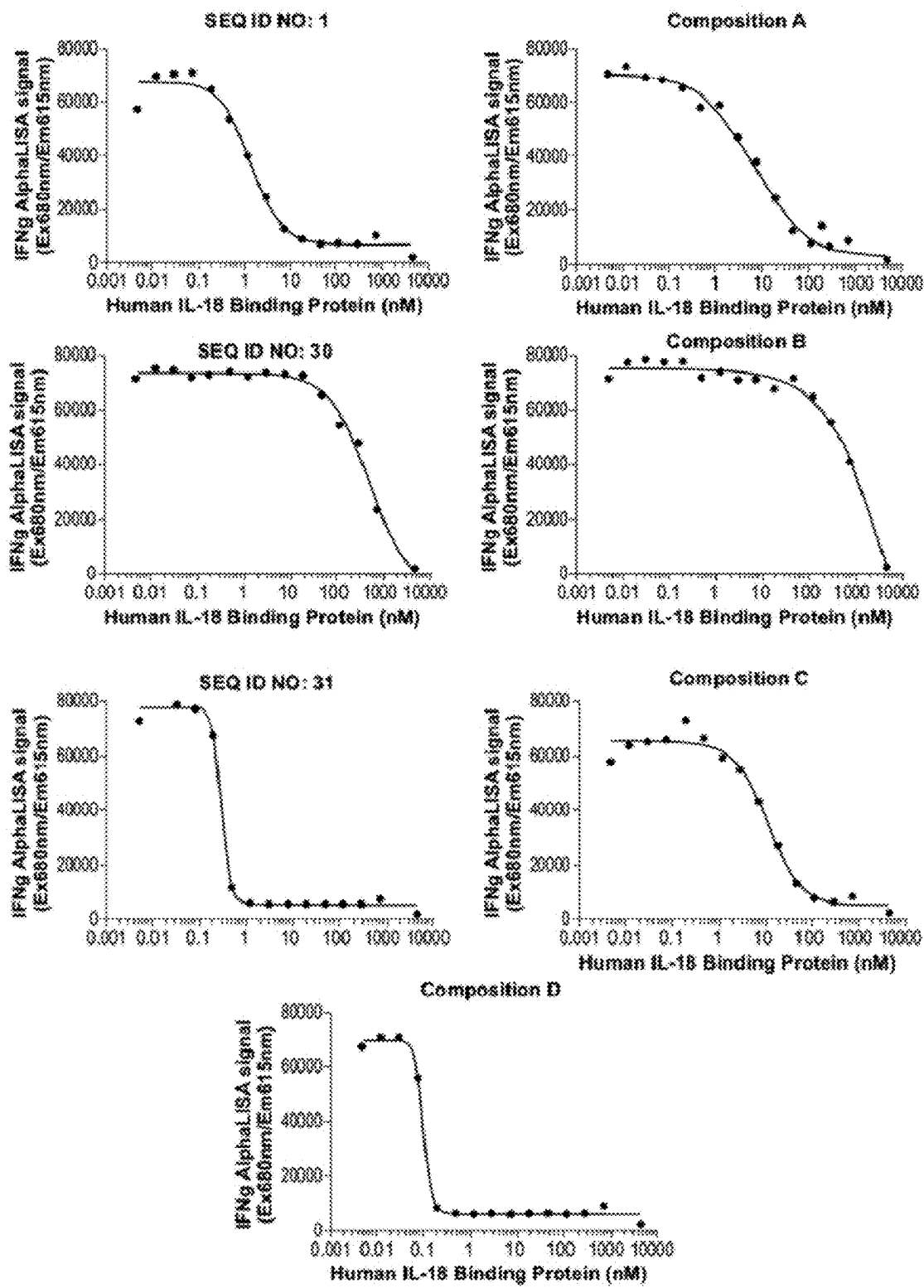


FIGURE 25

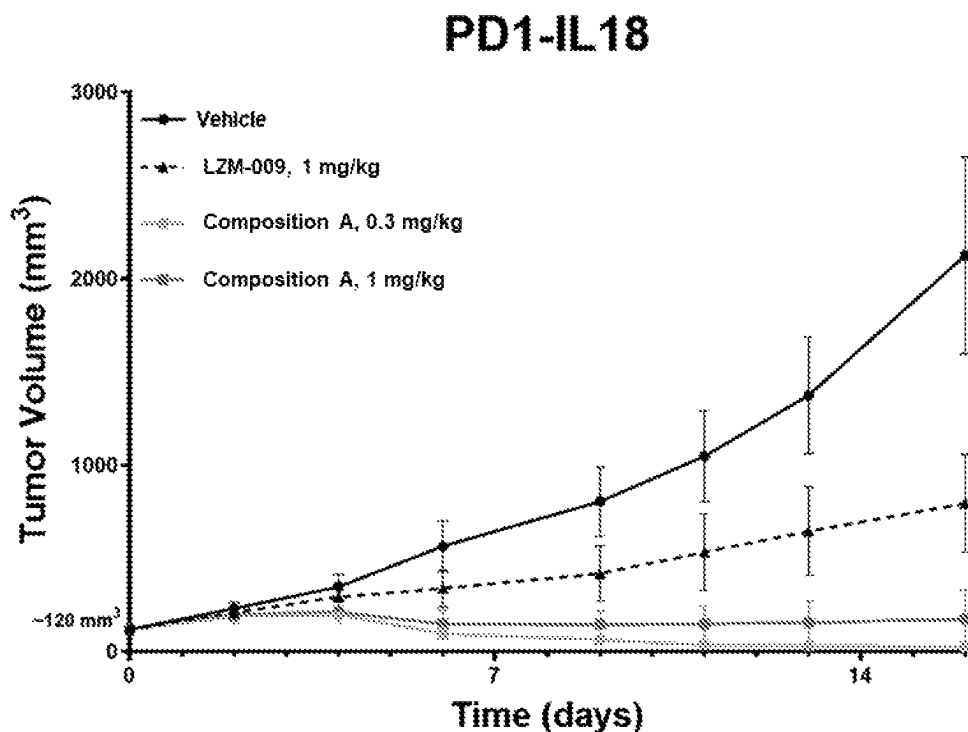


FIGURE 26A

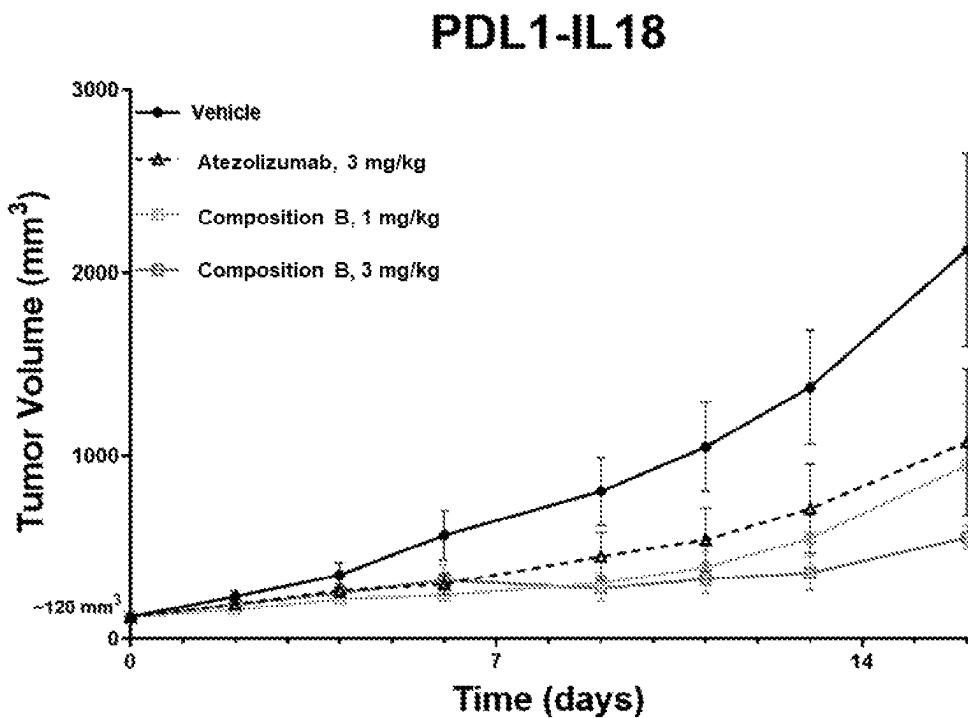


FIGURE 26B

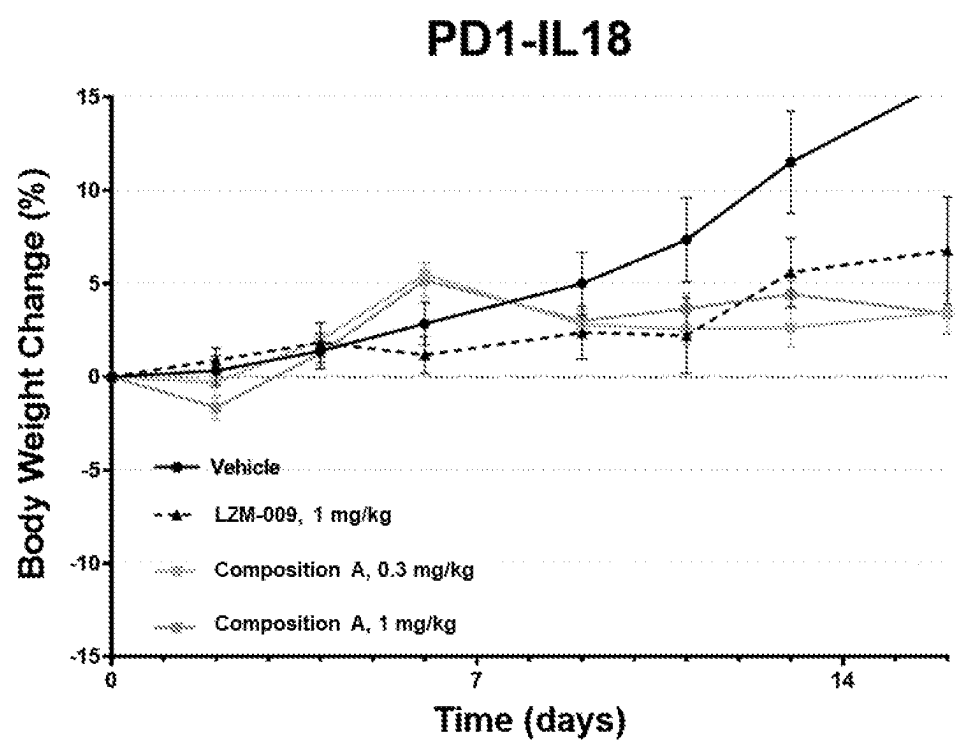
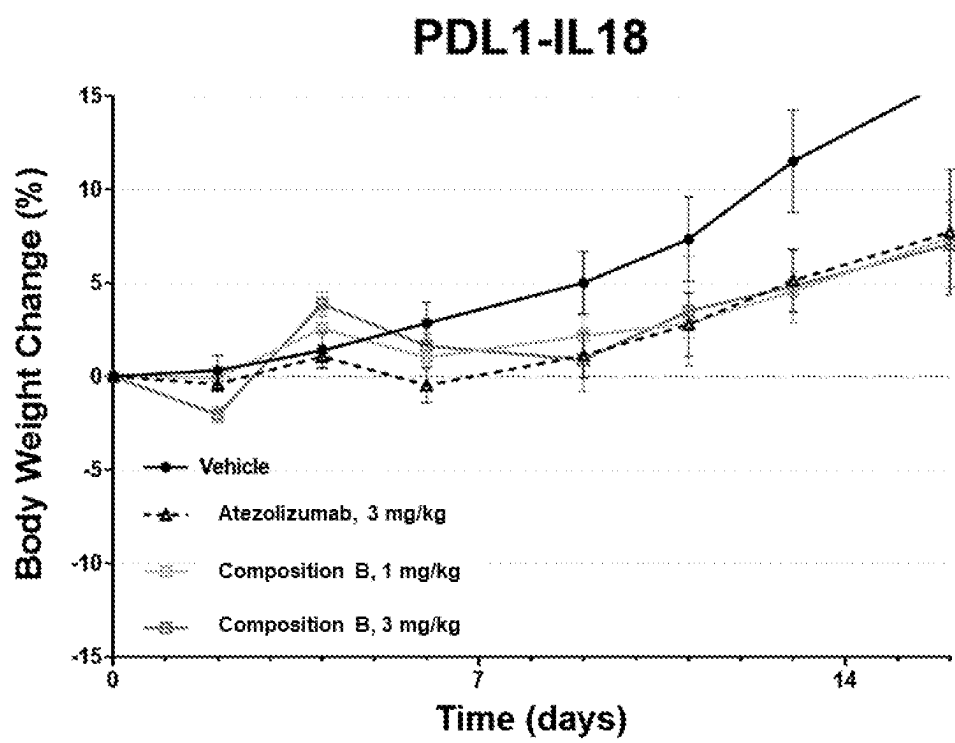


FIGURE 27A

**FIGURE 27B**

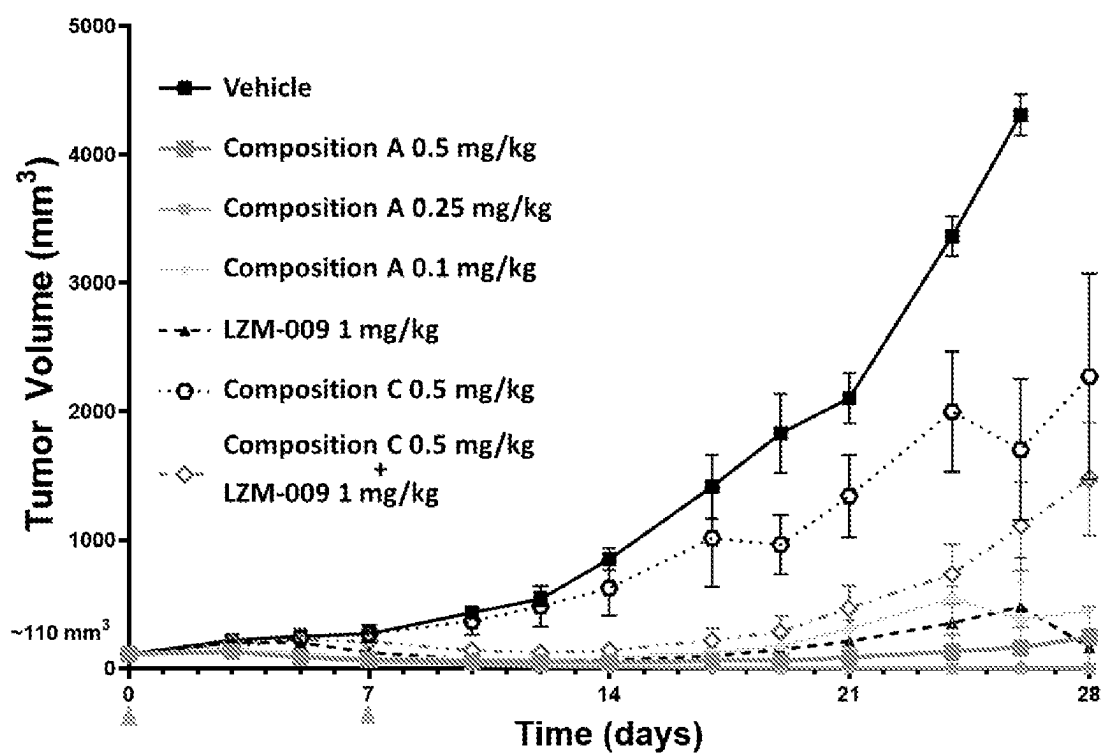


FIGURE 28A

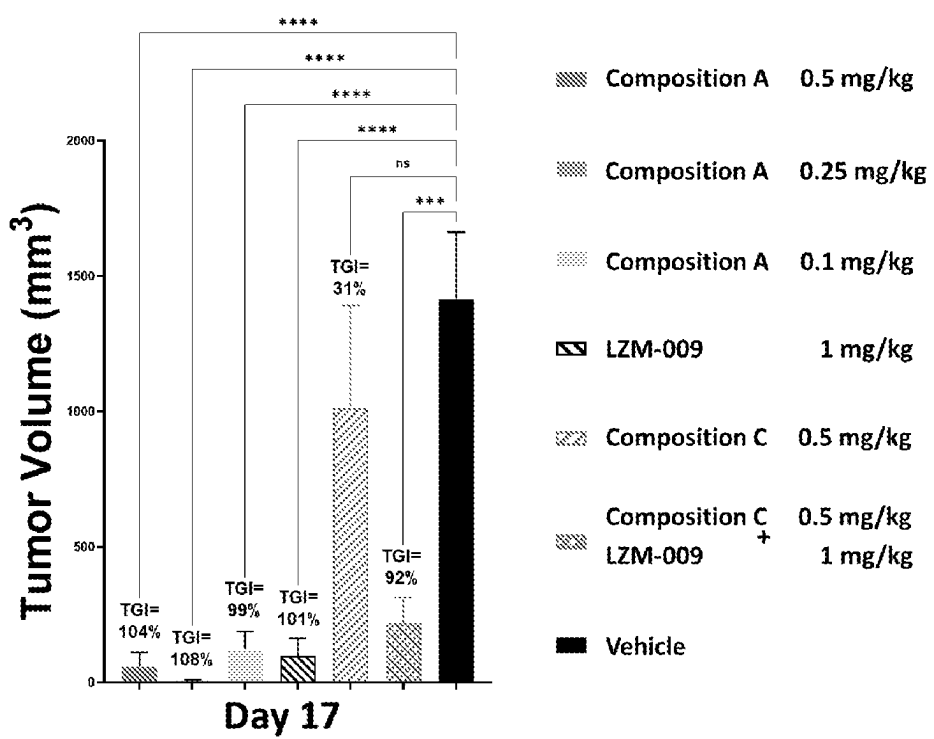


FIGURE 28B

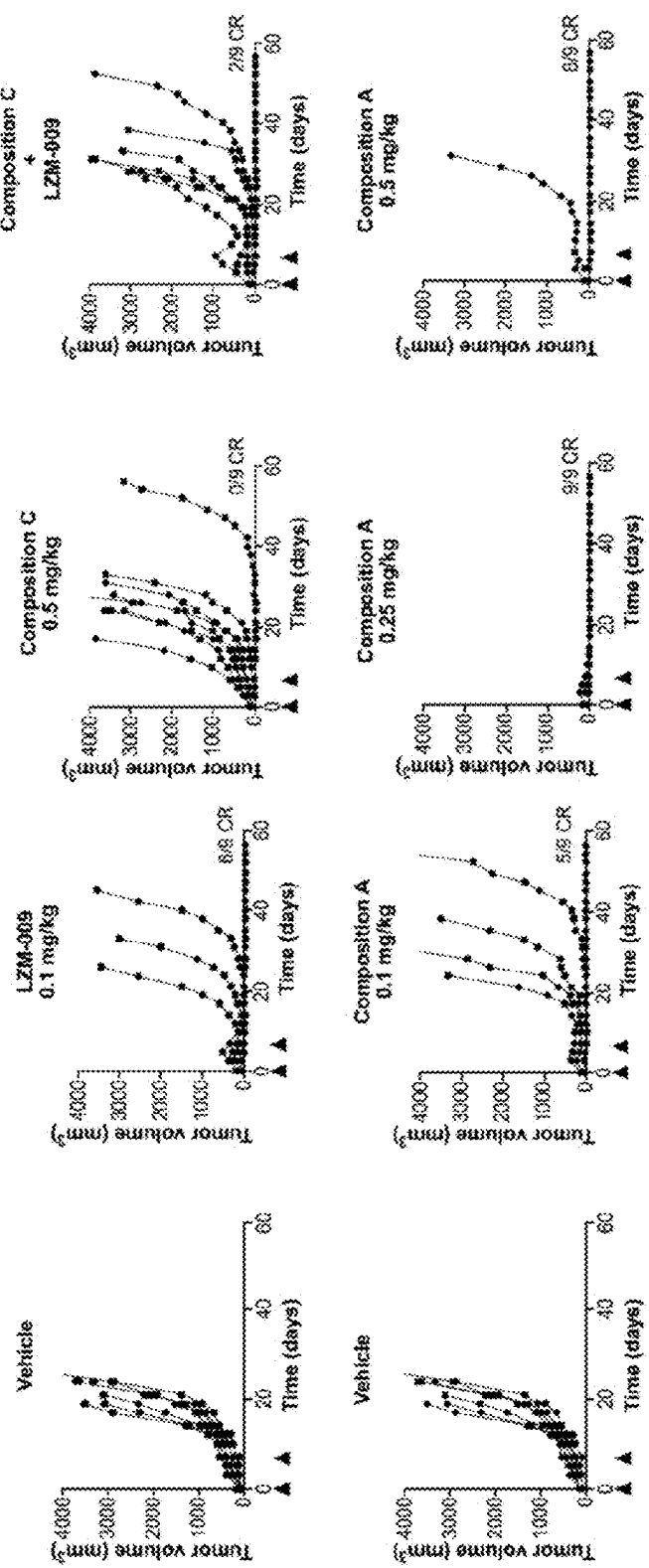


FIGURE 28C

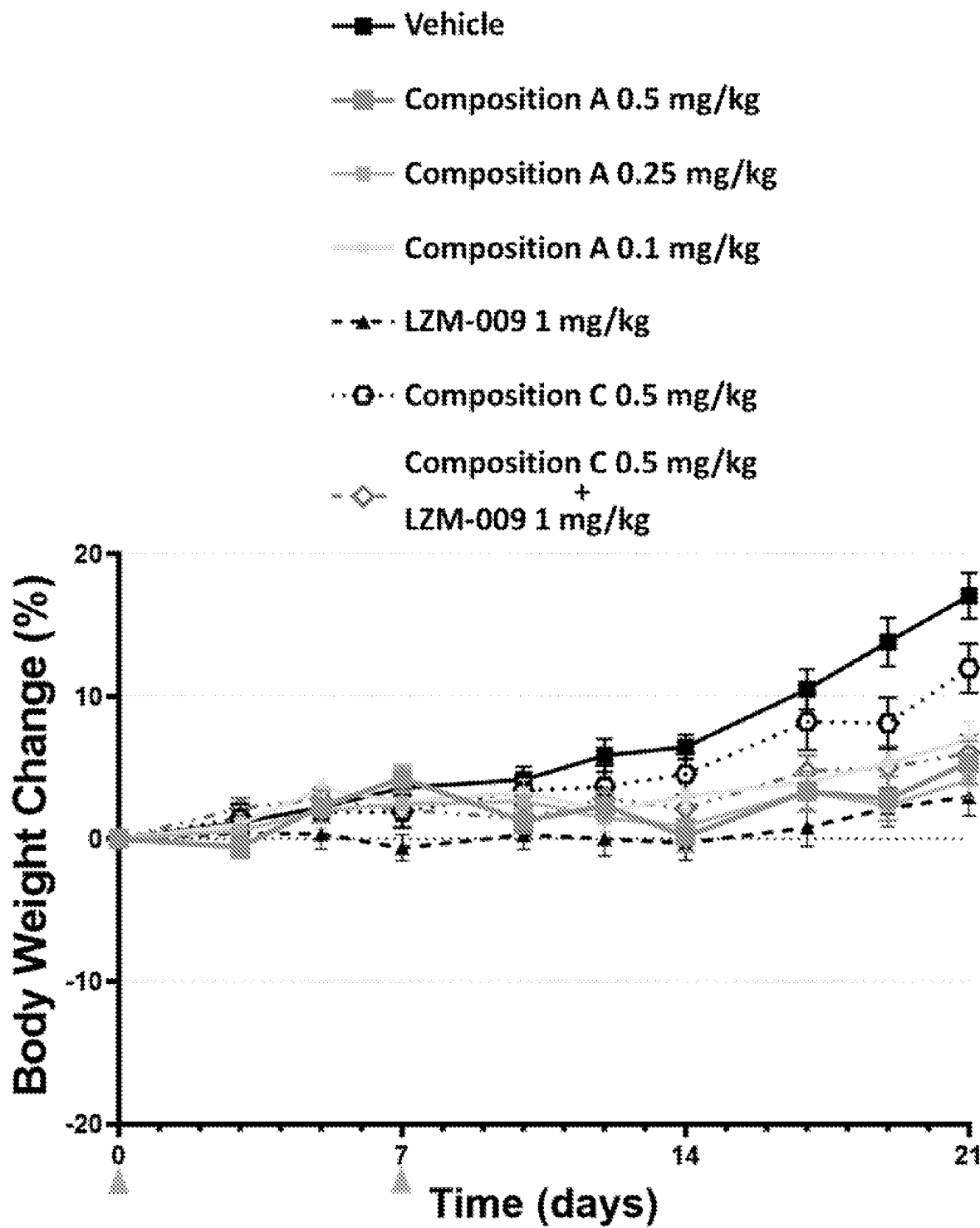


FIGURE 29

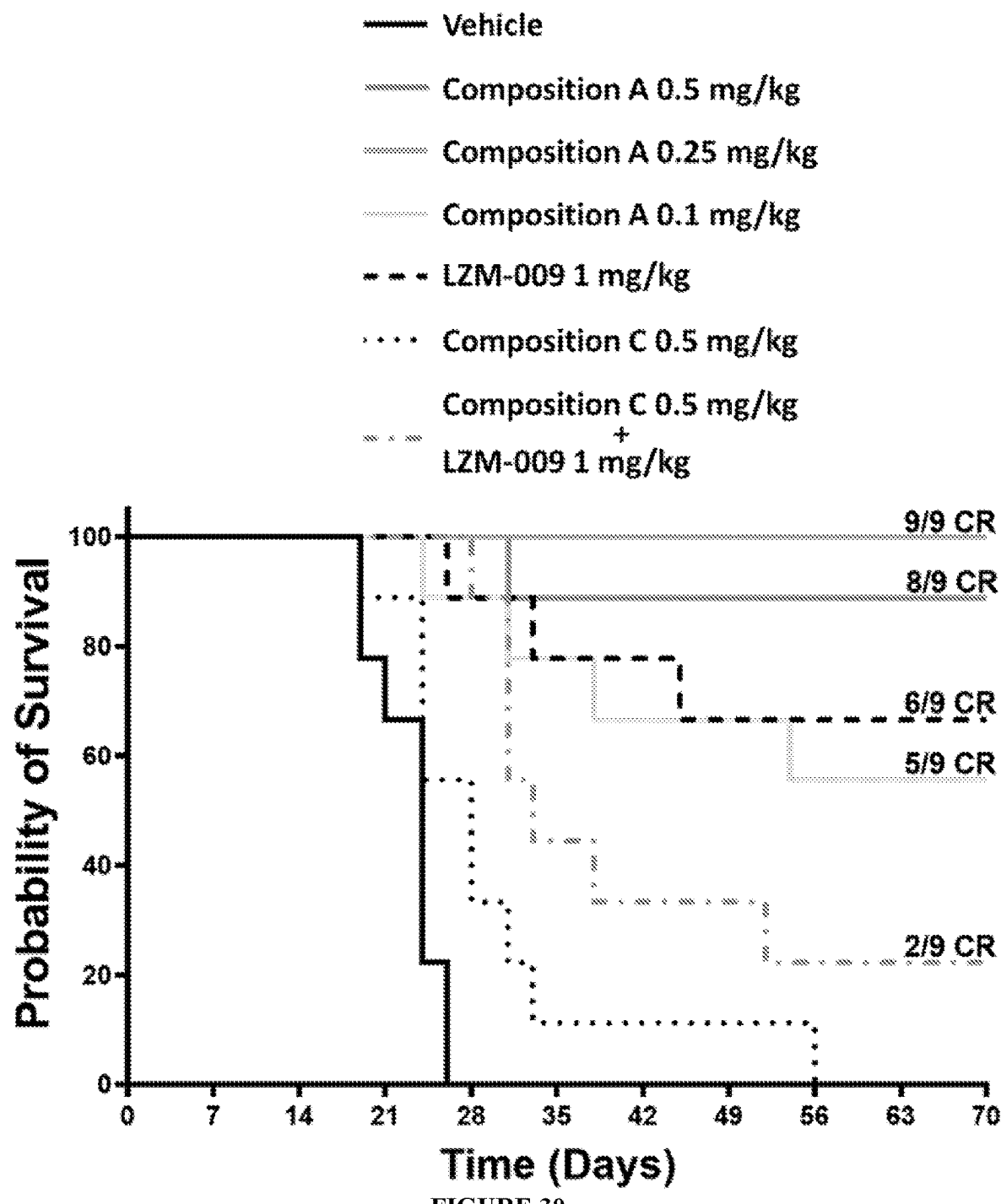


FIGURE 30

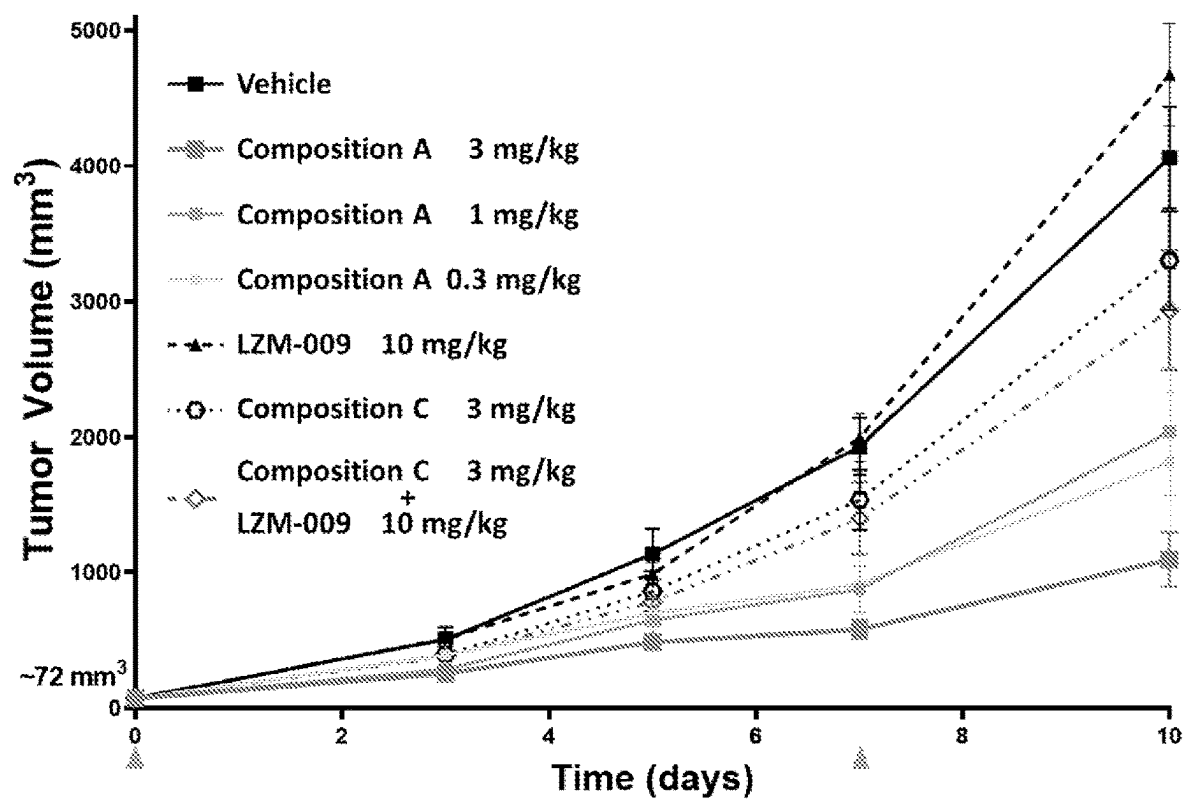


FIGURE 31A

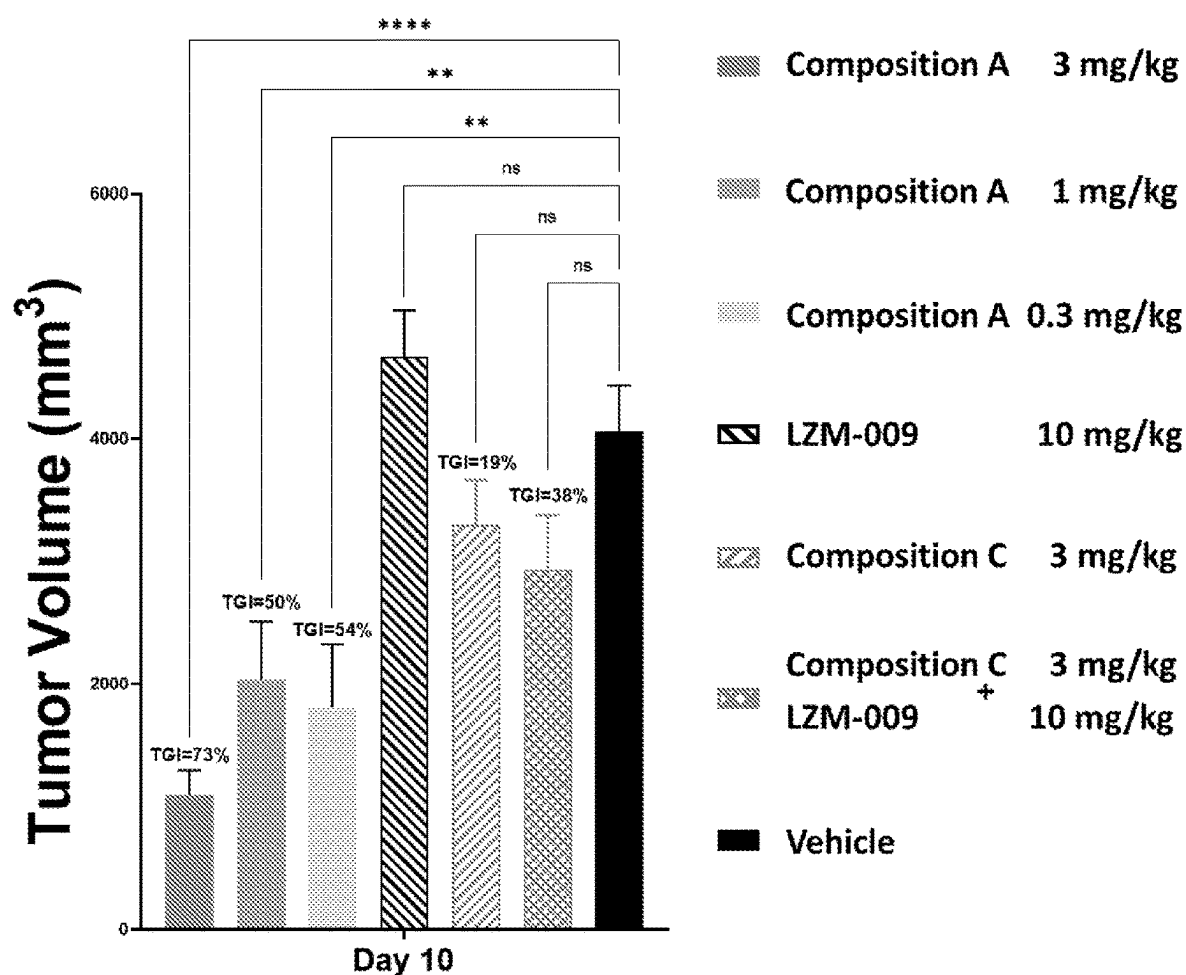


FIGURE 31B

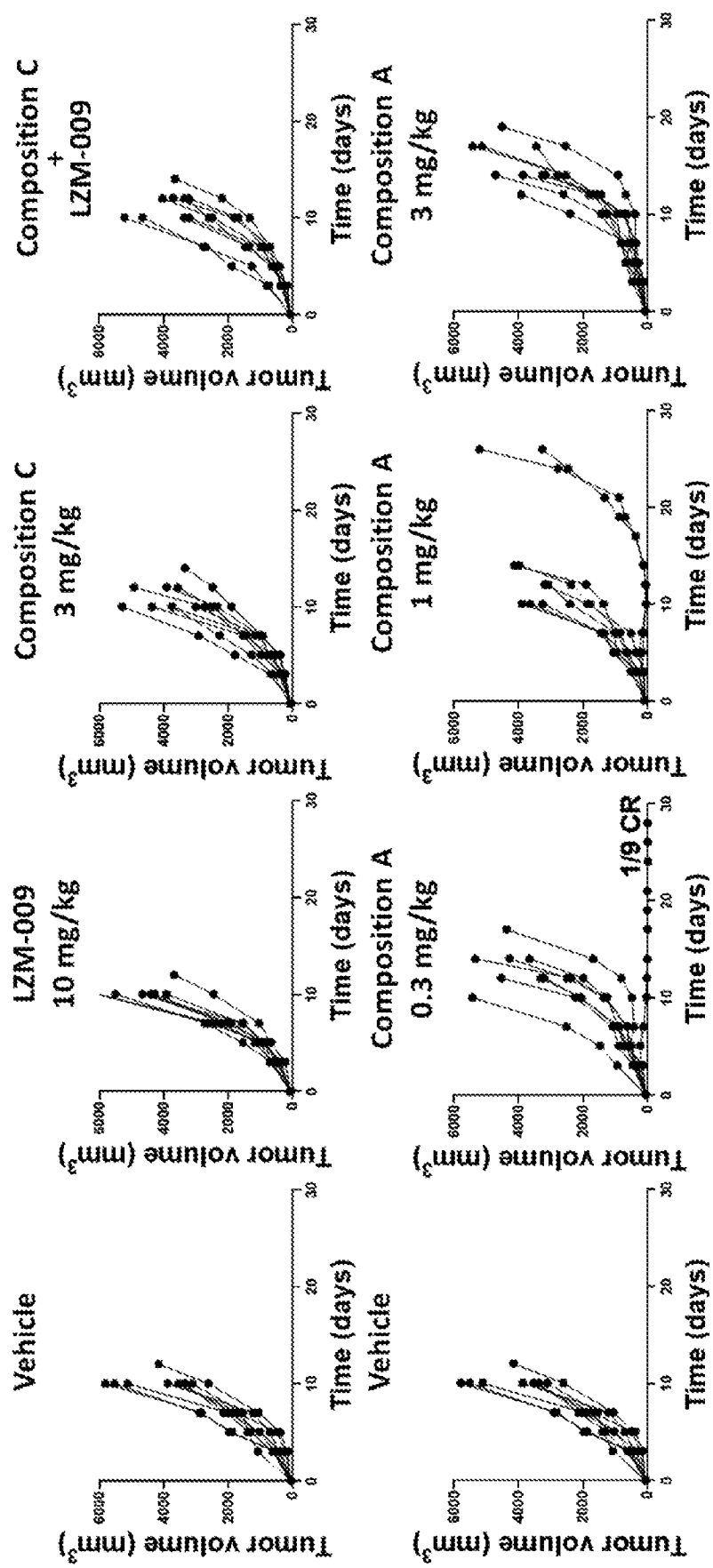


FIGURE 31C

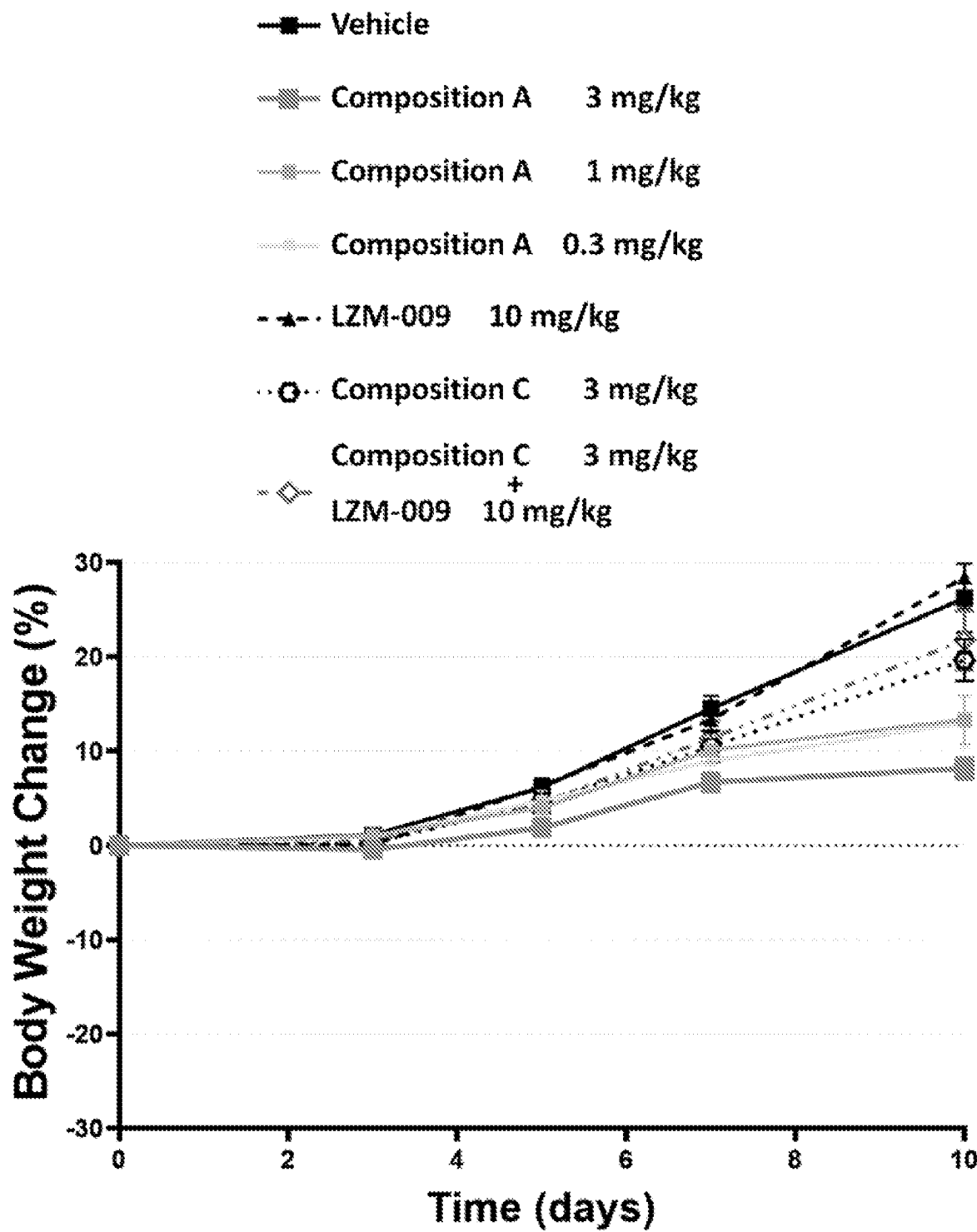


FIGURE 32

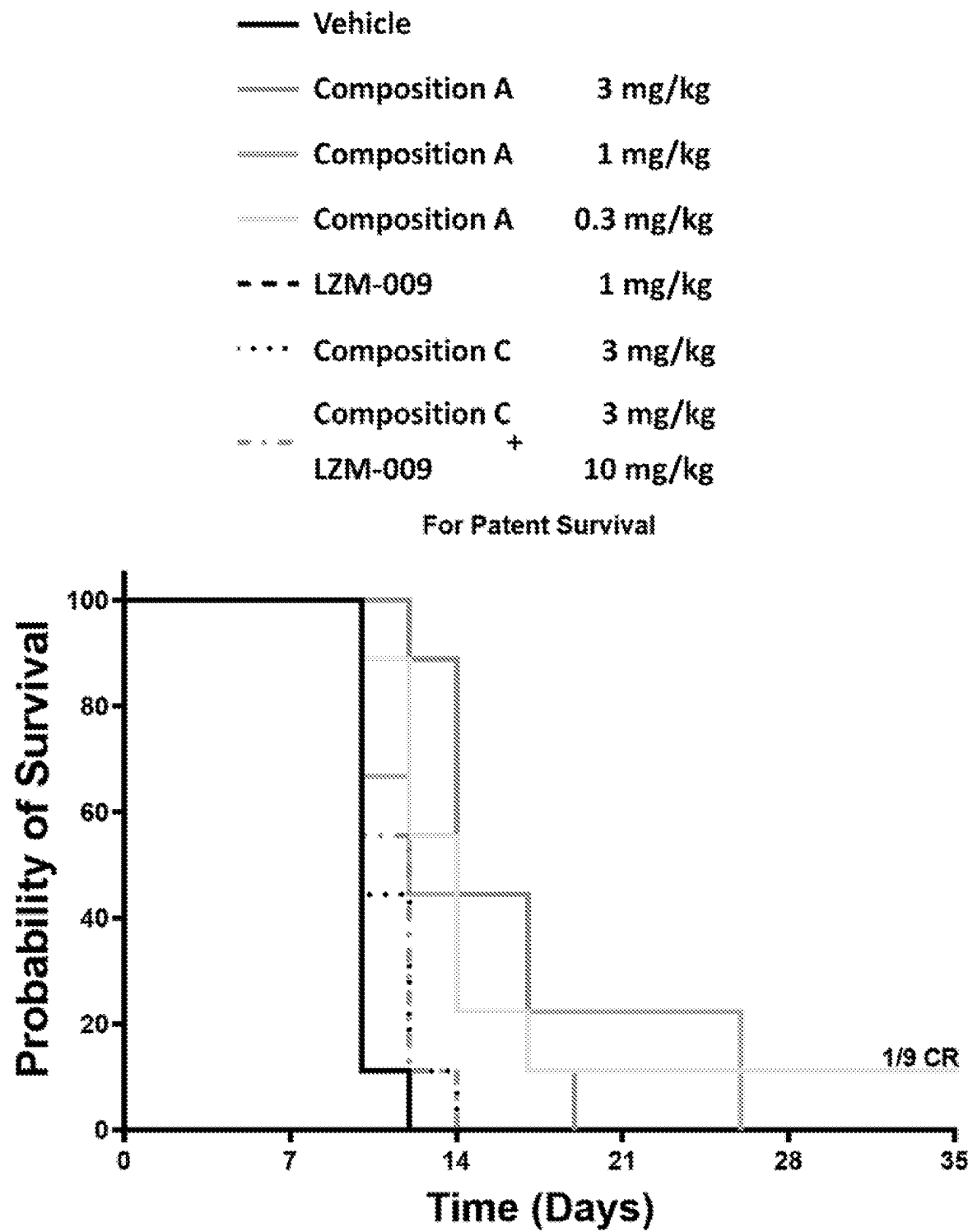


FIGURE 33

**IMMUNE ANTIGEN SPECIFIC IL-18  
IMMUNOCYTOKINES AND USES THEREOF**

CROSS REFERENCE

This application claims the benefit of U.S. Provisional Application No. 63/313,222 filed Feb. 23, 2022, and U.S. Provisional Application No. 63/479,529 filed Jan. 11, 2023, which applications are incorporated herein by reference in their entirety.

CROSS REFERENCE

The instant application contains a Sequence Listing which has been submitted electronically in XML file format and is hereby incorporated by references in its entirety. Said XML copy, created on Apr. 6, 2023, is named 94917-0082\_725201US\_SL.xml and is 52,712 bytes in size.

SEQUENCE LISTING

The instant application contains a Sequence Listing which has been submitted electronically in XML file format and is hereby incorporated by reference in its entirety. Said XML copy, created on Apr. 6, 2023, is named 94917-0082\_725201US\_SL.xml and is 252,712 bytes in size.

BACKGROUND

In 2022, an estimated 1.8 million new cases of cancer will be diagnosed in the United States, and over 600,000 people will die from the disease. Immunotherapies utilize the immune system of a subject to aid in the treatment of ailments. Immunotherapies can be designed to either activate or suppress the immune system depending on the nature of the disease being treated. A goal of various immunotherapies for the treatment of cancer is to stimulate the immune system so that it recognizes and destroys tumors or other cancerous tissue.

Immune cells implicated in response to various cancers express certain proteins that are implicated in the regulation of the immune response. These proteins, such as programmed cell death protein 1 (PD-1), programmed death-ligand 1 (PD-L1), and others, can downregulate the immune system and promote self-tolerance by suppressing T cell inflammatory activity. In light of these mechanisms, antibodies or antigen binding fragments which target these proteins have been identified as potential therapeutics. However, in some cases single mechanism therapies targeting these proteins alone are insufficient for treating cancer. Thus, there is a need for improved tools for cancer therapy.

BRIEF SUMMARY

Described herein are immunocytokine compositions which contain an immune cell associated antigen specific antibody or antigen binding fragment thereof and an interleukin 18 (IL-18) polypeptide.

In one aspect, provided herein, is an immunocytokine composition, comprising: an IL-18 polypeptide; and an antibody or an antigen binding fragment thereof specific for an immune cell associated antigen; and a linker, wherein the linker comprises: a first point of attachment to the IL-18 polypeptide; and a second point of attachment to the antibody or antigen binding fragment thereof.

In some embodiments, the linker comprises a polymer. In some embodiments, the polymer comprises a water-soluble

polymer. In some embodiments, the water-soluble polymer comprises poly(alkylene oxide), polysaccharide, poly(vinyl pyrrolidone), poly(vinyl alcohol), polyoxazoline, poly(acryloylmorpholine), or a combination thereof. In some embodiments, the polymer has a weight average molecular weight of at least about 0.1 kDa, at least about 0.5 kDa, at least about 1 kDa, at least about 5 kDa, at least about 10 kDa, at least about 20 kDa, at least about 30 kDa, at least 40 kDa, at least 50 kDa, at least 60 kDa, at least 70 kDa, at least 80 kDa, at least 90 kDa, at least 100 kDa, at least 120 kDa, at least 130 kDa, at least 140 kDa, at least 150 kDa, or more.

In some embodiments, the first point of attachment is at a residue which is not the N-terminus or the C-terminus of the IL-18 polypeptide. In some embodiments, the first point of attachment is to a residue in the region of residues 2-156 of the IL-18 polypeptide, wherein residue position numbering is based on SEQ ID NO: 1. In some embodiments, the first point of attachment is to a residue in the region of residues 38-144 of the IL-18 polypeptide. In some embodiments, the first point of attachment is at residue 38, 68, 69, 70, 76, 78, 85, 86, 95, 98, 121, 127, or 144 of the IL-18 polypeptide. In some embodiments, the first point of attachment is at residue 68, 69, 70, 85, 86, 95, or 98 of the IL-18 polypeptide. In some embodiments, the first point of attachment is at residue 68 of the IL-18 polypeptide. In some embodiments, the IL-18 polypeptide comprises an amino acid substitution at the first point of attachment. In some embodiments, the amino acid substitution at the first point of attachment is for an unnatural amino acid. In some embodiments, the amino acid substitution at the first point of attachment is for a natural amino acid. In some embodiments, the amino acid substitution at the first point of attachment is E69C, K70C, E85C, M86C, T95C, or D98C.

In some embodiments, the second point of attachment is at an amino acid residue in an Fc region of the antibody or antigen binding fragment. In some embodiments, the Fc region comprises an amino acid sequence having 90% or more identity to the amino acid sequence of SEQ ID NO: 260. In some embodiments, the second point of attachment is (a) an amino acid residue at positions 16 to 18 of SEQ ID NO: 260, (b) an amino acid residue at positions 58 to 60 of SEQ ID NO: 260, or (c) amino acid residue 87 of SEQ ID NO: 260. In some embodiments, the second point of attachment is to a lysine residue on the antibody or antigen binding fragment. In some embodiments, the IL-18 polypeptide displays reduced binding to IL-18 binding protein (IL-18BP) compared to a wild type IL-18 polypeptide of SEQ ID NO: 1. In some embodiments, the IL-18 polypeptide contains one or more amino acid substitutions that are located at residue positions selected from Y01, F02, E06, V11, C38, K53, D54, S55, T63, C76, and C127, wherein residue position numbering of the IL-18 polypeptides are based on SEQ ID NO: 1 as a reference sequence. In some embodiments, the one or more amino acid substitutions in the IL-18 polypeptide are selected from Y01G, F02A, E06K, V11, C38S, C38A, K53A, D54A, S55A, T63A, C68S, C76S, C76A, C127S, and C127A. In some embodiments, the IL-18 polypeptide contains E06K and K53A amino acid substitutions. In some embodiments, the IL-18 polypeptides contains a T63A amino acid substitution. In some embodiments, the IL-18 polypeptide is synthetic. In some embodiments, the IL-18 polypeptide comprises one or more amino acid substitutions selected from: (a) a homoserine residue located at any one of residues 26-36; (b) a homoserine residue located at any one of residues 45-67; (c) a homoserine residue located any one of residues 70-80; (d) a homoserine residue located at any one of residues 110-130; (e) a norleucine or

O-methyl-homoserine residue located at any one of residues 28-38; (f) a norleucine or O-methyl-homoserine residue located at any one of residues 46-56; g) a norleucine or O-methyl-homoserine residue located at any one of residues 54-64; (h) a norleucine or O-methyl-homoserine residue located at any one of residues 80-90; (i) a norleucine or O-methyl-homoserine residue located at any one of residues 108-118; and (j) a norleucine or O-methyl-homoserine residue located at any one of residues 145-155; wherein residue position numbering of the IL-18 polypeptide is based on SEQ ID NO: 1 as a reference sequence. In some embodiments, the IL-18 polypeptide comprises one or more amino acid substitutions selected from homoserine (Hse) 31, norleucine (Nle) 33, O-methyl-homoserine (Omh) 33, Hse50, Nle51, Omh51, Hse57, Nle60, Omh60, Hse63, Hse67, Hse75, Nle86, Omh86, Hse106, Nle113, Omh113, Hse116, Hse 121, Nle150, and Omh150.

In some embodiments, the antibody or antigen binding fragment thereof is a monoclonal antibody, a humanized antibody, a grafted antibody, a chimeric antibody, a human antibody, a de-immunized antibody, or a bispecific antibody. In some embodiments, the antibody or antigen binding fragment thereof an antigen binding fragment, wherein the antigen binding fragment comprises a Fab, a Fab", a F(ab')<sub>2</sub>, a bispecific F(ab')<sub>2</sub>, a trispecific F(ab')<sub>2</sub>, a variable fragment (Fv), a single chain variable fragment (scFv), a dsFv, a bispecific scFv, a variable heavy domain, a variable light domain, a variable NAR domain, bispecific scFv, a minobody, a diabody, a bispecific diabody, triabody, a tetrabody, a minibody, a maxibody, a camelid, a VHH, a minibody, an intrabody, fusion proteins comprising an antibody portion (a domain antibody), a single chain binding polypeptide, a scFv-Fc, a Fab-Fc, a bispecific T cell engager (BiTE), a tetravalent tandem diabody (TandAb), a Dual-Affinity Retargeting Antibody (DART), a bispecific antibody (bscAb), a single domain antibody (sdAb), a fusion protein, or a bispecific disulfide-stabilized Fv antibody fragment (dsFv-dsFv'). In some embodiments, the antibody or antigen binding fragment thereof comprises an IgG, an IgM, an IgE, an IgA, an IgD, or is derived therefrom. In some embodiments, the antibody or antigen binding fragment thereof comprises the IgG, and wherein the IgG is an IgG1, an IgG4, or is derived therefrom. In some embodiments, the immune cell associated antigen is 4-IBB, B7-H3, B7-H4, BTLA, CD3, CCR8, CD8A, CD8B, CD16A, CD27, CD28, CD33, CD38, CD39, CD40, CD47, CD70, CD80, CD86, CD96, CD163, CLEC-1, CLEVER-1, CTLA-4, D40, GITR, ICOS, ILT2/3/4, LAG-3, MHCI, MHCII, NKG2A, NKG2D, NKp30, NKp44, NKp46, OX40, PD-1, PD-L1, PD-L2, PSGL-1, SIGLEC-9, SIGLEC-15, SIRP- $\alpha$ , TCR, TIGIT, TIM-3, VISTA, or VSIG4.

In some embodiments, the immune cell associated antigen is PD-1. In some embodiments, the antibody or antigen binding fragment thereof comprises tislelizumab, baizean, OKVO411B3N, BGB-A317, hu317-1/IgG4mt2, sintilimab, tyvyt, IBI-308, toripalimab, TeRuiPuLi, terepril, tuoyi, JS-001, TAB-001, tamrelizumab, HR-301210, INCNSHR-01210, SHR-1210, temiplimab, cemiplimab-rwlc, 6QVL057INT, H4H7798N, REGN-2810, SAR-439684, lambrolizumab, pembrolizumab, MK-3475, SCH-900475, h409A11, nivolumab, nivolumab BMS, BMS-936558, MDX-1106, ONO-4538, prolgolimab, forteca, BCD-100, penpulimab, AK-105, zimberelimab, AB-122, GLS-010, WBP-3055, balstilimab, 1Q2QT5M7EO, AGEN-2034, AGEN-2034w, genolimzumab, geptanolimab, APL-501, CBT-501, GB-226, dostarlimab, ANB-011, GSK-4057190A, P0GVQ9A4S5, TSR-042, WBP-285, serplu-

limab, HLX-10, CS-1003, retifanlimab, 2Y3T5IF01Z, INCMGA-00012, INCMGA-0012, MGA-012, sasanlimab, LZZ0IC2EWP, PF-06801591, RN-888, spartalizumab, PDR-001, QOG25L6Z8Z, relatlimab/nivolumab, BMS-986213, cetrelimab, JNJ-3283, JNJ-63723283, LYK98WP91F, tebotelimab, MGD-013, BCD-217, BAT-1306, HX-008, MEDI-5752, JTX-4014, cadonilimab, AK-104, BI-754091, pidilizumab, CT-011, MDV-9300, YBL-006, AMG-256, RG-6279, RO-7284755, BH-2950, IBI-315, RG-6139, RO-7247669, ONO-4685, AK-112, 609-A, LY-3434172, T-3011, AMG-404, IBI-318, MGD-019, ONCR-177, LY-3462817, RG-7769, RO-7121661, F-520, XmAb-23104, Pd-1-pik, SG-001, S-95016, Sym-021, LZM-009, budigalimab, 6VDO4TY3OO, ABBV-181, PR-1648817, CC-90006, XmAb-20717, 2661380, AMP-224, B7-DC Ig, EMB-02, ANB-030, PRS-332, [89Zr]Deferoxamide-pembrolizumab, 89Zr-Df-Pembrolizumab, [89Zr] Df-Pembrolizumab, STI-1110, STI-A1110, CX-188, mPD-1, MCLA-134, 244C8, ENUM 224C8, ENUM C8, 388D4, ENUM 388D4, ENUM D4, MEDIO680, NVP-LZV-184, or AMP-514, or a modified version thereof. In some embodiments, the antibody or antigen binding fragment thereof comprises nivolumab, pembrolizumab, LZM-009, or cemiplimab, or a modified version thereof.

In some embodiments, the immune cell associated antigen is PD-L1. In some embodiments, the antibody is Avelumab (Bavencio, 451238, KXG2PJ551I, MSB-0010682, MSB-0010718C, PF-06834635, CAS 1537032-82-8), Durvalumab (Imfinzi, 28×28×90 KV (UNII code), MEDI-4736, CAS 1428935-60-7), Atezolizumab (Tecentriq, 52CM10WC3Y, MPDL-3280A, RG-7446, RO-5541267, CAS 1380723-44-3), Sugemalimab (CS-1001, WBP-3155), KN-046 (CAS 2256084-03-2), APL-502 (CBT-502, TQB-2450), Envafolimab (3D-025, ASC-22, KN-035, hu56V1-Fc-m1, CAS 2102192-68-5), Bintrafusp alfa (M-7824, MSB-0011359C, NW9K8C1JN3, CAS 1918149-01-5), STI-1014 (STI-A1014, ZKAB-001), PD-L1 t-haNk, A-167 (HBM-9167, KL-A167), IMC-001 (STI-3031, STI-A-1015, STI-A1015, s), HTI-1088 (SHR-1316), IO-103, CX-072 (CytomX Therapeutics), AUPM-170 (CA-170), GS-4224, ND-021 (NM21-1480, PRO-1480), BNT-311 (DuoBody-PD-L1x4-1BB, GEN-1046), BGB-A333, IBI-322, NM-01, LY-3434172, LDP, CDX-527, IBI-318, 89Zr-DFO-REGN3504, ALPN-202 (CD80 vIgD-Fc), INCB-086550, LY-3415244, SHR-1701, JS-003 (JS003-30, JS003-SD), HLX-20 (PL2 #3), ES-101 (INBRX-105, INBRX-105-1), MSB-2311, PD-1-Fc-OX40L (SL-279252, TAK-252), FS-118, FS118 mAb2, LAG-3/PD-L1 mAb2), FAZ-053 (LAE-005), Lodapolimab (LY-3300054, NR4MAD6PPB, CAS 2118349-31-6), MCLA-145, BMS-189, Cosibelimab (CK-301, TG-1501, CAS 2216751-26-5), IL-15Ralpha-SD/IL-15 (KD-033), WP-1066 (CAS 857064-38-1), BMS-936559 (MDX-1105), BMS-986192, RC-98, CD-200AR-L (CD200AR-L), ATA-3271, IBC-Ab002, BMX-101, AVA-04-VbP, ACE-1708, KY-1043, ACE-05 (YBL-013), ONC-0055 (ONC0055, PRS-344 S-095012), TLJ-1-CK, GR-1405, PD1ACR-T, N-809 (N-IL15/PDL1), CB-201, MEDI-1109, AVA-004 (AVA-04), CA-327, ALN-PDL, KY-1003, CD22(aPD-L1)CAR-T cells (SL-22P), ATA-2271 (M28z1XXPD1DNR CAR T cells), or a modified version thereof. In some embodiments, the antibody is durvalumab, atezolizumab, or avelumab, or a modified version thereof.

In some embodiments, the antibody or antigen binding fragment thereof comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%,

at least 97%, at least 98%, or at least 99% identical to an amino acid sequence of Table 1.

In some embodiments, the IL-18 polypeptide comprises an amino acid sequence having at least 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% sequence identity to the sequence set forth in SEQ ID NO: 1. In some embodiments, the IL-18 polypeptide of the immunocytokine provided herein comprises an amino acid sequence of any one of SEQ ID NOs: 2-67. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence of SEQ ID NO: 30. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence of SEQ ID NO: 68-72.

In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to at least one Fc receptor which is within 10-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits binding affinity ( $K_D$ ) to the antigen of the antibody which is within 5-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits an ability to induce IFN $\gamma$  production in a cell as measured by half-maximal effective concentration ( $EC_{50}$ ) which is within about 100-fold of the corresponding IL-18 polypeptide not comprised in the immunocytokine composition, and wherein the immunocytokine composition exhibits a lower  $EC_{50}$  than WT IL-18. In some embodiments, the immunocytokine composition exhibits enhanced anti-tumor growth inhibition compared to the antibody alone. In some embodiments, the immunocytokine composition exhibits enhanced anti-tumor growth inhibition compared to the antibody and the IL-18 polypeptide administered in combination.

In some embodiments, the immunocytokine composition provided herein, further comprising a second linker, wherein the second linker comprises a third point of attachment to the antibody or antigen binding fragment thereof, and a fourth point of attachment to an additional cytokine. In some embodiments, the additional cytokine is selected from a second IL-18 polypeptide, an IL-7 polypeptide, or an IL-2 polypeptide. In some embodiments, the additional cytokine is a second IL-18 polypeptide. In some embodiments, the second IL-18 polypeptide is substantially identical to the IL-18 polypeptide. In some embodiments, the third point of attachment is to a different heavy chain of the antibody or antigen binding fragment thereof from the second point of attachment.

In one aspect, provided herein, is a pharmaceutical composition comprising: an immunocytokine composition as provided herein, and one or more pharmaceutically acceptable carriers or excipients. In some embodiments, the pharmaceutical composition is formulated for parenteral or enteral administration. In some embodiments, the pharmaceutical composition is formulated for intravenous or subcutaneous administration. In some embodiments, the pharmaceutical composition is in a lyophilized form. In some embodiments, the one or more pharmaceutically acceptable carriers or excipients comprises one or more of each of: a carbohydrate, an inorganic salt, an antioxidant, a surfactant, a buffer, or any combination thereof. In some embodiments, the pharmaceutical composition comprises one, two, three, four, five, six, seven, eight, nine, ten, or more excipients.

In one aspect, provided herein, is a method of treating cancer in a subject in need thereof, comprising administering to the subject an effective amount of the immunocytokine composition as provided herein, or a pharmaceutical composition as provided herein. In some embodiments, the

cancer is a melanoma, a lung cancer, a bladder cancer (BC), a microsatellite instability high (MSI-H)/mismatch repair-deficient (dMMR) solid tumor, a tumor mutation burden high (TMB-H) solid tumor, a triple-negative breast cancer (TNBC), a gastric cancer (GC), a cervical cancer (CC), a pleural mesothelioma (PM), classical Hodgkin's lymphoma (cHL), a primary mediastinal large B cell lymphoma (PMBCL), or a combination thereof. In some embodiments, the cancer is a solid cancer consisting of bladder and ureteral cancer, bone cancer, brain and spinal cord cancer, breast cancer, cervical cancer, colorectal cancer, esophageal cancer, eye cancer, gallbladder or bile duct cancer, germ cell tumor, head and neck cancer, kidney cancer, liver cancer, lung cancer, metastatic brain tumor, ovarian cancer, pancreatic cancer, pediatric cancer, peripheral nerve sheath tumor, pituitary cancer, prostate cancer, skin cancer, stomach or gastric cancer, soft tissue cancer, testicular cancer, thyroid cancer, uterine cancer. In some embodiments, the cancer is a carcinoma, and wherein the carcinoma comprises a cutaneous squamous cell carcinoma (CSCC), a urothelial carcinoma (UC), a renal cell carcinoma (RCC), a hepatocellular carcinoma (HCC), a head and neck squamous cell carcinoma (HNSCC), an esophageal squamous cell carcinoma (ESCC), a gastroesophageal junction (GEJ) carcinoma, an endometrial carcinoma (EC), a Merkel cell carcinoma (MCC), or a combination thereof. In some embodiments, the cancer is a leukemia, lymphoma, myeloma, or a combination thereof. In some embodiments, the cancer is leukemia and the leukemia comprises, acute lymphoblastic leukemia, acute myeloid leukemia, chronic lymphocytic leukemia, small lymphocytic lymphoma, or chronic myeloid leukemia. In some embodiments, the cancer is lymphoma and the lymphoma comprises Burkitt lymphoma, chronic lymphocytic leukemia/small lymphocytic lymphoma, cutaneous lymphoma, diffuse large B cell lymphoma, follicular lymphoma, Hodgkin lymphoma, peripheral T cell lymphoma, Waldenstrom macroglobulinemia, lymphoplasmacytic lymphoma, marginal zone lymphoma, B cell cutaneous lymphoma, extranodal natural killer T cell lymphoma, T cell lymphoblastic lymphoma, peripheral T cell lymphoma, or T cell cutaneous lymphoma.

In some embodiments, the method of making an immunocytokine composition comprises a) covalently attaching a reactive group to a specific residue of the antibody or antigen binding fragment thereof, b) contacting the reactive group with a complementary reactive group attached to the IL-18 polypeptide; and c) forming the composition.

In one aspect, provided herein is a method of creating a composition comprising: an IL-18 polypeptide; and an antibody or an antigen binding fragment thereof specific for an immune cell antigen; and a linker, wherein the linker comprises: a first point of attachment to the IL-18 polypeptide; and a second point of attachment to the antibody or antigen binding fragment thereof, the method comprising: a) providing the antibody or antigen binding fragment, wherein the antibody or antigen binding fragment comprises at least one acceptor amino acid residue that is reactive with the linker in the presence of a coupling enzyme; and b) reacting said antibody or antigen binding fragment with the linker comprising a primary amine, wherein the linker comprises a reactive group (R), in the presence of an enzyme capable of causing the formation of a covalent bond between the at least one acceptor amino acid residue and the linker, wherein the covalent bond is not at the R moiety, and wherein the method is performed under conditions sufficient to cause the at least one acceptor amino acid residue to form a covalent bond to the reactive group via the linker, wherein the covalent bond comprises the second point of attachment of the linker. In

some embodiments, the enzyme comprises a transaminase. In some embodiments, the enzyme comprises a transglutaminase.

Additional aspects and advantages of the present disclosure will become readily apparent to those skilled in this art from the following detailed description, wherein only illustrative embodiments of the present disclosure are shown and described. As will be realized, the present disclosure is capable of other and different embodiments, and its several details are capable of modifications in various obvious respects, all without departing from the disclosure. Accordingly, the drawings and description are to be regarded as illustrative in nature, and not as restrictive.

#### INCORPORATION BY REFERENCE

All publications, patents, and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication, patent, or patent application was specifically and individually indicated to be incorporated by reference. To the extent publications and patents or patent applications incorporated by reference contradict the disclosure contained in the specification, the specification is intended to supersede and/or take precedence over any such contradictory material.

#### BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1A illustrates a non-limiting mechanism of action of an anti-PD-1 antibody/IL-18 immunocytokine provided herein, wherein an activated T cell shows enhanced activation through concurrent blockade of PD-1 and stimulation by IL-18.

FIG. 1B illustrates a non-limiting mechanism of action of an anti-PD-L1 antibody/IL-18 immunocytokine provided herein, wherein the immunocytokine disrupts PD-L1/PD-1 interaction and effectuates IL-18R signaling.

FIG. 1C illustrates a non-limiting mechanism of action of an anti-PD-L1 antibody/IL-18 immunocytokine acting on an immune cell expressing both IL-18R and PD-L1.

FIG. 1D illustrates an immunocytokine composition provided herein with two different protein payloads (e.g., IL-18 and IL-2) linked to an antibody (e.g., PD-L1 antibody) acting on an immune cell with simultaneous activation of two different receptors (e.g., IL-18R and IL-2R).

FIG. 2 shows a representative immunocytokine composition with a drug antibody ratio of 1 (DAR1).

FIG. 3 shows an illustration of a site selective introduction of a conjugation handle to an antibody as provided herein.

FIG. 4 illustrates the mechanism of action of IL-18 on IFN $\gamma$  and IL-18BP production, and IL-18 inhibitory activity by IL-18BP.

FIG. 5 shows a schematic of a synthetic route which can be used to synthesize an IL-18 polypeptide for conjugation to an antibody.

FIG. 6 shows a schematic representation of coupling of a bifunctional probe (also referred to as bifunctional linking reagent herein) to an IL-18 polypeptide provided herein.

FIG. 7 shows exemplary reaction scheme which can be used to prepare antibody/IL-18 immunocytokine compositions provided herein with drug antibody ratio of 1 (top) or 2 (bottom).

FIG. 8 shows a reverse phase HPLC chromatogram of a purified IL-18/anti-PD-1 antibody (LZM009) as provided herein with a DAR of 1 attached at residue K248 of the Fc region of the antibody (EU numbering).

FIG. 9 shows a size exclusion HPLC chromatogram of a purified IL-18/anti-PD-1 antibody (LZM009) as provided herein with a DAR of 1 attached at residue K248 of the Fc region of the antibody (EU numbering).

FIG. 10 shows a mass spectrometry trace (Q-TOF) of a purified IL-18/anti-PD-1 antibody (LZM009) as provided herein with a DAR of 1 attached at residue K248 of the Fc region of the antibody (EU numbering).

FIG. 11 shows the RP-HPLC chromatograms of purified composition A (detection: 280 nm).

FIG. 12 shows the Analytical SEC-HPLC chromatograms of purified composition A (detection: 280 nm).

FIG. 13 shows the Q-TOF mass spectrometry deconvolution chromatogram of purified composition A.

FIG. 14A and FIG. 14B show plots measuring ability of the unmodified and of conjugated anti-PD1 antibodies to bind with human PD1/CD279 ligand, with the figure showing ELISA signal on the y-axis and dosage of the biotinylated PD-1 protein on the x-axis.

FIG. 14A Shows the unconjugated reference antibodies. Tested in this figure are Pembrolizumab, LZM-009, Nivolumab, Atezolizumab, Durvalumab, and Avelumab.

FIG. 14B Shows the conjugated antibodies. Tested in this figure are compositions A and composition B.

FIG. 15A and FIG. 15B show plots measuring ability of the unmodified and of conjugated antibodies to bind with human PD-L1/B7-H1 ligand, with the figure showing ELISA signal on the y-axis and dosage of the biotinylated PD-L1 protein on the x-axis.

FIG. 15A shows the unconjugated reference antibodies. Tested in this figure are Pembrolizumab, LZM-009, Nivolumab, Atezolizumab, Durvalumab, and Avelumab.

FIG. 15B shows the conjugated antibodies. Tested in this figure are compositions A and composition B.

FIG. 16 shows plots measuring ability of the unmodified and of conjugated antibodies to bind to human PD-L1/B7-H1 ligand, with the figure showing net BioLayer interferometry shift in nanometer on the y-axis and time of incubation dosage of the biotinylated PD-L1 protein on the x-axis. The figure shows mean ELISA signal on the y-axis and dosage of the human Fc gamma receptors on the x-axis. The unconjugated reference antibodies are Pembrolizumab and LZM-009. The conjugated antibodies tested is Compositions A.

FIG. 17A and FIG. 17B show plots measuring ability of the unmodified and of conjugated anti-PD1 antibodies to interfere with PD1/PDL1 pathway, with the figure showing normalized luminescence intensity of effector cells NFAT-Lucia reporter on the y-axis and dosage of the unmodified and of conjugated anti-PD1 antibodies on the x-axis.

FIG. 17A shows the unconjugated reference antibodies are Pembrolizumab, LZM-009, Nivolumab, Atezolizumab, Durvalumab, and Avelumab.

FIG. 17B shows the conjugated antibodies tested. Tested in this figure are compositions A and composition B.

FIG. 18A shows plots measuring ability of the unmodified and of conjugated antibodies to bind to human Fc gamma receptor I (CD64). The figure shows mean ELISA signal on the y-axis and dosage of the human Fc gamma receptors on the x-axis. The unconjugated reference antibodies are LZM-009 and Atezolizumab. The conjugated antibodies tested are Compositions A and B.

FIG. 18B shows plots measuring ability of the unmodified and of conjugated antibodies to bind to human Fc gamma receptor IIIa (CD16). The figure shows mean ELISA signal on the y-axis and dosage of the human Fc gamma receptors on the x-axis. The unconjugated reference antibodies are

LZM-009 and Atezolizumab. The conjugated antibodies tested are Compositions A and B.

FIG. 19 shows plots measuring ability of the unmodified and of conjugated antibodies to bind to human Fc neonatal receptor. The figure shows mean AlphaLISA signal on the y-axis and dosage of the human Fc neonatal receptor (FcRn) on the x-axis. The unconjugated reference antibodies are LZM-009 and Atezolizumab. The conjugated antibodies tested are Compositions A and B.

FIG. 20 shows plots measuring the levels PD-1 and PD-L1 surface expression on NK92 cells.

FIG. 21 shows plots measuring ability of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines to stimulate the secretion of IFNgamma by NK92 cells. The figure shows mean IFNg alphaLISA signal on the y-axis and dosage of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO: 1), SEQ ID NO: 30, and SEQ ID NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

FIG. 22 shows plots measuring the levels PD-1 and PD-L1 surface expression on KG-1 cells.

FIG. 23 shows plots measuring the ability of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines to stimulate the secretion of IFNgamma by parental PD-1<sup>negative</sup> and by engineered PD-1<sup>positive</sup> KG-1 cells. The figure shows mean IFNg legendplex signal on the y-axis and dosage of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO: 1), SEQ ID NO: 30, and SEQ ID NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

FIG. 24 shows plots measuring the ability of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines to bind to the human IL-18 Binding Protein (IL-18BP). The figure shows mean free IL-18BP AlphaLISA signal on the y-axis and dosage of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO: 1), SEQ ID NO: 30, and SEQ ID NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

FIG. 25 shows plots measuring the ability of the human IL-18 Binding Protein to inhibit the secretion of IFNgamma by NK92 cells stimulated with 2 nM of unconjugated IL-18 variants and corresponding IL-18 immunocytokines. The figure shows mean IFNg alphaLISA signal on the y-axis and dosage of the human IL-18 Binding Protein on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO: 1), SEQ ID NO: 30, and SEQ ID NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

FIG. 26A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition A tested as a single agent at 0.3 and 1 mg/kg as two weekly i.v. injections. (n=9; mean±SEM).

FIG. 26B shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-L1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition B

tested as a single agent at 1 and 3 mg/kg as two weekly i.v. injections. (n=9; mean±SEM).

FIG. 27A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the body weight of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition A tested as a single agent at 0.3 and 1 mg/kg as two weekly i.v. injections. (n=9; mean±SEM).

FIG. 27B shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-L1 antibody on the body weight of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition B tested as a single agent at 1 and 3 mg/kg as two weekly i.v. injections. (n=9; mean±SEM).

FIG. 28A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 1 mg/kg (n=9; mean±SEM).

FIG. 28B shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows the mean tumor volume on day 17 post treatment initiation on the y-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; One-way Anova test \*\*\*P-value<0.001, \*\*P-value<0.01, \*P-value<0.1, ns not significant, TGI: Tumor Growth Inhibition).

FIG. 28C shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows the tumor volume of each individual animal on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 1 mg/kg (n=9; CR: Complete Response).

FIG. 29 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the body weight of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 1 mg/kg (n=9; mean±SEM).

## 11

FIG. 30 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the survival of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 1 mg/kg (n=9; mean±SEM; CR: Complete response).

FIG. 31A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of B16F10 syngeneic melanoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; mean±SEM).

FIG. 31B shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of B16F10 syngeneic melanoma tumors in hPD1 C57BL/6 mice. The figure shows the mean tumor volume on day 10 post treatment initiation on the y-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; TGI: Tumor Growth Inhibition).

FIG. 31C shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of B16F10 syngeneic melanoma tumors in hPD1 C57BL/6 mice. The figure shows the tumor volume of each individual animal on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; CR: Complete Response).

FIG. 32 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the body weight of B16F10 syngeneic melanoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; mean±SEM).

FIG. 33 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the survival of B16F10 syngeneic melanoma tumor-bearing hPD1 C57BL/6 mice. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; mean±SEM; CR: Complete response).

## DETAILED DESCRIPTION

Disclosed herein are antibodies or antigen binding fragments specific for immune cell associated antigens linked to

## 12

IL-18 polypeptides in immunocytokine compositions. In some instances, the immunocytokine compositions provided herein are useful as potent stimulators of one or more immune cell types (e.g., T cells, macrophages, etc.). In some embodiments, the immunocytokine compositions can act by one or more modes of action.

In some embodiments, the antibody of the immunocytokine composition allows for targeting of the immunocytokine composition to an immune cell. In some embodiments, 10 the immunocytokine composition can inhibit an activity of the immune cell associated antigen (e.g., inhibiting a checkpoint interaction such as a PD-1/PD-L1 interaction) through binding to the immune cell associated antigen. In some 15 embodiments, the immunocytokine compositions induce IFN $\gamma$  production in immune cells (e.g., T cells or NK cells). The antibody or antigen binding fragment-IL-18 immunocytokine compositions of the disclosure can have synergistic efficacy and improved tolerability by a subject. In some 20 embodiments, the antibody or antigen binding fragment-IL-18 immunocytokine compositions can significantly reduce the therapeutic dose of the antibody or antigen binding fragment, the IL-18 polypeptide, or both for a subject with a disease, such as a cancer, as compared to a treatment with one or both entities individually or in combination. In some 25 embodiments, the immunocytokine compositions provided herein are associated with fewer side effects than administration of one or both entities individually or in combination, potentially due to the targeting nature of the antibodies for an immune cell.

An exemplary, non-limiting mechanism of action of an immunocytokine provided herein is shown in FIG. 1A. In the exemplary embodiment, the immunocytokine composition comprises an anti-PD-1 antibody as the antibody or antigen binding fragment of the immunocytokine composition. 30 In this embodiment, the anti-PD-1 antibody portion of the immunocytokine selectively binds to PD-1 present on the surface of an activated T cell (e.g., a CD8 $+$  T cell). This binding prevents the checkpoint interaction of PD-1 and PD-L1/2, thus preventing attenuation of activity of the T cell. Concomitantly, the IL-18 portion of the immunocytokine, which is effectively in a high local concentration near the T-cell due to the linkage, further activates the T cell through IL-18R signaling. While the exemplary embodiment shows the mechanism of action of an anti-PD-1 35 antibody, antibodies or antigen binding fragments specific for other immune antigens provided herein can function according to a similar mechanism.

Another exemplary, non-limiting mechanism of action of an anti-PD-L1 antibody/IL-18 immunocytokine is shown in FIG. 1B. In the exemplary embodiment, the anti-PD-L1 antibody portion of the immunocytokine binds to PD-L1 expressed on the surface of a tumor cell. When a T cell comes into contact with the tumor cell, an interaction between PD-1 on the T cell and PD-L1 on the cell is blocked, 40 preventing attenuation of the activity of the T cell. Additionally, the IL-18 portion of the immunocytokine is free to signal through IL-18R on the immune cell, thereby inducing production of IFN $\gamma$  and further activation of the immune cell. Though this exemplary embodiment is demonstrated 45 for PD-L1, other immune antigens provided herein in immunocytokines may display similar mechanisms of action.

Also disclosed herein are methods of manufacturing the immunocytokine compositions provided herein. In some embodiments, the immunocytokine compositions are prepared 50 using chemical linkers which can attach the two moieties of immunocytokine composition to each other at pre-selected sites of each moiety with high fidelity. In some 55

## 13

embodiments, the methods provided herein can be used on a wide variety of antibodies or antigen binding fragments in order to rapidly and easily generate a wide variety of immune antigen specific antibody or antigen binding fragments linked to IL-18 polypeptides as immunocytokines. In some embodiments, the methods can be used on readily commercially available antibodies to allow for rapid linking with IL-18 polypeptides provided herein. One non-limiting illustration of an immunocytokine as provided herein is shown in FIG. 2, which depicts an IL-18 polypeptide linked to an antibody as provided herein with a point of attachment of the linker to the IL-18 polypeptide at a side chain of a residue and to a side chain of a residue in the Fc region.

The following description and examples illustrate embodiments of the present disclosure in detail. It is to be understood that this present disclosure is not limited to the particular embodiments described herein and as such can vary. Those of skill in the art will recognize that there are numerous variations and modifications of this present disclosure, which are encompassed within its scope.

Although various features of the present disclosure may be described in the context of a single embodiment, the features may also be provided separately or in any suitable combination. Conversely, although the present disclosure may be described herein in the context of separate embodiments for clarity, the present disclosure may also be implemented in a single embodiment.

The section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described.

#### Immune Antigen Specific Antibodies Linked to IL-18 Polypeptides as Immunocytokines

Provided herein are antibodies and antigen binding fragments which binds to immune cell associated antigens linked to IL-18 polypeptides as immunocytokine compositions. The immunocytokine compositions provided herein are effective for simultaneously delivering the IL-18 polypeptide and the antibody or antigen binding fragment to a target cell, such as an immune cell. This simultaneous delivery of both agents to the same cell has numerous benefits, including improved IL-18 polypeptide selectivity, enhanced therapeutic potential of the IL-18 polypeptide, and minimized risk of side effects from administering IL-18 therapies. In some embodiments, the immunocytokine compositions act through multiple modes of action, including without limitation disrupting an activity of the immune cell associated antigen (e.g., immune checkpoint evasion) and/or enhanced activation of immune cells in or around a tumor microenvironment.

The immunocytokine compositions provided herein utilize linkers to attach the antibody or antigen binding fragment to the IL-18 polypeptide. In some embodiments, the linkers are attached to each moiety (i.e., the antibody or antigen binding fragment and the IL-18 polypeptide) at specific residues or a specific subset of residues. In some embodiments, the linkers are attached to each moiety in a site-selective manner, such that a population of the immunocytokine compositions is substantially uniform. This can be accomplished in a variety of ways as provided herein, including by site-selectively adding reagents for a conjugation reaction to a moiety to be conjugated, synthesizing or otherwise preparing a moiety to be conjugated with a desired reagent for a conjugation reaction, or a combination of these two approaches. Using these approaches, the sites of attachment (such as specific amino acid residues) of the linker to each moiety can be selected with precision.

## 14

Additionally, these approaches allow a variety of linkers to be employed for the composition which are not limited to amino acid residues as is required for fusion proteins. This combination of linker choice and precision attachment to the moieties allows the linker to also, in some embodiments, perform the function of modulating the activity of one of the moieties, for example if the linker is attached to the IL-18 polypeptide at a position that interacts with a protein which binds to the IL-18 polypeptide (e.g., IL-18 binding protein).

In one aspect, provided herein, is an immunocytokine composition, comprising: an IL-18 polypeptide and an antibody or an antigen binding fragment thereof specific for an immune cell associated antigen. In some embodiments, the immunocytokine composition comprises a linker. In some embodiments, the linker comprises a first point of attachment to the IL-18 polypeptide. In some embodiments, the linker comprises a second point of attachment to the antibody or antigen binding fragment thereof.

#### Immune Cell Specific Antibodies

In some embodiments, an antibody or an antigen binding fragment of the disclosure specifically binds to an immune cell associated antigen. An immune cell associated antigen provided herein is an antigen expressed at elevated levels in cells of an immune cell relative to other cells. In some embodiments, the immune cell associated antigen is expressed at a level of at least 25% greater, at least 30% greater, at least 40% greater, at least 50% greater, at least 60% greater, at least 70% greater, at least 80% greater, at least 90% greater, or at least 100% greater in the immune cell than another cell. In some embodiments, the immune cell associated antigen is expressed at a level of at least 2-fold greater, at least 4-fold greater, at least 6-fold greater, at least 8-fold greater, or at least 10-fold greater in the immune cell than another cell.

An antibody selectively binds or preferentially binds to a target if it binds with greater affinity, avidity, more readily, and/or with greater duration than it binds to other substances. As such, “specific binding” or “preferential binding” does not necessarily require (although it can include) exclusive binding. Generally, but not necessarily, reference to specific binding means preferential binding where the affinity of the antibody, or antigen binding fragment thereof, is at least at least 2-fold greater, at least 3-fold greater, at least 4-fold greater, at least 5-fold greater, at least 6-fold greater, at least 7-fold greater, at least 8-fold greater, at least 9-fold greater, at least 10-fold greater, at least 20-fold greater, at least 30-fold greater, at least 40-fold greater, at least 50-fold greater, at least 60-fold greater, at least 70-fold greater, at least 80-fold greater, at least 90-fold greater, at least 100-fold greater, or at least 1000-fold greater than the affinity of the antibody for unrelated amino acid sequences. In some embodiments, an antibody or an antigen binding fragment of the disclosure can inhibit the action/activity of the substance to which it binds. In some embodiments, an antibody or antigen binding fragment of the disclosure can agonize the action/activity of the substance to which it binds (e.g., an immune cell agonist antibody or antigen binding fragment such as one specific for CD16A, NKG2D, NKp30, or other targets).

As used herein, the term “antibody” refers to an immunoglobulin (Ig), polypeptide, or a protein having a binding domain which is, or is homologous to, an antigen binding domain. The term further includes “antigen binding fragments” and other interchangeable terms for similar binding fragments as described below. Native antibodies and native immunoglobulins (Igs) are generally heterotetrameric glycoproteins of about 150,000 Daltons, composed of two

identical light chains and two identical heavy chains. Each light chain is typically linked to a heavy chain by one covalent disulfide bond, while the number of disulfide linkages varies among the heavy chains of different immunoglobulin isotypes. Each heavy and light chain also has regularly spaced intrachain disulfide bridges. Each heavy chain has at one end a variable domain ("V<sub>H</sub>") followed by a number of constant domains ("C<sub>H</sub>"). Each light chain has a variable domain at one end ("V<sub>L</sub>") and a constant domain ("C<sub>L</sub>") at its other end; the constant domain of the light chain is aligned with the first constant domain of the heavy chain, and the light-chain variable domain is aligned with the variable domain of the heavy chain. Particular amino acid residues are believed to form an interface between the light- and heavy-chain variable domains.

In some instances, an antibody or an antigen binding fragment comprises an isolated antibody or antigen binding fragment, a purified antibody or antigen binding fragment, a recombinant antibody or antigen binding fragment, a modified antibody or antigen binding fragment, or a synthetic antibody or antigen binding fragment.

Antibodies and antigen binding fragments herein can be partly or wholly synthetically produced. An antibody or antigen binding fragment can be a polypeptide or protein having a binding domain which can be, or can be homologous to, an antigen binding domain. In one instance, an antibody or an antigen binding fragment can be produced in an appropriate *in vivo* animal model and then isolated and/or purified.

Depending on the amino acid sequence of the constant domain of its heavy chains, immunoglobulins (Igs) can be assigned to different classes. There are five major classes of immunoglobulins: IgA, IgD, IgE, IgG, and IgM, and several of these may be further divided into subclasses (isotypes), e.g., IgG1, IgG2, IgG3, IgG4, IgA1 and IgA2. An Ig or portion thereof can, in some cases, be a human Ig. In some instances, a C<sub>H</sub>3 domain can be from an immunoglobulin. In some cases, a chain or a part of an antibody or antigen binding fragment, a modified antibody or antigen binding fragment, or a binding agent can be from an Ig. In such cases, an Ig can be IgG, an IgA, an IgD, an IgE, or an IgM, or is derived therefrom. In cases where the Ig is an IgG, it can be a subtype of IgG, wherein subtypes of IgG can include IgG1, an IgG2a, an IgG2b, an IgG3, or an IgG4. In some cases, a C<sub>H</sub>3 domain can be from an immunoglobulin selected from the group consisting of an IgG, an IgA, an IgD, an IgE, and an IgM, or is derived therefrom. In some embodiments, an antibody or antigen binding fragment described herein comprises an IgG or is derived therefrom. In some instances, an antibody or antigen binding fragment comprises an IgG1 or is derived therefrom. In some instances, an antibody or antigen binding fragment comprises an IgG4 or is derived therefrom. In some instances, an antibody or antigen binding fragment comprises an IgG2 or is derived therefrom. In some embodiments, an antibody or antigen binding fragment described herein comprises an IgM, is derived therefrom, or is a monomeric form of IgM. In some embodiments, an antibody or antigen binding fragment described herein comprises an IgE or is derived therefrom. In some embodiments, an antibody or antigen binding fragment described herein comprises an IgD or is derived therefrom. In some embodiments, an antibody or antigen binding fragment described herein comprises an IgA or is derived therefrom.

The "light chains" of antibodies (immunoglobulins) from any vertebrate species can be assigned to one of two clearly

distinct types, called kappa ("κ" or "K") or lambda ("λ"), based on the amino acid sequences of their constant domains.

A "variable region" of an antibody refers to the variable region of the antibody light chain or the variable region of the antibody heavy chain, either alone or in combination. The variable regions of the heavy and light chain each consist of four framework regions (FR) connected by three complementarity determining regions (CDRs) also known as hypervariable regions. The CDRs in each chain are held together in close proximity by the FRs and, with the CDRs from the other chain, contribute to the formation of the antigen binding site of antibodies. There are at least two techniques for determining CDRs: (1) an approach based on cross-species sequence variability (Kabat et al., Sequences of Proteins of Immunological Interest, 5th Ed., 1991, National Institutes of Health, Bethesda Md., pages 647-669; hereafter "Kabat"); and (2) an approach based on crystallographic studies of antigen-antibody complexes (Al-Lazikani et al. (1997) *J. Molec. Biol.* 273:927-948)). As used herein, a CDR may refer to CDRs defined by either approach or by a combination of both approaches.

With respect to antibodies, the term "variable domain" refers to the variable domains of antibodies that are used in the binding and specificity of each particular antibody for its particular antigen. However, the variability is not evenly distributed throughout the variable domains of antibodies. Rather, it is concentrated in three segments called hypervariable regions (also known as CDRs) in both the light chain and the heavy chain variable domains. More highly conserved portions of variable domains are called the "framework regions" or "FRs." The variable domains of unmodified heavy and light chains each contain four FRs (FR1, FR2, FR3, and FR4), largely adopting a β-sheet configuration interspersed with three CDRs which form loops connecting and, in some cases, part of the j-sheet structure. The CDRs in each chain are held together in close proximity by the FRs and, with the CDRs from the other chain, contribute to the formation of the antigen binding site of antibodies (see, Kabat).

The terms "hypervariable region" and "CDR" when used herein, refer to the amino acid residues of an antibody which are responsible for antigen binding. The CDRs comprise amino acid residues from three sequence regions which bind in a complementary manner to an antigen and are known as CDR1, CDR2, and CDR3 for each of the V<sub>H</sub> and V<sub>L</sub> chains. In the light chain variable domain, the CDRs typically correspond to approximately residues 24-34 (CDRL1), 50-56 (CDRL2), and 89-97 (CDRL3), and in the heavy chain variable domain the CDRs typically correspond to approximately residues 31-35 (CDRH1), 50-65 (CDRH2), and 95-102 (CDRH3) according to Kabat et al., *Id.* It is understood that the CDRs of different antibodies may contain insertions, thus the amino acid numbering may differ. The Kabat numbering system accounts for such insertions with a numbering scheme that utilizes letters attached to specific residues (e.g., 27A, 27B, 27C, 27D, 27E, and 27F of CDRL1 in the light chain) to reflect any insertions in the numberings between different antibodies. Alternatively, in the light chain variable domain, the CDRs typically correspond to approximately residues 26-32 (CDRL1), 50-52 (CDRL2), and 91-96 (CDRL3), and in the heavy chain variable domain, the CDRs typically correspond to approximately residues 26-32 (CDRH1), 53-55 (CDRH2), and 96-101 (CDRH3) according to Chothia and Lesk (*J. Mol. Biol.*, 196: 901-917 (1987)).

As used herein, “framework region,” “FW,” or “FR” refers to framework amino acid residues that form a part of the antigen binding pocket or groove. In some embodiments, the framework residues form a loop that is a part of the antigen binding pocket or groove and the amino acids residues in the loop may or may not contact the antigen. Framework regions generally comprise the regions between the CDRs. In the light chain variable domain, the FRs typically correspond to approximately residues 0-23 (FRL1), 35-49 (FRL2), 57-88 (FRL3), and 98-109 and in the heavy chain variable domain the FRs typically correspond to approximately residues 0-30 (FRH1), 36-49 (FRH2), 66-94 (FRH3), and 103-133 according to Kabat et al., Id. As discussed above with the Kabat numbering for the light chain, the heavy chain too accounts for insertions in a similar manner (e.g., 35A, 35B of CDRH1 in the heavy chain). Alternatively, in the light chain variable domain, the FRs typically correspond to approximately residues 0-25 (FRL1), 33-49 (FRL2) 53-90 (FRL3), and 97-109 (FRL4), and in the heavy chain variable domain, the FRs typically correspond to approximately residues 0-25 (FRH1), 33-52 (FRH2), 56-95 (FRH3), and 102-113 (FRH4) according to Chothia and Lesk, Id. The loop amino acids of a FR can be assessed and determined by inspection of the three-dimensional structure of an antibody heavy chain and/or antibody light chain. The three-dimensional structure can be analyzed for solvent accessible amino acid positions as such positions are likely to form a loop and/or provide antigen contact in an antibody variable domain. Some of the solvent accessible positions can tolerate amino acid sequence diversity and others (e.g., structural positions) are, generally, less diversified. The three-dimensional structure of the antibody variable domain can be derived from a crystal structure or protein modeling.

In the present disclosure, the following abbreviations (in the parentheses) are used in accordance with the customs, as necessary: heavy chain (H chain), light chain (L chain), heavy chain variable region (VH), light chain variable region (VL), complementarity determining region (CDR), first complementarity determining region (CDR1), second complementarity determining region (CDR2), third complementarity determining region (CDR3), heavy chain first complementarity determining region (VH CDR1), heavy chain second complementarity determining region (VH CDR2), heavy chain third complementarity determining region (VH CDR3), light chain first complementarity determining region (VL CDR1), light chain second complementarity determining region (VL CDR2), and light chain third complementarity determining region (VL CDR3).

The term “Fc region” is used to define a C-terminal region of an immunoglobulin heavy chain. The “Fc region” may be a native sequence Fc region or a variant Fc region. Although the boundaries of the Fc region of an immunoglobulin heavy chain might vary, the human IgG heavy chain Fc region is generally defined to stretch from an amino acid residue at position Cys226, or from Pro230, to the carboxyl-terminus thereof. The numbering of the residues in the Fc region is that of the EU index as in Kabat. The Fc region of an immunoglobulin generally comprises two constant domains,  $C_{H2}$  and  $C_{H3}$ .

“Antibodies” useful in the present disclosure encompass, but are not limited to, monoclonal antibodies, polyclonal antibodies, chimeric antibodies, bispecific antibodies, grafted antibodies, multispecific antibodies, heteroconjugate antibodies, humanized antibodies, human antibodies, deimmunized antibodies, mutants thereof, fusions thereof, immunoconjugates thereof, antigen binding fragments thereof,

and/or any other modified configuration of the immunoglobulin molecule that comprises an antigen recognition site of the required specificity, including glycosylation variants of antibodies, amino acid sequence variants of antibodies, and covalently modified antibodies. In certain embodiments of the methods and conjugates provided herein, the antibody requires an Fc region to enable attachment of a linker between the antibody and the protein (e.g., attachment of the linker using an affinity peptide, such as in AJICAP™ technology).

In some instances, an antibody is a monoclonal antibody. As used herein, a “monoclonal antibody” refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally-occurring mutations that may be present in minor amounts. In contrast to polyclonal antibody preparations, which typically include different antibodies directed against different determinants (epitopes), each monoclonal antibody is directed against a single determinant on the antigen (epitope). The modifier “monoclonal” indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies and is not to be construed as requiring production of the antibody by any particular method.

In some instances, an antibody is a humanized antibody. As used herein, “humanized” antibodies refer to forms of non-human (e.g., murine) antibodies that are specific chimeric immunoglobulins, immunoglobulin chains, or fragments thereof that contain minimal sequence derived from non-human immunoglobulin. For the most part, humanized antibodies are human immunoglobulins (recipient antibody) in which residues from a complementarity determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat, or rabbit having the desired specificity, affinity, and biological activity. In some instances, Fv framework region (FR) residues of the human immunoglobulin are replaced by corresponding non-human residues. Furthermore, the humanized antibody may comprise residues that are found neither in the recipient antibody nor in the imported CDR or framework sequences but are included to further refine and optimize antibody performance. In general, a humanized antibody comprises 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 FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region or domain (Fc), typically that of a human immunoglobulin. Antibodies may have Fc regions modified as described in, for example, WO 99/58572. Other forms of humanized antibodies have one or more CDRs (one, two, three, four, five, or six) which are altered with respect to the original antibody, which are also termed one or more CDRs “derived from” one or more CDRs from the original antibody.

If needed, an antibody or an antigen binding fragment described herein can be assessed for immunogenicity and, as needed, be deimmunized (i.e., the antibody is made less immunoreactive by altering one or more T cell epitopes). As used herein, a “deimmunized antibody” means that one or more T cell epitopes in an antibody sequence have been modified such that a T cell response after administration of the antibody to a subject is reduced compared to an antibody that has not been deimmunized. Analysis of immunogenicity

and T-cell epitopes present in the antibodies and antigen binding fragments described herein can be carried out via the use of software and specific databases. Exemplary software and databases include iTope™ developed by Antitope of Cambridge, England. iTope™, is an in silico technology for analysis of peptide binding to human MHC class II alleles. The iTope™ software predicts peptide binding to human MHC class II alleles and thereby provides an initial screen for the location of such "potential T cell epitopes." iTope™ software predicts favorable interactions between amino acid side chains of a peptide and specific binding pockets within the binding grooves of 34 human MHC class II alleles. The location of key binding residues is achieved by the in silico generation of 9mer peptides that overlap by one amino acid spanning the test antibody variable region sequence. Each 9mer peptide can be tested against each of the 34 MHC class II allotypes and scored based on their potential "fit" and interactions with the MHC class II binding groove. Peptides that produce a high mean binding score (>0.55 in the iTope™ scoring function) against >50% of the MHC class II alleles are considered as potential T cell epitopes. In such regions, the core 9 amino acid sequence for peptide binding within the MHC class II groove is analyzed to determine the MHC class II pocket residues (P1, P4, P6, P7, and P9) and the possible T cell receptor (TCR) contact residues (P-1, P2, P3, P5, P8). After identification of any T-cell epitopes, amino acid residue changes, substitutions, additions, and/or deletions can be introduced to remove the identified T-cell epitope. Such changes can be made so as to preserve antibody structure and function while still removing the identified epitope. Exemplary changes can include, but are not limited to, conservative amino acid changes.

An antibody can be a human antibody. As used herein, a "human antibody" means an antibody having an amino acid sequence corresponding to that of an antibody produced by a human and/or that has been made using any suitable technique for making human antibodies. This definition of a human antibody includes antibodies comprising at least one human heavy chain polypeptide or at least one human light chain polypeptide. One such example is an antibody comprising murine light chain and human heavy chain polypeptides. In one embodiment, the human antibody is selected from a phage library, where that phage library expresses human antibodies. Human antibodies can also be made by introducing human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Alternatively, the human antibody may be prepared by immortalizing human B lymphocytes that produce an antibody directed against a target antigen (such B lymphocytes may be recovered from an individual or may have been immunized in vitro).

Any of the antibodies herein can be bispecific. Bispecific antibodies are antibodies that have binding specificities for at least two different antigens and can be prepared using the antibodies disclosed herein. Traditionally, the recombinant production of bispecific antibodies was based on the co-expression of two immunoglobulin heavy chain-light chain pairs, with the two heavy chains having different specificities. Bispecific antibodies can be composed of a hybrid immunoglobulin heavy chain with a first binding specificity in one arm, and a hybrid immunoglobulin heavy chain-light chain pair (providing a second binding specificity) in the other arm. This asymmetric structure, with an immunoglobulin light chain in only one half of the bispecific

molecule, facilitates the separation of the desired bispecific compound from unwanted immunoglobulin chain combinations.

According to one approach to making bispecific antibodies, antibody variable domains with the desired binding specificities (antibody-antigen combining sites) are fused to immunoglobulin constant domain sequences. The fusion can be with an immunoglobulin heavy chain constant domain, comprising at least part of the hinge, CH2 and CH3 regions. 10 The first heavy chain constant region (CH1), containing the site necessary for light chain binding, can be present in at least one of the fusions. DNAs encoding the immunoglobulin heavy chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and 15 are co-transfected into a suitable host organism. This provides for great flexibility in adjusting the mutual proportions of the three polypeptide fragments in embodiments when unequal ratios of the three polypeptide chains used in the construction provide the optimum yields. It is, however, 20 possible to insert the coding sequences for two or all three polypeptide chains in one expression vector when the expression of at least two polypeptide chains in equal ratios results in high yields or when the ratios are of no particular significance.

25 In some instances, an antibody herein is a chimeric antibody. "Chimeric" forms of non-human (e.g., murine) antibodies include chimeric antibodies which contain minimal sequence derived from a non-human Ig. For the most part, chimeric antibodies are murine antibodies in which at 30 least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin, is inserted in place of the murine Fc. Chimeric or hybrid antibodies also may be prepared in vitro using suitable methods of synthetic protein chemistry, including those involving cross-linking agents.

35 Provided herein are antibodies and antigen binding fragments thereof, modified antibodies and antigen binding fragments thereof, and binding agents that specifically bind to one or more epitopes on one or more target antigens. In 40 one instance, a binding agent selectively binds to an epitope on a single antigen. In another instance, a binding agent is bivalent and either selectively binds to two distinct epitopes on a single antigen or binds to two distinct epitopes on two distinct antigens. In another instance, a binding agent is 45 multivalent (i.e., trivalent, quatrvivalent, etc.) and the binding agent binds to three or more distinct epitopes on a single antigen or binds to three or more distinct epitopes on two or more (multiple) antigens.

50 Antigen binding fragments of any of the antibodies herein are also contemplated. The terms "antigen binding portion of an antibody," "antigen binding domain," "antibody fragment," or a "functional fragment of an antibody" are used interchangeably herein to refer to one or more fragments of an antibody that retain the ability to specifically bind to an antigen. Representative antigen binding fragments include, but are not limited to, a Fab, a Fab', a F(ab')<sub>2</sub>, a bispecific F(ab')<sub>2</sub>, a trispecific F(ab')<sub>2</sub>, a variable fragment (Fv), a single chain variable fragment (scFv), a dsFv, a bispecific scFv, a variable heavy domain, a variable light domain, a variable NAR domain, bispecific scFv, a minibody, a diabody, a bispecific diabody, triabody, a tetrabody, a minibody, a maxibody, a camelid, a VH, a minibody, an intrabody, fusion proteins comprising an antibody portion (e.g., a domain antibody), a single chain binding polypeptide, a scFv-Fc, a Fab-Fc, a bispecific T cell engager (BiTE; two scFvs produced as a single polypeptide chain, where each scFv comprises an amino acid sequences a combination of

21

CDRs or a combination of VL/VL described herein), a tetravalent tandem diabody (TandAb; an antibody fragment that is produced as a non-covalent homodimer folder in a head-to-tail arrangement, e.g., a TandAb comprising an scFv, where the scFv comprises an amino acid sequences a combination of CDRs or a combination of VL/VL described herein), a Dual-Affinity Re-targeting Antibody (DART; different scFvs joined by a stabilizing interchain disulphide bond), a bispecific antibody (bscAb; two single-chain Fv fragments joined via a glycine-serine linker), a single domain antibody (sdAb), a fusion protein, a bispecific disulfide-stabilized Fv antibody fragment (dsFv-dsFv'; two different disulfide-stabilized Fv antibody fragments connected by flexible linker peptides). In certain embodiments of the invention, a full length antibody (e.g., an antigen binding fragment and an Fc region) are preferred.

Heteroconjugate polypeptides comprising two covalently joined antibodies or antigen binding fragments of antibodies are also within the scope of the disclosure. Suitable linkers may be used to multimerize binding agents. Non-limiting examples of linking peptides include, but are not limited to, (GS)<sub>n</sub> (SEQ ID NO: 224), (GGS)<sub>n</sub> (SEQ ID NO: 225), (GGGS)<sub>n</sub> (SEQ ID NO: 226), (GGSG)<sub>n</sub> (SEQ ID NO: 227), or (GGSGG)<sub>n</sub> (SEQ ID NO: 228), (GGGGS)<sub>n</sub> (SEQ ID NO: 229), wherein n is 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10. For example, a linking peptide can be (GGGGS)<sub>3</sub> (SEQ ID NO: 230) or (GGGGS)<sub>4</sub> (SEQ ID NO: 231). In some embodiments, a linking peptide bridges approximately 3.5 nm between the carboxy terminus of one variable region and the amino terminus of the other variable region. Linkers of other sequences have been designed and used. Linkers can in turn be modified for additional functions, such as attachment of drugs or attachment to solid supports.

As used herein, the term "avidity" refers to the resistance of a complex of two or more agents to dissociation after dilution. Apparent affinities can be determined by methods such as an enzyme-linked immunosorbent assay (ELISA) or any other suitable technique. Avidities can be determined by methods such as a Scatchard analysis or any other suitable technique.

As used herein, the term "affinity" refers to the equilibrium constant for the reversible binding of two agents and is expressed as K<sub>D</sub>. The binding affinity (K<sub>D</sub>) of an antibody or antigen binding fragment herein can be less than 500 nM, 475 nM, 450 nM, 425 nM, 400 nM, 375 nM, 350 nM, 325 nM, 300 nM, 275 nM, 250 nM, 225 nM, 200 nM, 175 nM, 150 nM, 125 nM, 100 nM, 90 nM, 80 nM, 70 nM, 50 nM, 50 nM, 49 nM, 48 nM, 47 nM, 46 nM, 45 nM, 44 nM, 43 nM, 42 nM, 41 nM, 40 nM, 39 nM, 38 nM, 37 nM, 36 nM, 35 nM, 34 nM, 33 nM, 32 nM, 31 nM, 30 nM, 29 nM, 28 nM, 27 nM, 26 nM, 25 nM, 24 nM, 23 nM, 22 nM, 21 nM, 20 nM, 19 nM, 18 nM, 17 nM, 16 nM, 15 nM, 14 nM, 13 nM, 12 nM, 11 nM, 10 nM, 9 nM, 8 nM, 7 nM, 6 nM, 5 nM, 4 nM, 3 nM, 2 nM, 1 nM, 990 pM, 980 pM, 970 pM, 960 pM, 950 pM, 940 pM, 930 pM, 920 pM, 910 pM, 900 pM, 890 pM, 880 pM, 870 pM, 860 pM, 850 pM, 840 pM, 830 pM, 820 pM, 810 pM, 800 pM, 790 pM, 780 pM, 770 pM, 760 pM, 750 pM, 740 pM, 730 pM, 720 pM, 710 pM, 700 pM, 690 pM, 680 pM, 670 pM, 660 pM, 650 pM, 640 pM, 630 pM, 620 pM, 610 pM, 600 pM, 590 pM, 580 pM, 570 pM, 560 pM, 550 pM, 540 pM, 530 pM, 520 pM, 510 pM, 500 pM, 490 pM, 480 pM, 470 pM, 460 pM, 450 pM, 440 pM, 430 pM, 420 pM, 410 pM, 400 pM, 390 pM, 380 pM, 370 pM, 360 pM, 350 pM, 340 pM, 330 pM, 320 pM, 310 pM, 300 pM, 290 pM, 280 pM, 270 pM, 260 pM, 250 pM, 240 pM, 230 pM, 220 pM, 210 pM, 200 pM, 190 pM, 180 pM, 170 pM, or any integer therebetween. Binding affinity

22

may be determined using surface plasmon resonance (SPR), KINEXA® Biosensor, scintillation proximity assays, enzyme linked immunosorbent assay (ELISA), ORIGEN immunoassay (IGEN), fluorescence quenching, fluorescence transfer, yeast display, or any combination thereof. Binding affinity may also be screened using a suitable bioassay.

Also provided herein are affinity matured antibodies. The following methods may be used for adjusting the affinity of an antibody and for characterizing a CDR. One way of characterizing a CDR of an antibody and/or altering (such as improving) the binding affinity of a polypeptide, such as an antibody, is termed "library scanning mutagenesis." Generally, library scanning mutagenesis works as follows. One or more amino acid position in the CDR is replaced with two or more (such as 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20) amino acids. This generates small libraries of clones (in some embodiments, one for every amino acid position that is analyzed), each with a complexity of two or more members (if two or more amino acids are substituted at every position). Generally, the library also includes a clone comprising the native (unsubstituted) amino acid. A small number of clones, for example, about 20-80 clones (depending on the complexity of the library), from each library can be screened for binding specificity or affinity to the target polypeptide (or other binding target), and candidates with increased, the same, decreased, or no binding are identified. Binding affinity may be determined using Biacore surface plasmon resonance analysis, which detects differences in binding affinity of about 2-fold or greater.

In some instances, an antibody or antigen binding fragment is bispecific or multispecific and can specifically bind to more than one antigen. In some cases, such a bispecific or multispecific antibody or antigen binding fragment can specifically bind to 2 or more different antigens. In some cases, a bispecific antibody or antigen binding fragment can be a bivalent antibody or antigen binding fragment. In some cases, a multi specific antibody or antigen binding fragment can be a bivalent antibody or antigen binding fragment, a trivalent antibody or antigen binding fragment, or a quadrivalent antibody or antigen binding fragment.

An antibody or antigen binding fragment described herein can be isolated, purified, recombinant, or synthetic.

It is contemplated that generic or biosimilar versions of the named antibodies herein which share the same amino acid sequence as the indicated antibodies are also encompassed when the name of the antibody is used.

The antibodies described herein may be made by any suitable method. Antibodies can often be produced in large quantities, particularly when utilizing high level expression vectors.

In one embodiment, an antibody or an antigen binding fragment of the disclosure comprises a fusion protein or a peptide immunotherapeutic agent.

#### Immune Cell Associated Antigen Specific Antibodies

In some embodiments, the antibody or antigen binding fragment thereof is specific for an immune cell associated antigen. In some embodiments, the immune cell associated antigen is associated with an immune cell subtype (e.g., lymphocyte, neutrophil, macrophage, etc.). In some embodiments, the immune cell associated antigen is associated with a T cell, a monocyte, and/or a natural killer (NK) cell. In embodiments, the immune cell antigen is associated with a T cell. In some embodiments, the immune cell antigen is associated with an effector T cell, a cytotoxic T cell, a helper T cell, a regulatory T cell, and/or a memory T cell.

In some embodiments, the immune cell associated antigen is an immune checkpoint molecule. In some embodiments, the immune cell associated antigen is a costimulatory antigen. In some embodiments, the immune cell associated antigen is a macrophage cell surface antigen. In some embodiments, the immune cell associated antigen is an NK cell surface antigen. In some embodiments, the immune cell associated antigen is a T cell surface antigen (e.g., CD8A, CD8B).

In some embodiments, the immune cell associated antigen is 4-IBB, B7-H3, B7-H4, BTLA, CD3, CCR8, CD8A, CD8B, CD16A, CD27, CD28, CD33, CD38, CD39, CD40, CD47, CD70, CD80, CD86, CD96, CD163, CLEC-1, CLEVER-1, CTLA-4, D40, GITR, ICOS, ILT2/3/4, LAG-3, MHC<sub>I</sub>, MHC<sub>II</sub>, NKG2A, NKG2D, NKp30, NKp44, NKp46, OX40, PD-1, PD-L1, PD-L2, PSGL-1, SIGLEC-9, SIGLEC-15, SIRP- $\alpha$ , TCR, TIGIT, TIM-3, VISTA, or VSIG4. In some embodiments, the immune cell associated antigen is PD-1. In some embodiments, the immune cell associated antigen is CCR8, CD8A, CD8B, CD16A, CD96, CD226, CTLA-4, ICOS, LAG-3, NKG2A, NKG2D, NKp30, NKp44, NKp46, PD-1, PD-L1, TIGIT, or TIM-3.

In some embodiments, the immune cell associated antigen is 4-1BB. In some embodiments, the immune cell associated antigen is B7-H3. In some embodiments, the immune cell associated antigen is B7-H4. In some embodiments, the immune cell associated antigen is BTLA. In some embodiments, the immune cell associated antigen is CD3. In some embodiments, the immune cell associated antigen is CCR8. In some embodiments, the immune cell associated antigen is CD8A. In some embodiments, the immune cell associated antigen is CD8B. In some embodiments, the immune cell associated antigen is CD16A. In some embodiments, the immune cell associated antigen is CD27. In some embodiments, the immune cell associated antigen is CD33. In some embodiments, the immune cell associated antigen is CD38. In some embodiments, the immune cell associated antigen is CD39. In some embodiments, the immune cell associated antigen is CD40. In some embodiments, the immune cell associated antigen is CD47. In some embodiments, the immune cell associated antigen is CD80. In some embodiments, the immune cell associated antigen is CD86. In some embodiments, the immune cell associated antigen is CD96. In some embodiments, the immune cell associated antigen is CD163. In some embodiments, the immune cell associated antigen is CLEC-1. In some embodiments, the immune cell associated antigen is CLEVER-1. In some embodiments, the immune cell associated antigen is CTLA4. In some embodiments, the immune cell associated antigen is D40. In some embodiments, the immune cell associated antigen is GITR. In some embodiments, the immune cell associated antigen is ICOS. In some embodiments, the immune cell associated antigen is ILT2/3/4. In some embodiments, the immune cell associated antigen is LAG-3. In some embodiments, the immune cell associated antigen is MHC<sub>I</sub>. In some embodiments, the immune cell associated antigen is MHC<sub>II</sub>. In some embodiments, the immune cell associated antigen is NKG2A. In some embodiments, the immune cell associated antigen is NKp30. In some embodiments, the immune cell associated antigen is NKp44. In some embodiments, the immune cell associated antigen is NKp46. In some embodiments, the immune cell associated antigen is OX40. In some embodiments, the immune cell associated antigen is PD-1. In some embodiments, the immune cell associated antigen is PD-L1. In some embodiments, the immune cell associated antigen is PD-L2. In some embodiments, the immune cell associated antigen is PSGL-1. In some embodiments, the

immune cell associated antigen is SIGLEC-9. In some embodiments, the immune cell associated antigen is SIGLEC-15. In some embodiments, the immune cell associated antigen is SIRP- $\square$ . In some embodiments, the immune cell associated antigen is TCR. In some embodiments, the immune cell associated antigen is TIGIT. In some embodiments, the immune cell associated antigen is TIM-3. In some embodiments, the immune cell associated antigen is VISTA. In some embodiments, the immune cell associated antigen is VSIG4.

In some embodiments, the antibody or antigen binding fragment thereof is an anti-PD-1 antibody or antigen binding fragment. Programmed cell death protein 1 (also known as PD-1 and CD279), is a cell surface receptor that plays a role in down-regulating the immune system and promoting self-tolerance by suppressing T cell inflammatory activity. PD-1 is an immune cell inhibitory molecule that is expressed on activated B cells, T cells, and myeloid cells. PD-1 represents an immune checkpoint and guards against autoimmunity via a dual mechanism of promoting apoptosis (programmed cell death) in antigen-specific T-cells in lymph nodes while reducing apoptosis in regulatory T cells. PD-1 is a member of the CD28/CTLA-4/ICOS costimulatory receptor family that delivers negative signals that affect T and B cell immunity. PD-1 is monomeric both in solution as well as on cell surface, in contrast to CTLA-4 and other family members that are all disulfide-linked homodimers. Signaling through the PD-1 inhibitory receptor upon binding its ligand, PD-L1, suppresses immune responses against autoantigens and tumors and plays a role in the maintenance of peripheral immune tolerance. The interaction between PD-1 and PD-L1 results in a decrease in tumor infiltrating lymphocytes, a decrease in T cell receptor mediated proliferation, and immune evasion by the cancerous cells. A non-limiting, exemplary, human PD-1 amino acid sequence is

MQIPQAPWPVVWAVLQLGWRPGWFLD-  
SPDRPWNPPFTSPALLVVTEGDNATFTCSFSN  
TSESFVLNWYRMSPSNQTDKLAAPFEDR-  
SQPGQDCRFRVTQLPNGRDFHMSVVRARRN DSG-  
TYLCGAISLAPKAQIKESLRAELRVTERRAEVP-  
TAHPSPSPRPGQFQTLVVGVGG  
LLGSLVLLVVWLVAVICSRAARGTI-  
GARRTGQPLKEDPSAVPVF SVDYGELDFQWREKTP  
EPPVPCVPEQTEYATIVFPMSGTSSPARRGSADGPR-  
SAQPLRPEDGHCSWPL (SEQ ID NO: 331).

In one embodiment, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment of the disclosure comprises a combination of a heavy chain variable region (VH) and a light chain variable region (VL) described herein. In another embodiment, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment of the disclosure comprises a combination of complementarity determining regions (VH CDR1, VH CDR2, VH CDR3, VL CDR1, VL CDR2, and VL CDR3) described herein. In one embodiment, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment of the disclosure comprises a modified Tislelizumab, Baizean, 0KV0411B3N, BGB-A317, hu317-1/IgG4mt2, Sintilimab, Tyvyt, IBI-308, Toripalimab, TeRuiPuLi, Terepril, Tuoyi, JS-001, TAB-001, Camrelizumab, HR-301210, INCSHR-01210, SHR-1210, Cemiplimab, Cemiplimab-rwlc, LIBTAYO®, 6QVLO57INT, H4H7798N, REGN-2810, SAR-439684, Avelumab, BAVENCIO®, 451238, KXG2PJ551I, MSB-0010682, MSB-0010718C, PF-06834635, Durvalumab, IMFINZI®, 28x28x90 KV, MEDI-4736, Lambrolizumab, Pembrolizumab, KEYTRUDA®, MK-3475, SCH-900475, h409A11, Nivolumab, Nivolumab BMS, OPDIVO®, BMS-936558,

MDX-1106, ONO-4538, Prolgolimab, Forteca, BCD-100, Penpulimab, AK-105, Zimberelimab, AB-122, GLS-010, WBP-3055, Balstilimab, 1Q2QT5M7EO, AGEN-2034, AGEN-2034w, Genolizumab, Geptanolimab, APL-501, CBT-501, GB-226, Dostarlimab, ANB-011, GSK-4057190A, P0GVQ9A4S5, TSR-042, WBP-285, Serplulimab, HLX-10, CS-1003, Retifanlimab, 2Y3T5IF0IZ, INCMGA-00012, INCMGA-0012, MGA-012, Sasanlimab, LZZOIC2EW, PF-06801591, RN-888, Spartalizumab, NVP-LZV-184, PDR-001, QOG25L6Z8Z, Relatlimab/nivolumab, BMS-986213, Cetrelimab, JNJ-3283, JNJ-63723283, LYK98WP91F, Tebotelimab, MGD-013, BCD-217, BAT-1306, HX-008, MEDI-5752, JTX-4014, Cadonilimab, AK-104, BI-754091, Pidilizumab, CT-011, MDV-9300, YBL-006, AMG-256, RG-6279, RO-7284755, BH-2950, IBI-315, RG-6139, RO-7247669, ONO-4685, AK-112, 609-A, LY-3434172, T-3011, MAX-10181, AMG-404, IBI-318, MGD-019, INCB-086550, ONCR-177, LY-3462817, RG-7769, RO-7121661, F-520, XmAb-23104, Pd-1-pik, SG-001, S-95016, Sym-021, LZM-009, Budigalimab, 6VDO4TY3OO, ABBV-181, PR-1648817, CC-90006, XmAb-20717, 2661380, AMP-224, B7-DCIg, EMB-02, ANB-030, PRS-332, [89Zr]Deferoxamido-pembrolizumab, 89Zr-Df-Pembrolizumab, [89Zr]Df-Pembrolizumab, STI-1110, STI-A1110, CX-188, mPD-1 Pb-Tx, MCLA-134, 244C8, ENUM 224C8, ENUM C8, 388D4, ENUM 388D4, ENUM D4, MEDIO680, or AMP-514.

In some embodiments, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment of the disclosure comprises a Tislelizumab, Sintilimab, Toripalimab, Terepril, Camrelizumab, Cemiplimab, Pembrolizumab Nivolumab, Prolgolimab, Penpulimab, Zimberelimab, Balstilimab, Genolizumab, Geptanolimab, Dostarlimab, Serplulimab, Retifanlimab, Sasanlimab, Spartalizumab, Cetrelimab, Tebotelimab, Cadonilimab, A Pidilizumab, LZM-009, or Budigalimab. In one embodiment, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment of the disclosure comprises a modified Tislelizumab, Sintilimab, Toripalimab, Terepril, Camrelizumab, Cemiplimab, Pembrolizumab Nivolumab, Prolgolimab, Penpulimab, Zimberelimab, Balstilimab, Genolizumab, Geptanolimab, Dostarlimab, Serplulimab, Retifanlimab, Sasanlimab, Spartalizumab, Cetrelimab, Tebotelimab, Cadonilimab, A Pidilizumab, LZM-009, or Budigalimab.

In some embodiments, the anti-PD-1 polypeptide is Nivolumab, Pembrolizumab, LZM-009, Dostarlimab, Sintilimab, Spartalizumab, Tislelizumab, or Cemiplimab. In some embodiment, the anti-PD-1 polypeptide is Dostarlimab, Sintilimab, Spartalizumab, or Tislelizumab. In some embodiments, the anti-PD-1 polypeptide is Nivolumab, Pembrolizumab, LZM-009, or Cemiplimab.

In some embodiments, the anti-PD-1 antibody is Pembrolizumab. In some embodiments, the anti-PD-1 antibody is modified Pembrolizumab.

In some embodiments, the anti-PD-1 antibody is a biosimilar of Tislelizumab, Sintilimab, Toripalimab, Terepril, Camrelizumab, Cemiplimab, Pembrolizumab Nivolumab, Prolgolimab, Penpulimab, Zimberelimab, Balstilimab, Genolizumab, Geptanolimab, Dostarlimab, Serplulimab, Retifanlimab, Sasanlimab, Spartalizumab, Cetrelimab, Tebotelimab, Cadonilimab, A Pidilizumab, LZM-009, or Budigalimab. In some embodiments, the anti-PD-1 antibody is a biosimilar of any one of the antibodies provided herein.

TABLE 1 provides the sequences of exemplary anti-PD-1 antibodies and anti-PD-1 antigen binding fragments that can be modified to prepare anti-PD-1 immunoconjugates. TABLE 1 also shows provides combinations of CDRs that

can be utilized in a modified anti-PD-1 immunoconjugate. Reference to an anti-PD-1 antibody herein may alternatively refer to an anti-PD-1 antigen binding fragment.

In some instances, the SEQ ID NOs listed in Table 1 contain full-length heavy or light chains of the indicated antibodies with the VH or VL respectively indicated in bold. Where there is a reference herein to a VH or VL of a SEQ ID NO in Table 1 which contains a full-length heavy or light chain, it is intended to reference the bolded portion of the sequence. For example, reference to “a VH having an amino acid sequence shown in SEQ ID NO: 332” refers to the bolded portion of SEQ ID NO: 332 in Table 1.

An anti-PD-1 antibody or an anti-PD-1 antigen binding fragment can comprise a VH having an amino acid sequence of any one of SEQ ID NOS: 332, 334, 336, 338, 340, 346, 348, 350, 352, 354, 356, 358, 360, 362, 364, 366, 368, 370, 372, 374, 376, and 378. An anti-PD-1 antibody or an anti-PD-1 antigen binding fragment can comprise a VL having an amino acid sequence of any one of SEQ ID NOS: 333, 335, 337, 339, 341, 347, 349, 351, 353, 355, 357, 359, 361, 363, 365, 367, 369, 371, 373, 375, 377, and 379.

An anti-PD-1 antibody or an anti-PD-1 antigen binding fragment can comprise a heavy chain or VH having an amino acid sequence of any one of SEQ ID NOS: 332, 334, 336, 338, 340, 346, 348, 350, 352, 354, 356, 358, 360, 362, 364, 366, 368, 370, 372, 374, 376, and 378, or a portion corresponding to a VH thereof. An anti-PD-1 antibody or an anti-PD-1 antigen binding fragment can comprise a light chain or VL having an amino acid sequence of any one of SEQ ID NOS: 333, 335, 337, 339, 341, 347, 349, 351, 353, 355, 357, 359, 361, 363, 365, 367, 369, 371, 373, 375, 377, and 379, or a portion corresponding to a VL thereof.

In one instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 332, and a VL having an amino acid sequence shown in SEQ ID NO: 333. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 334, and a VL having an amino acid sequence shown in SEQ ID NO: 335. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 336, and a VL having an amino acid sequence shown in SEQ ID NO: 337. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 338, and a VL having an amino acid sequence shown in SEQ ID NO: 339. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 340, and a VL having an amino acid sequence shown in SEQ ID NO: 341. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 346, and a VL having an amino acid sequence shown in SEQ ID NO: 347. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 348, and a VL having an amino acid sequence shown in SEQ ID NO: 349. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 350, and a VL having an amino acid sequence shown in SEQ ID NO: 351.

TABLE 1 also shows provides combinations of CDRs that

In one instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence of SEQ ID NO: 332, and a VL having an amino acid sequence of SEQ ID NO: 333. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence of SEQ ID NO: 334, and a VL having an amino acid sequence of SEQ ID NO: 335. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence of SEQ ID NO: 336, and a VL having an amino acid sequence of SEQ ID NO: 337. In another instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH having an amino acid sequence of SEQ ID NO:

29

binding fragment comprises a VH having an amino acid sequence of SEQ ID NO: 378, and a VL having an amino acid sequence of SEQ ID NO: 379.

In one instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH CDR1 having an amino acid sequence of SEQ ID NO: 380, a VH CDR2 having an amino acid sequence of SEQ ID NO: 381, a VH CDR3 having an amino acid sequence of SEQ ID NO: 382, VL CDR1 having an amino acid sequence of SEQ ID NO: 383, a VL CDR2 having an amino acid sequence of SEQ ID NO: 384, and a VL CDR3 having an amino acid sequence of SEQ ID NO: 385. In one instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH CDR1 having an amino acid sequence of SEQ ID NO: 386, a VH CDR2 having an amino acid sequence of SEQ ID NO: 387, a VH CDR3 having an amino acid sequence of SEQ ID NO: 388, VL CDR1 having an amino acid sequence of SEQ ID NO: 389, a VL CDR2 having an amino acid sequence of SEQ ID NO: 390, and a VL CDR3 having an amino acid sequence of SEQ ID NO: 391. In one instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH CDR1 having an amino acid sequence of SEQ ID NO: 392, a VH CDR2 having an amino acid sequence of SEQ ID NO: 393, a VH CDR3 having an amino acid sequence of SEQ ID NO: 394, VL CDR1 having an amino acid sequence of SEQ ID NO: 395, a VL CDR2 having an amino acid sequence of SEQ ID NO: 396, and a VL CDR3 having an amino acid sequence of SEQ ID NO: 397. In one instance, an anti-PD-1 antibody or an anti-PD-1 antigen binding fragment comprises a VH CDR1 having an amino acid sequence of SEQ ID NO: 398, a VH CDR2 having an amino acid sequence of SEQ ID NO: 399, a VH CDR3 having an amino acid sequence of SEQ ID NO: 400, VL CDR1 having an amino acid sequence of SEQ ID NO: 401, a VL CDR2 having an amino acid sequence of SEQ ID NO: 402, and a VL CDR3 having an amino acid sequence of SEQ ID NO: 403.

In one instance, an anti-PD-1 antibody comprises a fusion protein. Such fusion protein can be, for example, a two-sided Fc fusion protein comprising the extracellular domain (ECD) of programmed cell death 1 (PD-1) and the ECD of tumor necrosis factor (ligand) superfamily member 4 (TNFSF4 or OX40L) fused via hinge-CH2-CH3 Fc domain of human IgG4, expressed in CHO-K1 cells, where the fusion protein has an exemplary amino acid sequence of SEQ ID NO: 104.

In some embodiments, the antibody or antigen binding fragment thereof is an anti-PD-L1 antibody or antigen binding fragment. Programmed death-ligand 1 (PD-L1) is a ligand for an immunosuppressive receptor “programmed death receptor 1 (PD-1)” that is predominantly expressed in activated T and B cells, which can negatively regulate antigen receptor signaling. The ligands (PD-L1 and PD-L2) for PD-1 may be constitutively expressed or may be derived into a number of cell types, including non-hematopoietic cell tissues and various tumor types. PD-L1 is expressed in B cells, T cells, bone marrow cells and dendritic cells (DCs), but also on non-lymphatic organs such as peripheral cells, pseudo-vascular endothelial cells and heart, lungs, etc. A non-limiting, exemplary, human PD-L1 amino acid sequence is MRIFAVFIFMTYWHLLNAFTVTVPKDLY-

30

VVEYGSNMTIECKFPVEKQLDLAALIVYWE  
MEDKNIIQFVH-  
GEEDLKVKHQSSYRQRARLLKDQLSLG-  
NAALQITDVKLQDAGVYRCMI SYGGA-  
5 DYKRITVKVNAPYNKINQRILVVDPVTSEHELTCAE-  
GYPKAEVIWTSSDHQVL SGKTTTNSKREKEKFNVTLRINTTTNEIFYC-  
TFRRLDPEENHTAELVPIELPLAHPPNE RTHLVIL-  
10 GAILLCLGVALTFI-  
FRLRKGRMMDVKKCGIQDTNSKKQSDTHLEET (SEQ  
ID NO: 330)

In one embodiment, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment of the disclosure comprises a combination of a heavy chain variable region (VH) and a light chain variable region (VL) described herein. In another embodiment, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment of the disclosure comprises a combination of complementarity determining regions (VH CDR1, VH CDR2, VH CDR3, VL CDR1, VL CDR2, and VL CDR3) described herein. In one embodiment, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment of the disclosure comprises a modified Modified Avelumab 15 (Bavencio, 451238, KXG2PJ551I, MSB-0010682, MSB-0010718C, PF-06834635, CAS 1537032-82-8: EMD Serono, Merck & Co., Merck KGaA, Merck Serono, National Cancer Institute (NCI), Pfizer), Durvalumab (Imfinzi, 28x28x90 KV (UNII code), MEDI-4736, CAS 1428935-60-7: AstraZeneca, Celgene, Children’s Hospital Los Angeles (CHLA), City of Hope National Medical Center, MedImmune, Memorial Sloan-Kettering Cancer Center, Mirati Therapeutics, National Cancer Institute (NCI), Samsung Medical Center (SMC), Washington University), Atezolizumab (Tecentriq, 52CMI0WC3Y, MPDL-3280A, RG-7446, RO-5541267, CAS 1380723-44-3: Academisch Medisch Centrum (AMC), Chugai Pharmaceutical, EORTC, Genentech, Immune Design (Merck & Co.), Memorial 20 Sloan-Kettering Cancer Center, National Cancer Institute (NCI), Roche, Roche Center for Medical Genomics), Sugemalimab (CS-1001, WBP-3155: CStone Pharmaceuticals, EQRx, Pfizer), KN-046 (CAS 2256084-03-2: Jiangsu Alphamab Biopharmaceuticals, Sinovent), APL-502 (CBT-502, TQB-2450: Apollomics, Jiangsu Chia Tai Tianqing Pharmaceutical), Envafolimab (3D-025, ASC-22, KN-035, hu56V1-Fc-m1, CAS 2102192-68-5: 3D Medicines, Ascletis, Jiangsu Alphamab Biopharmaceuticals, Suzhou Alphamab, Tracon Pharmaceuticals, Inc.), Bintrafusp alfa 25 (M-7824, MSB-0011359C, NW9K8C1JN3, CAS 1918149-01-5: EMD Serono, GlaxoSmithKline, Merck KGaA, National Cancer Institute (NCI)), STI-1014 (STI-A1014, ZKAB-001: Lee’s Pharmaceutical, Sorrento Therapeutics), PD-L1 t-haNK (ImmunityBio, NantKwest), A-167 (HBM-9167, KL-A167: Harbour BioMed, Sichuan Kelun-Biotech Biopharmaceutical), IMC-001 (STI-3031, STI-A-1015, STI-A1015, ImmuneOncia Therapeutics, Sorrento Therapeutics), HTI-1088 (SHR-1316: Atridox, Jiangsu Hengrui), IO-103 (IO Biotech), CX-072 (CytomX Therapeutics), AUPM-170 (CA-170: Aurigen, Curis), GS-4224 (Gilead), ND-021 (NM21-1480, PRO-1480: CStone Pharmaceuticals, Numab Therapeutics), BNT-311 (DuoBody-PD-L1x4-1BB, GEN-1046: BioNTech, Genmab), BGB-A333 (BeiGene), IBI-322 (Innovent Biologics), NM-01 (Nanomab Technol-

ogy, Shanghai First People's Hospital), LY-3434172 (Eli Lilly), LDP (Dragonboat Biopharmaceutical), CDX-527 (Celldex Therapeutics), IBI-318 (Innovenet Biologics, Lilly), 89Zr-DFO-REGN3504 (Regeneron), ALPN-202 (CD80 vIgD-Fc: Alpine Immune Sciences), INCB-086550 (Incyte), LY-3415244 (Eli Lilly), SHR-1701 (Jiangsu Hengrui), JS-003 (JS003-30, JS003-SD: Shanghai Junshi Biosciences), HLX-20 (PL2 #3: Henlix Biotech, Shanghai Henlius Biotech), ES-101 (INBRX-105, INBRX-105-1: Elpiscience BioPharma, Inhibrix), MSB-2311 (MabSpace Biosciences), PD-1-Fc-OX40L (SL-279252, TAK-252: Heat Biologics, Shattuck Labs, Takeda), FS-118, FS118 mAb2, LAG-3/PD-L1 mAb2: F-star Therapeutics, Merck & Co., Merck KGaA), FAZ-053 (LAE-005: Laekna Therapeutics, Novartis), Lodapolimab (LY-3300054, NR4MAD6PPB, CAS 2118349-31-6: Eli Lilly), MCLA-145 (Incyte, Merus), BMS-189 (BMS-986189, PD-L1-Milla from Bristol-Myers Squibb), Cosibelimab (CK-301, TG-1501, CAS 2216751-26-5: Checkpoint Therapeutics, Dana-Farber Cancer Institute, Samsung Biologics, TG Therapeutics), IL-15Ralpha-S/D/IL-15 (KD-033: Kadmon), WP-1066 (CAS 857064-38-1: M.D. Anderson Cancer Center, Moleculin Biotech), BMS-936559 (MDX-1105: Bristol-Myers Squibb, Medarex, National Institute Allergy Infect Dis.), BMS-986192 (Bristol-Myers Squibb), RC-98 (RemeGen), CD-200AR-L (CD200AR-L: OX2 Therapeutics, University of Minnesota), ATA-3271 (Atara Biotherapeutics), IBC-Ab002 (ImmunoBrain Checkpoint), BMX-101 (Biomunex Pharmaceuticals), AVA-04-VbP (Avacta), ACE-1708 (Acepodia Biotech), KY-1043 (Kymab, Provenance Biopharmaceuticals), ACE-05 (YBL-013: Y-Biologics), ONC-0055 (ONC0055, PRS-344 S-095012: Pieris Pharmaceuticals, Servier), TLJ-1-CK (I-Mab Biopharma), GR-1405 (Chinese Academy of Medical Sciences), PD-1ACR-T (Taipei Medical University), N-809 (N-IL15/PD-L1: ImmunityBio), CB-201 (Crescendo Biologics), MEDI-1109 (MedImmune), AVA-004 (AVA-04: Avacta), CA-327 (Aurigene, Curis), ALN-PDL (Alnylam Pharmaceuticals), KY-1003 (Kymab), CD22(aPD-L1)CAR-T cells (SL-22P: Hebei Senlang Biotechnology), ATA-2271 (M28zLXXPD-1DNR CAR T cells: Atara Biotherapeutics), and Zeushield cytotoxic T lymphocytes (Second Xiangya Hosp Central South Univ.).

In some embodiments, the anti-PD-L1 antibody is Avelumab, Durvalumab, Atezolizumab, Sugemalimab, Envafolimab, Lodapolimab, or Cosibelimab, or a modified version thereof. In some embodiments, the anti-PD-L1 antibody is Avelumab, Durvalumab, Atezolizumab, Sugemalimab, Envafolimab, Lodapolimab, or Cosibelimab. In some embodiments, the antibody is a biosimilar of Avelumab, Durvalumab, Atezolizumab, Sugemalimab, Envafolimab, Lodapolimab, or Cosibelimab.

TABLE 1 provides the sequences of exemplary anti-PD-L1 antibodies and anti-PD-L1 antigen binding fragments that can be modified to prepare anti-PD-L1 immunoconjugates. TABLE 1 also provides exemplary combinations of CDRs that can be utilized in a modified anti-PD-L1 immunoconjugate. Reference to an anti-PD-L1 antibody herein may alternatively refer to an anti-PD-L1 antigen binding fragment.

In some embodiments, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy

chain or VH having an amino acid sequence of any one of SEQ ID NOS: 232, 234, 236, 238, 242, 244, or 248. An anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of any one of SEQ ID NOS: 233, 235, 237, 239, 243, 245, or 249. In one instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of SEQ ID NO: 232, and a light chain or VL having an amino acid sequence of SEQ ID NO: 233. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of SEQ ID NO: 234, and a light chain or VL having an amino acid sequence of SEQ ID NO: 235. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of SEQ ID NO: 236, and a light chain or VL having an amino acid sequence of SEQ ID NO: 237. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of SEQ ID NO: 238, and a light chain or VL having an amino acid sequence of SEQ ID NO: 239. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a VH having an amino acid sequence of SEQ ID NO: 242, and a VL having an amino acid sequence of SEQ ID NO: 243. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a VH having an amino acid sequence of SEQ ID NO: 244, and a VL having an amino acid sequence of SEQ ID NO: 245. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of SEQ ID NO: 248, and a light chain or VL having an amino acid sequence of SEQ ID NO: 249.

In some embodiments, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence of any one of SEQ ID NOS: 232, 234, 236, 238, 242, 244, or 248, or a portion corresponding to a VH thereof. An anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a light chain or VL having an amino acid sequence of any one of SEQ ID NOS: 233, 235, 237, 239, 243, 245, or 249, or a portion corresponding to a VL thereof. In one instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence shown in SEQ ID NO: 232, and a light chain or VL having an amino acid sequence shown in SEQ ID NO: 233. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence shown in SEQ ID NO: 234, and a light chain or VL having an amino acid sequence shown in SEQ ID NO: 235. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence shown in SEQ ID NO: 236, and a light chain or VL having an amino acid sequence shown in SEQ ID NO: 237. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence shown in SEQ ID NO: 238, and a light chain or VL having an amino acid sequence shown in SEQ ID NO: 239. In another instance, an anti-

PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 242, and a VL having an amino acid sequence shown in SEQ ID NO: 243. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a VH having an amino acid sequence shown in SEQ ID NO: 244, and a VL having an amino acid sequence shown in SEQ ID NO: 245. In another instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a heavy chain or VH having an amino acid sequence shown in SEQ TD NO: 248, and a light chain or VL having an amino acid sequence shown in SEQ TD NO: 249.

In one instance, an anti-PD-L1 antibody or an anti-PD-L1 antigen binding fragment comprises a VH CDR1 having an

amino acid sequence of SEQ ID NO: 250, a VH CDR2 having an amino acid sequence of SEQ ID NO: 251, a VH CDR3 having an amino acid sequence of SEQ ID NO: 252, VL CDR1 having an amino acid sequence of SEQ ID NO: 253, a VL CDR2 having an amino acid sequence of SEQ TD NO: 254, and a VL CDR3 having an amino acid sequence of SEQ TD NO: 255.

In one instance, an anti-PD-L1 antibody comprises a single domain binding antibody having an amino acid sequence of SEQ TD NO: 256, a anti-specific fusion single chain antibody construct having an amino acid sequence of SEQ ID NO: 257, or a bispecific tetrameric antibody like engager having an amino acid sequence of SEQ TD NO: 258.

TABLE 1

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Tislelizumab, Baizean, OKVO411B3N, BGB-A317, hu317- 1/IgG4mt2 Heavy Chain (VH in Bold)	PD-1	QVQLQESGPGLVKPSETLSLTCTVSGFSILSYGVHWIRQPPKG GLEWIGIVIYADGSTNYNPSLKSRTISKDTSKNQVSLKLSSVT AADTAVYYCARAYGNYWYIDVWGGTTVTVSSASTKGPSVFP LAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHTFPA VLQSSGLYSLSSVTVTPSSSLGTTKTYTCNVDHKPSNTKVDKRVES KYGPPCPCCPAPPVAGGPSVFLFPPPKPKDTLMISRTPEVTCVVAV SQEDPEVQFNWYVDGVEVHNAKTKPREEQFNSTYRVSVLTVV HQDWLNGKEYCKVSNKGLPSSIETKISKAKGQPREPQVTLP QEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPV DSDGSFFLYSKLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSL LSLGK	332
Tislelizumab, Baizean, OKVO411B3N, BGB-A317, hu317- 1/IgG4mt2 Light Chain (VL in Bold)	PD-1	DIVMTQSPDSLAVSLGERATINCKSSESVSNDVAVYQQKPGQP PKLLINYAFHRFTGVDFRFSGGYGTDFLTISLQAEDVAVY YCHQAYSSPYTFGQGTKEIKRTVAAPSVFIFPPSDEQLKSGTAS VVCCLNNFYPREAKVQWVKDNAQSGNSQESVTEQDSKDSTYS LSSTLTSKADYEHKVYACEVTHQGLSSPVTKSFNRGEC	333
Sintilimab, Tyvyt, IBI- 308 Heavy Chain (VH in Bold)	PD-1	QVQLVQSGAEVKKPGSSVKVSCKASGGTFSSYAIISWVRQAPG QGLEWMGLIIPMFDTAGYAQKFQGRVAITVDESTSTAYMELS SLRSEDTAVYYCARAEHSSTGTFDYWGGQGLTVTVSSASTKGPS VFP LAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHT FPAVLQSSGLYSLSSVTVTPSSSLGTTKTYTCNVDHKPSNTKVDKR VESKYGPPCPCCPAPFELGGPSVFLFPPPKPKDTLMISRTPEVTCVV VDVSKQEDPEVQFNWYVDGVEVHNAKTKPREEQFNSTYRVSVL TVLHQDWLNGKEYCKVSNKGLPSSIETKISKAKGQPREPQVT LPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKT PVLDSDGSSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQK SLSLSLGK	334
Sintilimab, Tyvyt, IBI- 308 Light Chain (VL in Bold)	PD-1	DIQMTQSPSSVSASVGDRVTITCRASQGISSWLAZYQQKPGK APKLLISAASSLQSGVPSRFSRGSGSGTDFLTISLQPEDFATY CQQANHLPFTFGGGTKVEIKRTVAAPSVFIFPPSDEQLKSGTAS VVCCLNNFYPREAKVQWVKDNAQSGNSQESVTEQDSKDSTYS LSSTLTSKADYEHKVYACEVTHQGLSSPVTKSFNRGEC	335
Toripalimab, TeRuiPuLi, Terepril, Tuoyi, JS- 001, TAB- 001 Heavy Chain (VH in Bold)	PD-1	QGQLVQSGAEVKKPGASVKVSCKASGYTFTDYEMHWVRQA PIHGLEWIGVIESETGGTAYNQFKGRVTITADKSTSTAYMEL SSLRSEDTAVYYCAREGITTATVATTYYWYFDVWGGQTTVTVSS ASTKGPSVPLAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSG LTSGVHTFPAVLQSSGLYSLSSVTVTPSSSLGTTKTYTCNVDHKPS NTKVDKRVESKGYPCCPCCPAPFELGGPSVFLFPPPKPKDTLMISRT PEVTCVVVDVSKQEDPEVQFNWYVDGVEVHNAKTKPREEQFNST YRVSVLTVLHQDWLNGKEYCKVSNKGLPSSIETKISKAKGQP REPQVYTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE NNYKTTPPVLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEAL HNHYTQKSLSLSLGK	336

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Toripalimab, TeRuPuLi, Terepril, Tuoyi, JS-001, TAB-001 Light Chain (VL in Bold)	PD-1	DVVMTQSPLSLPVTLGQPAISCRSSQSIVHSNGNTYLEWYLQ KPGQSPQLLIYKVSNRFSGVPDRFSGSGCTDFTLKRISRVEAED VGVYYCFQGSHVPLTFGQGTTKLEIKRTVAAPSVIFPPSDEQLK SGTASVVCLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSK DSTYLSLTLTKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC	337
Camrelizumab, HR-301210, INC SHR-01210, SHR-1210 Heavy Chain (VH in Bold)	PD-1	EVQLVESGGGLVQPGGSLRLSCAASGFTFSSYMMWSVRQAP GKGLEWVATISGGANTYYPDSDKRFTISRDNAKNSLYLQM NSLRAEDTAVYYCARQLYYFDYWGQGTTVVSASTKGPSVF PLAPCSRSTSEESTAALGLCLVKDYFPEPVTVWSNSGALTSGVHTFP AVLQSSGLYSLSSVTPVSSSLGTTKTYTCNVDHKPSNTKVDKRVE SKYGPCCPCCPAPEFLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSQEDPEVQFNWYWDGVEVHNNAKTKPREEQFNSTYRVVSVLTV LHQDWLNGKEYKCKVSNKGLPSSIETKTISKAKGQPREQVYTLPP SQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPV LDSDGSSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSL SLSLGK	338
Camrelizumab, HR-301210, INC SHR-01210, SHR-1210 Light Chain (Light Chain in Bold)	PD-1	DIQMTQSPSSLSASVGDRVTITCLASQTIGTWLTYQQKPGK APKLLIYTATSLADGVPSRFSGSGSGTDFTLTISLQPEDFATYY YCQQVYSIPWTFGGGTKVEIKRTVAAPSVFIFPPSDEQLKSGTA SVVCLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYS LSSTLTLKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC	339
Cemiplimab, Cemiplimab-rwlc, LIBTAYO®, 6QVL057INT, H4H7798N, REGN-2810, SAR-439684 Heavy Chain (VH in Bold)	PD-1	EVQLLESGGGLVQPGGSLRLSCAASGFTFSNFGMTWVRQAPG KGLEWVSGISGGGRDTYFADSVKRFTISRDNASKNTLYLQMN SLKGEDTAVYYCVKWGNIYFDYWGQGTLTVVSASTKGPSVF PLAPCSRSTSEESTAALGLCLVKDYFPEPVTVWSNSGALTSGVHTFP AVLQSSGLYSLSSVTPVSSSLGTTKTYTCNVDHKPSNTKVDKRVE SKYGPCCPCCPAPEFLGGPSVFLFPPKPKDTLMISRTPEVTCVVVD VSQEDPEVQFNWYWDGVEVHNNAKTKPREEQFNSTYRVVSVLTV LHQDWLNGKEYKCKVSNKGLPSSIETKTISKAKGQPREQVYTLPP SQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPPV LDSDGSSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSL SLSLGK	340
Cemiplimab, Cemiplimab-rwlc, LIBTAYO®, 6QVL057INT, H4H7798N, REGN-2810, SAR-439684 Light Chain (VL in Bold)	PD-1	DIQMTQSPSSLSASVGDSITITCRASLSINTFLNWYQQKPGKAP NLLIYAASSLHGGVPSRFSGSGSGTDFTLTIRTLQPEDFATYYC QQSSNTPFTFGPGTVDFRTTVAPASVIFPPSDEQLKSGTASVW CLLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYSLSS TLTLTSKADYEKHKVYACEVTHQGLSSPVTKSFRGEC (Disulfide bridge) H22-H96, H131-L214, H144-H200, H223-H'223, H226- H'226, H258-H318, H364-H422, H'22-H'96, H'131-L'214, H'114- H'200, H'258-H'318, H'364-H'422, L23-L88, L134-L194, L'23- L'88, L'134-L'194)	341
Lambrolizumab, Pembrolizumab, 1 KEYTRUDA®, MK-3475, SCH-900475, h409A11 Heavy Chain (VH in Bold)	PD-1	QVQLVQSGVEVKPGASVKVSCKASGYTFTNYYMWWVRQAP GQGLEWMGGINPSNNGTNFNEKFKNRVTLLTDSTTTAYME LKSILQFDTTAVYYCARRDYRFDMGFDYWGQGTTVVSAST KGPSVFPLACSRSTSEESTAALGLCLVKDYFPEPVTVWSNSGALT GVHTFPALQSSGLYSLSSVTPVSSSLGTTKTYTCNVDHKPSNTK VDKRVESKYGPCCPCCPAPEFLGGPSVFLFPPKPKDTLMISRTPEV TCVVVDVSQEDPEVQFNWYWDGVEVHNNAKTKPREEQFNSTYRV VSVLTVLHQDWLNKEYKCKVSNKGLPSSIETKTISKAKGQPREQ VYTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNY KTTPPVLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHN YTQKSLSLSLGK	346
Lambrolizumab, Pembrolizumab, KEYTRUDA®	PD-1	EIVLTQSPATLSLSPGERATLSCRASKGVSTSGSYSYLHWYQQK PGQAPRLLIYLASYLESGVPARFSGSGSGTDFTLTISLLEPEDF AVYYCQHSRDLPLTFGGGTKVEIKRTVAAPSVFIFPPSDEQLKS	347

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
MK-3475, SCH-900475, h409A11 Light Chain (VL in Bold)		GTASVVCLNNFYPREAKVQWKVDNALQSGNSQESVTBQDSKD STYSLSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC	
Lambrolizumab, Pembrolizumab, KEYTRUDA®, MK-3475, SCH-900475, h409A11 VH	PD-1	QVQLVQSGVEVKPGASVKVSCKASGYTFTNYYMWVRQAPG QGLEWMGGFPSPNGTNFNEKFKNRVTLTTDSSTTAYMELKSLQ FDDTAVYYCARRDYRFDMGFDYWGQGTTVTVSS (Disulfide bridge) H22-H96, H134-L218, H147-H203, H226-H'226, H229-H'229, H261-H321, H367-H425, H'22-H'96, H'134-L'218, H'147-H'203, H'261-H'321, H'367-H'425, L23-L92, L138-L198, L'23-L'92, L'138-L'198)	348
Lambrolizumab, Pembrolizumab, KEYTRUDA®, MK-3475, SCH-900475, h409A11 VL	PD-1	EIVLTQSPATLSLSPGERATLSCRASKGVSTSGYSYLYHWWYQQKPG QAPRLLIYLASYLESGVPARFSGSGSGTDFTLTISLEPEDFAVYYC QHSRDLPLTFGGGTKEIK	349
Nivolumab, Nivolumab BMS, OPDIVO®, BMS-936558, MDX-1106, ONO-4538 VH	PD-1	QVQLVESGGVVQPGRSRLLDCKASGITFSNSGMHWVRQAPKG GLEWVAIVWDGSKRYYADSVKGRTFISRDNSKNTLFLQMNLSR AEDTAVYYCATNDDYWGQGTLVTVSS	350
Nivolumab, Nivolumab BMS, OPDIVO®, BMS-936558, MDX-1106, ONO-4538 VL	PD-1	EIVLTQSPATLSLSPGERATLSCRASQSVSSYLAWYQQKPGQAPR LLIYDASNRATGIPARFSGSGSGTDFTLTISLEPEDFAVYYCQOSS NWPPRTFCQCQTKVEIK (Disulfide bridge) H22-H96, H127-L214, H140-H196, H219-H'219, H222-H'222, H254-H314, H360-H418, H'22-H'96, H'127-L'214, H'140-H'196, H254-H'314, H'360-H'418, L-23-L88, L134-L194, L'23-L'88, L'134-L'194)	351
Prolgolimab, Forteca, BCD-100 Heavy Chain (VH in Bold)	PD-1	QVQLVQSGGGLVQPGGSLRLSCAASGFTFSSYWMHWVRQVP GKGLEWVAIDTGGGRTYYADSVKGRFAISRVNAKNTMYLQ MNSLRAEDTAVYYCARDEGGGTGWGVLKDWPYGLDAWQ GTLIVTVSSASTKGPSVFPLAPSSKSTSGGTAALGCLVKDYFPEPV TVSWNSGALTSGVHTFPVALQSSGLYSLSSVVTVPSSSLGTQTYIC VNHNKPSNTKVDKVEPKSCDKTHTCPPCPAPEAAGGPSVFLPP KPKDTLMISRTPETVCVVVDVSHEDPEVFKFNWYDGVEVHN TKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPI EKTISKAKGQPREFQVYTLPPSRREMTKNQVSLTCLVKGFYPSDI AVEWESNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKSRWQQGN VFSCSVMHEALHNHYTQKSLSLSPGK	352
Prolgolimab, Forteca, BCD-100 Light Chain (VL in Bold)	PD-1	QPVLQTPLSVSVALGQTARITCGGNIGSKNVHWYQQKPGQ APVILVIYRDSNRPSGIPERFSGNSGNATLTISRAQAGDEADY YCQVWDSSATAFGTGTKLTVLQRTVAAPSVFIFPPSDEQLKSGT ASVCLNNFYPREAKVQWKVDNALQSGNSQESVTBQDSKDST YSLSSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC	353
Balstilimab, 1Q2QT5M7EO, AGEN-2034, AGEN-2034w Heavy Chain (VH in Bold)	PD-1	QVQLVESGGVVQPGRSRLSCAASGFTFSSYGMHWVRQAP GKGLEWVAIVWDGSNKYYADSVKGRFTISRDNSKNTLFLQ MNSLRAEDTAVYYCASNGDHWGQGTLVTVSSASTKGPSVFPL APCSRSTSEESTAALGCLVKDYFPEPVTVWSWNSGALTSGVHTFP LQSSGLYSLSSVVTVPSSSLGTKYTCNVDHKPSNTKVDKRVESK YGPVPCPPCAPEFLGGPSVFLFPKPKDTLMISRTPETVCVVVDV QEDPEVQFNWYVDGVEVHNAKTPREEQFNSTYRVVSVLTVLH QDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREFQVYTLPPSQ	354

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Balstilimab, 1Q2QT5M7EO, AGEN-2034, AGEN-2034w Light Chain (VL in Bold)	PD-1	EEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTPPVLD SDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLSL SLG	
Dostarlimab, ANB-011, GSK- 4057190A, POGVQ9A4S5, TSR-042, WBP- 285 Heavy Chain (VH in Bold)	PD-1	EIVMTQSPATLSVPGERATLSCRASQS VSSNLAWYQQKPGQ APRLLIYGASTRATGIPARFSGSGSGTEFTLTISLQSEDFAVYY CQQYNNWPRTFGQGT KVEIKRTVAAPS VIFPPSDEQLKSGTAS VVCLNNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYS LSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKS FNRGEC	355
Dostarlimab, ANB-011, GSK- 4057190A, POGVQ9A4S5, TSR-042, WBP- 285 Light Chain (VL in Bold)	PD-1	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYDMSWVRQAPG KGLEWVSTISGGGSYTYYQDSVKGRTFISRDNKNTLYLQMN SLRAEDTA VYYCASPYYAMDYWGQTTTVVSSASTKGPSVFP LAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHTPPA VLQSSGLYSLSSVVTVPSSSLGTKTYTCNVVDHKPSNTKVDKRVES KYGPPCPCCPAPEFLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV SQEDPEVQFNWYV DGEVHVNAKTKPREEQFNSTYRVSVLTVL HQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPS QEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTPPVLD DSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLS LSLGK	356
Dostarlimab, ANB-011, GSK- 4057190A, POGVQ9A4S5, TSR-042, WBP- 285 Light Chain (VH in Bold)	PD-1	DIQLTQSPSFLSAYVGDRV TITCKASQDV GTA VAWYQQKPGK APKLLIYWA STLHTGVPSRSFSGSGSGTEFTLTISLQPEDFATY YCQHYSSYPWTFGQGT KLEIKRTVAAPS VIFPPSDEQLKSGTAS SVVCLNNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYS LSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKS FNRGEC	357
Serplulimab, HLX-10 Heavy Chain (VH in Bold)	PD-1	QVQLVESGGGLVKP GGSRLSCAASGFTFSNYGMSWIRQAPG KGLEWSTISGGGSNIYYADSVKGRTFISRDNKNSLYLQMN RAEDTA VYYCVSYYGYIDFWGQGT STVVSSASKYGPSVFPLAPC SRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHTFP AVLQS SGLYSLSSVVTVPSSSLGTKTYTCNVVDHKPSNTKVDKRVESKYGP PCPPCPA PPEFLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV S QED PEVQFNWYV DGEVHVNAKTKPREEQFNSTYRVVSVVLTVLHQD WLN GKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTLPPS QEE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTPPVLD GSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKSLSL GK	358
Serplulimab, HLX-10 Light Chain (VL in Bold)	PD-1	DIQMTQSPSFLSASVGDRV TITCKASQDVTTAVAWYQQKPGK APKLLIYWA STRH TGVPSRSFSGSGSGTDFTLTISLQPEDFATY YCQHYTI PWTFGGGT KLEIKRTVAAPS VIFPPSDEQLKSGTAS SVVCLNNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYS LSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKS FNRGEC	359
Retifanlimab, 2Y3T5IF01Z, INCMGA-00012, INCMGA- 0012, MGA- 012 Heavy Chain (VH in Bold)	PD-1	QVQLVQSGAEVKKPGASVVKVSCKASGY SFTSYWMNWVRQAP GQGLEWIGVIHPDSSETWL DQKF KDRV TITV DKST STAYMEL SSLRSEDTAVYYCAREHYG TS PFA YWGQ GTLTVVSSASTKGPS VFPLAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHT FPAVLQSSGLYSLSSVVTVPSSSLGTKTYTCNVVDHKPSNTKVDKR VESKYGPPCPCCPAPEFLGGPSVFLFPPKPKDTLMISRTPEVTCVV VDV S QEDPEVQFNWYV DGEVHVNAKTKPREEQFNSTYRVVSVL TVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVY LPPS QEE MTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTP PVLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQK SLSL SLG	360

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Retifanlimab, 2Y3T5IF01Z, INCmGA- 0012, INCmGA- 0012, MGA- 012 Light Chain (VL in Bold)	PD-1	<b>EIVLTQSPATLSLSPGERATLSCRASESVDNYGMSFMNWPFQQ</b> KPGQPCKLIIHAASNQGSGVPSSRSGSGSGTDFTLTISSLEPED <b>FAVYFCQQSKVEPVTFGGGTKEIKRTVAAPSVFIFPPSDEQLK</b> SGTASVVCCLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSK DSTYSLSSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNRGEC	361
Sasanlimab, LZZ01C2EWP, PF- 06801591, RN-888 Heavy Chain (VH in Bold)	PD-1	<b>QVQLVQSGAEVKPGASVKVSCKASGYTFTSYWINWVRQAP</b> GQGLEWMGNIYPGSSLTNYNEKFKNRVTMTRDTSTSTVYME <b>LSSLRSEDTAVYYCARLSTGTFAWQGQTLTVVSSASTKGPSV</b> FPLAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHTF PAVLQSSGLYSLSSVTPVSSSLGTTCTCNVDHKPSNTKVDKR ESKYGPPCPCCPAPEFLGGPSVFLFPPPKDITLMISRTPETCVVV DVSDQEDPEVQFNWYVDGVEVHNAAKTKPREEQFNSTYRVVSVL VLHQDWLNKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYTL PSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPP VLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQKS LSLSLGK	362
Sasanlimab, LZZ01C2EWP, PF- 06801591, RN-888 Light Chain (VL in Bold)	PD-1	<b>DIVMTQSPDSLAVSLGERATINCKSSQSLWDSGNQNKNFLTWY</b> QQKPGQPPKLLIYWTSYRESGVPDFSGSGSGTDFTLTISSLQ <b>AEDAVVYVQNDYFPHTFGGTKEIKRTVAAPSVFIFPPSDE</b> QLKSGTASVVCCLNNFYPREAKVQWKVDNALQSGNSQESVTEQ DSKDSTYSLSSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFNR GEC	363
Spatializumab, NVP- LZV-184, PDR-001, QO25L6Z8Z Heavy Chain (VH in Bold)	PD-1	<b>EVQLVQSGAEVKPGESLRISCKGSGYTFTTYWMHWVRQAT</b> GQGLEWMGNIYPGTTGSNFDEKFKNRVTITADKSTSTAYME <b>LSSLRSEDTAVYYCTRWTGTGAYWQGQTTVTVSSASTKGPS</b> VFPLAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSGVHT FPAVLQSSGLYSLSSVTPVSSSLGTTCTCNVDHKPSNTKVDKR VESKYGPPCPCCPAPEFLGGPSVFLFPPPKDITLMISRTPETCVV DVSDQEDPEVQFNWYVDGVEVHNAAKTKPREEQFNSTYRVVSVL TVLHQDWLNKEYKCKVSNKGLPSSIEKTISKAKGQPREPQVYT LPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYKTTPP VLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHYTQK SLSLSLG	364
Spatializumab, NVP- LZV-184, PDR-001, QO25L6Z8Z Light Chain (VL in Bold)	PD-1	<b>EIVLTQSPATLSLSPGERATLSCRASSQSLLDGNGNQNKNFLTWYQ</b> QKPGQAPRLLIYWASTRESGVPDFSGSGSGTDFTLTISSLEAE <b>DAATYYCQNDYSPYTFGGTKEIKRTVAAPSVFIFPPSDEQL</b> KSGTASVVCCLNNFYPREAKVQWKVDNALQSGNSQESVTEQDS KDSTYSLSSSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFRGE C	365
Cetrelimab, JNJ-3283, JNJ- 63723283, LYK98WP91F Heavy Chain (VH in Bold)	PD-1	<b>QVQLVQSGAEVKPGSSVKVSCKASGGTFSSYAIISWVRQAPG</b> QGLEWMGGIIPIDTANYAQKFQGRVTITADESTSTAYME <b>LRSEDTAVYYCARPLAAYDTGSLDYWGQGTLTVVSSASTK</b> GPSVPLAPCSRSTSESTAALGCLVKDYFPEPVTVWSNSGALTSG VHTFPVLQSSGLYSLSSVTPVSSSLGTTCTCNVDHKPSNTK DKRVESKYGPPCPCCPAPEFLGGPSVFLFPPPKDITLMISRTPET CVVVDVSQEDPEVQFNWYVDGVEVHNAAKTKPREEQFNSTYRVV SVLTVLHQDWLNKEYKCKVSNKGLPSSIEKTISKAKGQPREPQV YTLPPSQEEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENNYK TTPPVLDSDGSFFLYSRLTVDKSRWQEGNVFSCSVMHEALHNHY TQKSLSLSLGK	366
Cetrelimab, JNJ-3283, JNJ- 63723283, LYK98WP91F Light Chain (VL in Bold)	PD-1	<b>EIVLTQSPATLSLSPGERATLSCRASQSVRSYIAWYQQKPGQA</b> PRLLIYDASNRATGIPARFSGSGSGTDFTLTISSLEPEDFAVYYC <b>QQRNYWPLTFGGTKEIKRTVAAPSVFIFPPSDEQLKSGTASV</b> VCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYSL SSTLTLSKADYEKHKVYACEVTHQGLSSPVTKSFRGE	367

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Tebotelimab, MGD-013 Heavy Chain (VL in Bold)	PD-1	<b>D</b> IQLTQSPSSLSAVGDRVTITCRASQDVSSVVAWYQQKPGK <b>A</b> PKLLI <b>S</b> ASYRYTGVP <i>S</i> RFS <i>G</i> SGSTDFTLT <b>I</b> SSLQPEDFATY <b>Y</b> CQQHYS <i>P</i> WT <i>F</i> GGGT <i>K</i> LEIKGGGSGGGQVQLVQSGAEVKK PGASVKVS <b>C</b> ASGY <i>F</i> TS <i>Y</i> WMN <i>V</i> RQAP <i>Q</i> GLEWIGV <b>I</b> HPSDSE TWLDQKFDRVTITV <b>D</b> KST <b>T</b> AYMELSSLRSED <i>T</i> AVYYCAREHY GTSP <i>F</i> AYWGQGT <i>L</i> TVSS <i>G</i> CGGG <i>E</i> VAAC <i>E</i> KEVAALKEVAAL EKEVAALKE <i>E</i> SKYGGPPCP <i>C</i> PAPE <i>F</i> LGGS <i>P</i> SVFLFPP <i>P</i> KDTLY <i>I</i> TR EPEVTCVVVD <i>V</i> DVSQED <i>P</i> EVQFN <i>W</i> DGV <i>V</i> HNAKT <i>K</i> PREEQFNS TYRVSVLTVLHQDWLNGKEYKCKVSNKGLP <i>S</i> SI <i>E</i> KT <i>I</i> SKAKGQ PREPVYTL <i>L</i> PP <i>S</i> QEEMTKNQVSL <i>T</i> CLVKG <i>F</i> YPS <i>D</i> IAVEWESNGQP ENNYTT <i>TP</i> PLSDGS <i>F</i> FLYSR <i>L</i> TVDKSRW <i>Q</i> EGNV <i>F</i> SCVMHEA LHNHTQ <i>K</i> SL <i>S</i> LS <i>L</i> SG	368
Tebotelimab, MGD-013 Light Chain (VL in Bold)	PD-1	<b>E</b> IVLTQSPATL <i>S</i> LP <i>G</i> ERA <i>T</i> LS <i>C</i> RASE <i>S</i> VD <i>N</i> YGM <i>S</i> FM <i>N</i> WF <i>Q</i> Q KPGPP <i>K</i> L <i>I</i> HAAS <i>N</i> Q <i>G</i> V <i>P</i> S <i>R</i> F <i>S</i> GS <i>G</i> STDFTLT <b>I</b> SSL <i>E</i> PED <b>F</b> AV <i>V</i> YC <i>Q</i> Q <i>S</i> KE <i>V</i> P <i>T</i> Y <i>F</i> GG <i>G</i> TK <i>V</i> E <i>I</i> KGGGSGGG <i>Q</i> V <i>Q</i> LS <i>G</i> SA EV <i>K</i> PG <i>A</i> SV <i>K</i> V <i>S</i> CK <i>A</i> SG <i>Y</i> FT <i>D</i> Y <i>M</i> WD <i>W</i> R <i>Q</i> AP <i>Q</i> GLE <i>W</i> MG <i>D</i> i NP <i>D</i> NG <i>V</i> TI <i>Y</i> N <i>Q</i> K <i>F</i> GR <i>V</i> T <i>M</i> TD <i>T</i> ST <i>S</i> TA <i>M</i> EL <i>S</i> LR <i>S</i> DD <i>T</i> AVYY CAREADYFYFDYWGQGTT <i>L</i> TVSS <i>G</i> CGGG <i>K</i> VAAC <i>E</i> KEVAALKE KVAALKE <i>V</i> KAALKE	369
Pidilizumab, CT-011, MDV-9300 Heavy Chain (VH in Bold)	PD-1	<b>Q</b> VLVQSGSELKK <i>P</i> GA <i>S</i> V <i>K</i> IS <i>C</i> KA <i>S</i> GY <i>T</i> FT <i>N</i> YGM <i>N</i> W <i>V</i> R <i>Q</i> AP G <i>Q</i> GL <i>Q</i> WM <i>G</i> W <i>N</i> INT <i>D</i> SG <i>E</i> TY <i>A</i> EE <i>F</i> K <i>G</i> R <i>F</i> V <i>F</i> SL <i>D</i> T <i>S</i> V <i>N</i> T <i>A</i> Y <i>L</i> Q <i>I</i> <b>T</b> SL <i>T</i> AED <i>T</i> GM <i>Y</i> FC <i>V</i> R <i>G</i> Y <i>D</i> AL <i>D</i> W <i>Q</i> GT <i>L</i> TV <i>S</i> AST <i>K</i> GPS <i>V</i> F <i>P</i> LA <i>P</i> SS <i>K</i> ST <i>S</i> GG <i>T</i> AA <i>L</i> GL <i>V</i> K <i>D</i> Y <i>F</i> PE <i>P</i> PT <i>V</i> WS <i>N</i> GA <i>L</i> T <i>S</i> GV <i>H</i> TF <i>V</i> PA <i>V</i> L <i>V</i> Q <i>S</i> SG <i>L</i> Y <i>S</i> LS <i>S</i> V <i>V</i> T <i>V</i> P <i>S</i> SS <i>L</i> GT <i>Q</i> T <i>Y</i> LC <i>N</i> V <i>N</i> H <i>K</i> PS <i>N</i> T <i>K</i> DK <i>R</i> V <i>C</i> EP <i>K</i> SC <i>D</i> DK <i>T</i> HT <i>C</i> PP <i>C</i> PA <i>E</i> LL <i>G</i> GP <i>S</i> V <i>F</i> LF <i>PP</i> K <i>P</i> K <i>D</i> T <i>L</i> M <i>I</i> SR <i>T</i> PE <i>V</i> T <i>C</i> VV <i>V</i> D <i>V</i> SH <i>E</i> DP <i>E</i> VK <i>P</i> N <i>W</i> Y <i>D</i> G <i>V</i> E <i>V</i> H <i>N</i> A <i>K</i> T <i>K</i> P <i>R</i> EE <i>Q</i> Y <i>N</i> ST <i>Y</i> RV <i>V</i> V <i>L</i> TV <i>L</i> H <i>Q</i> D <i>W</i> L <i>N</i> KG <i>B</i> Y <i>K</i> C <i>K</i> V <i>S</i> N <i>K</i> AL <i>P</i> AP <i>E</i> KT <i>I</i> SK <i>A</i> KG <i>Q</i> RE <i>P</i> OV <i>V</i> Y <i>T</i> L <i>P</i> PS <i>R</i> EE <i>M</i> TK <i>N</i> Q <i>V</i> SL <i>T</i> CL <i>V</i> K <i>G</i> F <i>Y</i> PS <i>D</i> IAVE <i>W</i> ES <i>N</i> Q <i>P</i> ENNY <i>K</i> T <i>T</i> PP <i>V</i> L <i>D</i> S <i>G</i> FF <i>L</i> Y <i>S</i> KL <i>T</i> VD <i>K</i> SR <i>W</i> QQ <i>G</i> N <i>V</i> F <i>C</i> SV <i>M</i> HEAL <i>H</i> N <i>Y</i> T <i>Q</i> KS <i>L</i> SL <i>S</i> PG <i>K</i>	370
Pidilizumab, CT-011, MDV-9300 Light Chain (VL in Bold)	PD-1	<b>E</b> IVLTQSPSSLSAVGDRVTITCS <i>A</i> RSS <i>S</i> VS <i>Y</i> M <i>H</i> WF <i>Q</i> KPG <i>K</i> AP K <i>L</i> W <i>I</i> Y <i>R</i> TS <i>N</i> LA <i>S</i> GV <i>P</i> S <i>R</i> F <i>S</i> GS <i>G</i> CT <i>S</i> Y <i>C</i> LT <i>I</i> NS <i>L</i> Q <i>P</i> ED <i>F</i> AT <i>Y</i> <i>C</i> <b>Q</b> QRSS <i>F</i> PL <i>T</i> GG <i>G</i> TK <i>V</i> E <i>I</i> K <i>R</i> TV <i>A</i> APS <i>V</i> F <i>F</i> PP <i>S</i> DE <i>Q</i> L <i>K</i> SG <i>T</i> AS <i>V</i> CL <i>L</i> NN <i>F</i> Y <i>P</i> REAK <i>V</i> W <i>K</i> V <i>D</i> NAL <i>Q</i> CS <i>N</i> Q <i>E</i> S <i>V</i> TE <i>O</i> DS <i>K</i> D <i>T</i> SY <i>S</i> L <i>S</i> T <i>L</i> TL <i>S</i> KAD <i>Y</i> E <i>K</i> H <i>V</i> Y <i>A</i> CE <i>V</i> TH <i>Q</i> GL <i>S</i> SP <i>V</i> T <i>K</i> SF <i>N</i> R <i>G</i> EC	371
SG-001 VH	PD-1	QVQLVESGGVVQ <i>P</i> GR <i>S</i> L <i>R</i> LT <i>C</i> AS <i>G</i> LT <i>F</i> SS <i>S</i> GM <i>H</i> W <i>V</i> R <i>Q</i> AP <i>G</i> GLE <i>W</i> VA <i>V</i> I <i>W</i> D <i>G</i> K <i>R</i> YY <i>A</i> DS <i>V</i> K <i>G</i> R <i>F</i> T <i>I</i> SR <i>D</i> NS <i>K</i> N <i>T</i> LF <i>Q</i> M <i>N</i> SL <i>R</i> AED <i>T</i> AVYY <i>C</i> AT <i>NN</i> ND <i>Y</i> WGQ <i>G</i> TL <i>V</i> T <i>V</i> SS	372
SG-001 VL	PD-1	<b>E</b> IVLTQSPATL <i>S</i> LP <i>G</i> ERA <i>T</i> LS <i>C</i> RAS <i>Q</i> S <i>V</i> S <i>S</i> Y <i>L</i> AW <i>Y</i> QQ <i>K</i> PG <i>Q</i> AP <i>R</i> LL <i>I</i> Y <i>T</i> AS <i>N</i> R <i>A</i> T <i>G</i> i <i>P</i> AR <i>F</i> SG <i>S</i> GS <i>G</i> TD <i>F</i> TL <i>T</i> ISS <i>L</i> PE <i>D</i> FAVYY <i>C</i> Q <i>Y</i> NW <i>P</i> RT <i>F</i> G <i>Q</i> GT <i>K</i> VE <i>I</i> K	373
mpLZM-009 VH (Murine Precursor of LZM-009)	PD-1	EVQLQQSG <i>P</i> VL <i>V</i> K <i>P</i> GA <i>S</i> V <i>K</i> MS <i>C</i> AS <i>G</i> Y <i>T</i> FT <i>S</i> YY <i>M</i> Y <i>W</i> V <i>K</i> Q <i>S</i> H <i>G</i> K S <i>L</i> EW <i>IG</i> GV <i>N</i> PS <i>N</i> GG <i>T</i> N <i>F</i> NE <i>K</i> FK <i>S</i> K <i>A</i> TL <i>T</i> VD <i>K</i> SS <i>S</i> TA <i>M</i> EL <i>N</i> SL <i>T</i> ED <i>S</i> AVYY <i>C</i> ARR <i>D</i> Y <i>R</i> Y <i>D</i> M <i>G</i> F <i>D</i> Y <i>W</i> Q <i>G</i> TT <i>L</i> TV <i>SS</i>	374
mpLZM-009 VL (Murine Precursor of LZM-009)	PD-1	<b>Q</b> IVLTQSPA <i>M</i> S <i>A</i> P <i>G</i> E <i>K</i> V <i>T</i> M <i>T</i> CR <i>A</i> SK <i>G</i> V <i>S</i> T <i>S</i> GY <i>S</i> Y <i>L</i> HW <i>Y</i> QQ <i>K</i> P G <i>S</i> SP <i>R</i> LL <i>I</i> Y <i>L</i> AS <i>Y</i> LE <i>S</i> GV <i>P</i> V <i>R</i> F <i>S</i> GS <i>G</i> GT <i>S</i> Y <i>S</i> L <i>T</i> IR <i>M</i> EA <i>D</i> AT <i>Y</i> YC <i>Q</i> H <i>S</i> RE <i>L</i> PL <i>T</i> FG <i>T</i> GT <i>R</i> LE <i>I</i> K	375
LZM-009 VH	PD-1	QVQLVQSGAE <i>V</i> KK <i>P</i> GA <i>S</i> V <i>K</i> VS <i>C</i> AS <i>G</i> Y <i>T</i> FT <i>S</i> YY <i>M</i> Y <i>W</i> V <i>R</i> Q <i>A</i> PG Q <i>G</i> LE <i>W</i> VG <i>N</i> PS <i>N</i> GG <i>T</i> N <i>F</i> NE <i>K</i> FK <i>S</i> R <i>V</i> T <i>I</i> TAD <i>K</i> ST <i>S</i> TA <i>M</i> EL <i>S</i> SS R <i>SE</i> DT <i>A</i> VYY <i>C</i> ARR <i>D</i> Y <i>R</i> Y <i>D</i> M <i>G</i> F <i>D</i> Y <i>W</i> Q <i>G</i> TT <i>L</i> TV <i>SS</i>	376
LZM-009 VL	PD-1	<b>E</b> IVLTQSPATL <i>S</i> LP <i>G</i> ERA <i>T</i> LS <i>C</i> RAS <i>Q</i> S <i>V</i> S <i>S</i> Y <i>L</i> AW <i>Y</i> QQ <i>K</i> PG <i>Q</i> AP <i>R</i> Q <i>A</i> P <i>R</i> LL <i>I</i> Y <i>L</i> AS <i>Y</i> LE <i>S</i> GV <i>P</i> AR <i>F</i> SG <i>S</i> GS <i>G</i> GT <i>S</i> Y <i>S</i> L <i>T</i> IR <i>M</i> EA <i>D</i> AT <i>Y</i> Q <i>H</i> SR <i>E</i> LP <i>L</i> PL <i>T</i> FG <i>T</i> GT <i>K</i> VE <i>I</i> K	377
Budigalimab, 6VDO4TY300, ABBV-181, PR-	PD-1	<b>E</b> IQLVQSGAE <i>V</i> KK <i>P</i> GS <i>S</i> V <i>K</i> VS <i>C</i> AS <i>G</i> Y <i>T</i> FT <i>H</i> Y <i>G</i> M <i>N</i> W <i>V</i> R <i>Q</i> AP <i>R</i> Q <i>G</i> GLE <i>W</i> VG <i>W</i> V <i>N</i> TY <i>T</i> GE <i>P</i> TY <i>A</i> DD <i>F</i> K <i>G</i> R <i>L</i> T <i>F</i> L <i>D</i> T <i>S</i> TA <i>M</i> EL <i>N</i> SL <i>T</i> <b>L</b> SS <i>L</i> R <i>S</i> ED <i>T</i> AVYY <i>C</i> TR <i>E</i> GE <i>G</i> LG <i>F</i> GD <i>W</i> Q <i>G</i> TT <i>L</i> TV <i>S</i> AS <i>T</i> K <i>G</i> P SV <i>F</i> PL <i>A</i> PS <i>S</i> K <i>T</i> SG <i>G</i> TA <i>A</i> LG <i>C</i> LV <i>K</i> D <i>Y</i> F <i>P</i> EP <i>V</i> T <i>V</i> W <i>N</i> GA <i>L</i> T <i>S</i> GV <i>K</i>	378

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
1648817 Heavy Chain (VH in Bold)		TFPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTKVDK KVEPKSCDKTHTCPPCPAPEAAGGPSVFLFPPKPDKTLMISRTPEV TCVVVVDVSHEDEPKFVNWVVGVEVHNAKTKPREEQYNSTYRV VSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIKAKGQPREP QVYTLPPLPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPENN YKTTTPVLDSDGSFFLYSKLTVDKSRWQQGNVFCSVMEALHN HYTQKSLSLSPGE	
Budigalimab, 6VDO4TY300, ABBV- 181, PR- 1648817 Light Chain (VL in Bold)	PD-1	DVVMTOQSPLSLPVTPGEPASISCRSSQSIVHSHGDITLEWYLQ KPGQSPQLLIYKVSNRFSGVPDRFSGSGSGTDFTLKISRVEAED VGVIYCFQGSHIPVIFGGQGTKLEIKRTVAAPSVFIFPPSDEQLK GTASVVCLLNPFYPREAKVQWKVDNALQSGNSQESVTEQDSKD STYSLSSTLTL SKADYEKHKVYACEVT HQGLSSPVT KSFNR GEC	379
Lambrolizumab, Pembrolizumab, KEYTRUDA ®, MK- 3475, SCH- 900475, h409A11 VH CDR1	PD-1	NYYMY	380
Lambrolizumab, Pembrolizumab, KEYTRUDA ®, MK- 3475, SCH- 900475, h409A11 VH CDR2	PD-1	GINPSNGGTNFNEKFKN	381
Lambrolizumab, Pembrolizumab, KEYTRUDA ®, MK- 3475, SCH- 900475, h409A11 VH CDR3	PD-1	RDYRFDMGFDY	382
Lambrolizumab, Pembrolizumab, KEYTRUDA ®, MK- 3475, SCH- 900475, h409A11 VL CDR1	PD-1	RASKGVSTSGYSYLN	383
Lambrolizumab, Pembrolizumab, KEYTRUDA ®, MK- 3475, SCH- 900475, h409A11 VL CDR2	PD-1	LASYLES	384
Lambrolizumab, Pembrolizumab, KEYTRUDA ®, MK- 3475, SCH- 900475, h409A11 VL CDR3	PD-1	QHSRDLPLT	385

TABLE 1-continued

<u>Exemplary Antibodies for Immune Cell Associated Antigens</u>			
Antibody or Ag- binding fragment	Antigen Bound	Sequence	SEQ ID NO
Nivolumab, Nivolumab BMS, OPDIVO ®, BMS- 936558, MDX-1106, ONO-4538 VH CDR1	PD-1	NSGMH	386
Nivolumab, Nivolumab BMS, OPDIVO ®, BMS- 936558, MDX-1106, ONO-4538 VH CDR2	PD-1	VIWYDGSKRYYADSVKG	387
Nivolumab, Nivolumab BMS, OPDIVO ®, BMS- 936558, MDX-1106, ONO-4538 VH CDR3	PD-1	NDDY	388
Nivolumab, Nivolumab BMS, OPDIVO ®, BMS- 936558, MDX-1106, ONO-4538 VL CDR1	PD-1	RASQSVSSYLA	389
Nivolumab, Nivolumab BMS, OPDIVO ®, BMS 936558, MDX-1106, ONO-4538 VL CDR2	PD-1	DASN RAT	390
Nivolumab, Nivolumab BMS, OPDIVO ®, BMS- 936558, MDX-1106, ONO-4538 VL CDR3	PD-1	QQSSNWPRT	391
Serplulimab, HLX-10 VH CDR1	PD-1	FTFSNYGMS	392
Serplulimab, HLX-10 VH CDR2	PD-1	TISGGGSNIY	393
Serplulimab, HLX-10 VH CDR3	PD-1	VSYYYGIDF	394

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Serplulimab, HLX-10 VL CDR1	PD-1	KASQDVTTAVA	395
Serplulimab, HLX-10 VL CDR2	PD-1	WASTRHT	396
Serplulimab, HLX-10 VL CDR3	PD-1	QQHYTIPWT	397
SG-001 VH CDR1	PD-1	GLTFSSSG	398
SG-001 VH CDR2	PD-1	IWYDGSKR	399
SG-001 VH CDR3	PD-1	ATNNDY	400
SG-001 VL CDR1	PD-1	RASQSVSSYLA	401
SG-001 VL CDR2	PD-1	TASN RAT	402
SG-001 VL CDR3	PD-1	QQYSNWPRT	403
PD-1-Fc- OX40L (Code) , SL- 279252 (Code) , TAK-252 (Code)	PD-1	MQIQPAPWPWAVLQLGWRPGFWLDSPDRPWNPTFSPALLVV TEGDNATFTCSFSNTSESFPVNLWYRMSPSNQTDKLAAPFEDRSQP GQDCRFRVTQLPNGRDFHMSVVRARRNDSGTLYCGAISLAPKAQ IKESLRAELRVTERRAEVPTAHPSPSPRPAGQFQSKYGPCCPSCPA PEFLGGPSVFLFPKPKDTLMISRTPEVTCWVDVSQEDPEVQFW YVDGVEVHNAKTKPREEQFNSTYRVVSVLTVLHQDWLSGKEYK CKVSSKGLPSSIEKTISNATGQPREPQVYTLPPSQEEMTQNVS LT CLVKGFYPSDIAVEWESNGOPENNYKTPPVLDGSFFLYSRLT VDKSSWQEGNVFSCVMHEALHNHYTQKSLSLKGIEGRMDQ VSHRYPRIQSIIKVQFTEYKKEKGFI LTSQKEDEIMKVNQNSVI INC DGFYLISLKGYFSQEVNISLHYQKDEEPLFQLKKVRSVNSLMVAS LTYDKVYLNVTDTNTSLDDFHVNNGELILIHQNPGEFCVLMQIP QAPWPWWAVLQLGWRPGFWLDSPDRPWNPTFSPALLVVTEGD NATFTCSFSNTSESFPVNLWYRMSPSNQTDKLAAPFEDRSQPQGDC RFRVQLPNGRDFHMSVVRARRNDSGTLYCGAISLAPKAQIKESL RAELRVTERRAEVPTAHPSPSPRPAGQFQVSHRYPRIQSIIKVQFT EYKEKGFI LTSQKEDEIMKVNQNSVI INCDGFYIISLKGYFSQEV NISLHYQKDEEPLFQLKKVRSVNSLMVASLTYKDVKVYLNVTTDN TSLLDFHVNGGELILIHQNPGEFCVL	404
Avelumab (Generic) Bavencio (Brand) 451238 KXG2PJ551I MSB-0010682 MSB-0010718C PF-06834635 Heavy Chain (VH in Bold)	PD-L1	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYIMMWVRQAPG KGLEWVSSIYPGGITFYADTVKGRFTISRDNSKNTLYLQMNS LRAEDTAVYYCARIKLGTVTTVDYWGQGTLVTVS SASTKGPS VFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVWSNSGALTSGVHT FPAVLQSSGLYSSLSSVTVPSSSLGTQTYICNVNHKPSNTKVDKK VEPKSCDKTHTCP CPCPAPEELLGGPSVFLFPPKPKDTLMISRTPEV CVVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVV SVLTVLHQDWLNGEYKCKVSNKALPAPI EKTISKAKGQPREPQ VYTLPPSRDELTKNQVSLTCLVKGFYPSDIAVEWESNGOPENNYK TTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFSCVMHEALHNHY TQKSLSLSPGK	232

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Avelumab (Generic) Bavencio (Brand) 451238 KXG2PJ551I MSB-0010682 MSB-0010718C PF-06834635 Light Chain (VL in Bold)	PD-L1	QSALTQPASVSGSPGQSITISCTGTSSDVGGYNVWSYQQHPG KAPKLMIYDVSNRPSGVSNRFSGSKSGNTASLTISGLQAEDEA DYYCSSYTSSSTRVFGTGTKVTLGQPKANPTVTLFPPSSEELQA NKATLVCLISDFYFPGAVTVAKADGSPVKAGVETTKPSKQSNNK YAASSYLSLTPEQWKSHRSYSQCQVTHEGSTVEKTVAPETECS	233
Durvalumab (Generic) Imfinzi (Brand) 28X28X9OKV (UNII code) MEDI-4736 Heavy Chain (VH in Bold)	PD-L1	EVQLVESGGGLVQPGGSLRLSCAASGFTFSRYWMSWVRQAP GKGLIEWVANIKQDGSEKYYVDSVKGRFTISRDNAKNSLYLQ MNSLRAEDTAVYYCAREGGWFGEIAFDYWGQGTLVTVSSAS TKGPSPVFLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSGALT SGVHTFPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTK VDKRVEPKSCDKTHTCPPCPAPEFEFGGPSPVLFPPPKDTLMSRT PEVTCVVVDVSHEDPEVKPNWYVGVEVHNAKTKPREEQYNST YRVSVLTVLHQDWLNGKEYKCKVSNKALPASIEKTISKAKGQP REPVQYTLPPSREEMTKNQVSLSCLVKGFYPSDIAVEWESNGOPE NNYKTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEAL HNHYTQKSLSLSPKG	234
Durvalumab (Generic) Imfinzi (Brand) 28X28X9OKV (UNII code) MEDI-4736 Light Chain (VL in Bold)	PD-L1	EIVLTQSPGTLSLSPGERATLSCRASQRVSSSYLAWYQQKPGQ APRLLIYDASSRATGIPDRFSGSGSGTDFTLTISLEPEDFAVYY CQQYGS LPWTFGQGTTKVEIKRTVAAPSVFIFPPSDEQLKSGTAS VVCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYS LSSTLTL SKADYEKKVYACEVTHQGLSSPVTKSFNRGEC	235
Atezolizumab (Generic) Tecentriq (Brand) 52CM10WC3Y MPDL-3280A RG-7446 RO-5541267 Heavy Chain (VH in Bold)	PD-L1	EVQLVESGGGLVQPGGSLRLSCAASGFTFSDSWIHWVRQAPG KGLEWVAISPYGGSTYYADSVKGRFTISADTSKNTAYLQMN SLRAEDTAVYYCARRHWPGGF DYWGQGTLVTVSSASTKGPS VFPLAPSSKSTSGGTAALGCLVKDYFPEPVTVSWNSGALTSGVHT FPAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTKVDK VEPKSCDKTHTCPPCPAPEELLGGPSVFLFPPPKDTLMSRTPEV CVVVDVSHEDPEVKPNWYVGVEVHNAKTKPREEQYASTYRVV SVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTISKAKGQPREPQ VYTLPPSREEMTKNQVSLSCLVKGFYPSDIAVEWESNGOPE KTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHN YTQKSLSLSPKG	236
Atezolizumab (Generic) Tecentriq (Brand) 52CM10WC3Y MPDL-3280A RG-7446 RO-5541267 Light Chain (VL in Bold)	PD-L1	DIQMKTQSPSSL SASVGDRV TITCRASQDV STAVAWYQQKPGK APKLLIYASFLYSGVPSRFSFGSGSGTDFTLTISLQPEDFATYY CQQYLHPATFGQGTTKVEIKRTVAAPSVFIFPPSDEQLKSGTAS VVCLLNNFYPREAKVQWKVDNALQSGNSQESVTEQDSKDSTYS LSSTLTL SKADYEKKVYACEVTHQGLSSPVTKSFNRGEC	237
Sugemalimab (Generic) CS-1001 WBP-3155	PD-L1	EVQLLESGGGLVQPGGSLRLSCAASGFTFSSYAMSWVRQAPG KGLEWVSGISGSGFTYYADSVKGRFTISRDNSKNTLYLQMN SLRAEDTAVYYCAKPPRGYNYGPF DYWGQGTLVTVSSASTKG PSVFPLAPCSRSTSEESTAALGCLVKDYFPEPVTVSWNSGALTSGV	238

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
Heavy Chain (VH in Bold)		HTPPAVLQSSGLYSLSSVTVPPSSLGTTKTYTCNVDHKPSNTKVD KRVESKYGPPCPCPAPEFLGGPSVFLFPPKPKDTLMISRTPEVTC VVVIVDSQEDPEVQPNWYDVGVEVHNAKTKPREEQFNSTYRVVS VLTVLHQDWLNGKEYKCKVSNKGLPSSIEKTISKAKGQPREPQV YTLPSPSQEEMTKNQVSLTCLVKGYPSPDIAVEWESENQOPENNYK TPPVLDSDGSFFLYSRSLTVDKSRWQEGNVFSCSVMHEALHNHY TQKSLSLSLGK	
Sugemalimab (Generic) CS-1001 WBP-3155 Light Chain (VL in Bold)	PD-L1	SYVLTQPPSVSVAQGQTARITCGGNНИGSKSVHWWQQKPGQA PVLVVYDDSDRPSGIPERFSGNSGNTATLTISRVEAGDEADYY CQVWDSSSDHVVFGGGTKLTVLGQPKAAPSVTLFPPSSEELQA NKATLVCLISDFYPGAVTVAWKADSSPVKAGVETTPSKQSNNK YAASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPTECS	239
JS-003 JS003-30 JS003-SD VH	PD-L1	QGQLQESGPSPSLVKPKSQTLSLTCTVSGDSITRGYWNVIRKHKGKL EYIGYIYSTYGTYSNLSLKSRVIIIRDTSKNQYYLKLSSVTAADTA VYYCATSTGWLDPPDVYWGQGTLTVSS	242
JS-003 JS003-30 JS003-SD VL	PD-L1	DIVMTQSPDSLAVSLGERATINCKASQNVDTSVAWFQQKPGQPP KALIYSASFRYSGVPDRFSGSGSTDFLTITSSLQAEDVAVYFCQQ YYGYPFTFGQQGKLEIK	243
HLX-20 PL2#3 VH	PD-L1	EVQLVQSGGGLVKPGGSLRLSCAASGFTFSSYTMNWVRQAPKG GLEWVSSIIGSDYLYYADSVKGRTFISRDNAKNSLYLQMNSLRA EDTAVYYCARNELRWPQAGAFDRWGQGTMVTVSS	244
HLX-20 PL2#3 VL	PD-L1	QSVVTQPPSMSAAPGQRVTISCGSSSYIESSYVGWYQQLPGTAP RLLIYDDMRPSGIPDRFSGSKSGTSATLAITGLQTGDEADYYCEI WRSLGLGVFGGGTKLTVL	245
Lodapolimab (Generic) LY-3300054 NR4MAD6 PPB Heavy Chain (VH in Bold)	PD-L1	QVQLVQSGAEVKKPGSSVKVSCKASGGTFSSYAIISWVRQAPG QGLEWMGGIIPIFGTANYAQKFQGRVITADKSTSTAYMELESS LRSEDTAVYYCARSPDYSPYYYYGMDVWGQGTTVTVSSASTK GPSVPLAPSSKSTSGGAALGCLVKDVFPEPVTVWSNSGALTSG VHTFPVAVLQSSGLYSLSSVVTVPSSSLGTQTYICNVNHKPSNTKV DKRVEPKSCDKTHTCPPCPAPEAEGAPSVFLFPPKPKDTLMISRTP EVTCVVVDVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTY RVVSVLTVLHQDWLNGKEYKCKVSNKALPSSIEKTISKAKGQPR EPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESENQOPEN NYKTPPVLDSDGSFFLYSKLTVDKSRWQOGNVFSCSVMHEALH NHYTQKSLSLSPGK	248
Lodapolimab (Generic) LY-3300054 NR4MAD6 PPB Light Chain (VL in Bold)	PD-L1	QSVLTQPPSASGTPGQRVTISCGSSSNIGSNTVNWYQQLPGT APKLLIYGNNSNRPSPDRFSGSKSGTSASLAIISGLQSEDEADYY YCQSYDSSLGSGVFGGGIKLTVLGQPKAAPSVTLFPPSSEELQAN KATLVCLISDFYPGAVTVAWKADSSPVKAGVETTPSKQSNNKY AASSYLSLTPEQWKSHRSYSCQVTHEGSTVEKTVAPAEC	249
HLX-20 PL2#3 VH CDR1	PD-L1	SYTMN	250
HLX-20 PL2#3 VH CDR2	PD-L1	SISSGSDYLYYADSVKG	251
HLX-20 PL2#3 VH CDR3	PD-L1	NELRWYPQAGAFDR	252
HLX-20 PL2#3 VL CDR1	PD-L1	SGSSSYIESSYVG	253
HLX-20 PL2#3 VL CDR2	PD-L1	DDDMRPS	254

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens			
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO
HLX-20 PL2#3 VL CDR3	PD-L1	EIWRSGLGGV	255
Envafolimab (Generic) 3D-025 ASC-22 KN-035 hu56V1-Fc-m1 single-domain antibody (VH in Bold)	PD-L1	QVQLVESGGGLVQPGGSLRLSCAASGKMSRRCMAWFRQAP GKERERVAKLLTTSGSTYLADSVKGRFTISRDNSKNTVYLOM NSLRAEDTAVYYCAADSFEDPTCTLTSSGAFQYWQGQTLVT VSSEPKSSDKTHTCPGCPAPEELLGGPSVFLFPKPDKTLMISRPEV TCVVVAVSHEDPEVKFNWYVGVEVHNNAKTKPREEQYNSTYRV VSVLTVLHQDWLNKEYCKVSNKALPAGIEKTISKAKGQPREP QVYTLPSSRDELTKNQVSLTCLVKGFYPSDIAVEWESNGQOPENNY KTTTPVLDSDGSFPLYSKLTVDKSRWQQGNVFSCSVMHEALHNH YTQKSLSLSPGK	256
ND-021 NM21-1480 PRO-1480 Tri-specific fusion single-chain antibody construct	PD-L1	DIQMTQSPASLSASAVGDRVITTCQASQSIGTYLAQKPGKPK LLIYRAFILASGVPSRFSGSGSGTDFLTITSSLQPEDFATYYCQSNF YSDSTTIGPNAFGTGTKVTVLGGGGSEVQLVESGGLVQPGGS LRLSCAASGFSFSANYYPCWVRQAPGKLEWIGCIYGGSDITYD ANWTKGRFTISRDNSKNTVYLOQMSLRAEDTAVYYCARSAWYS GWGDLWGQCTLTVSSGGGGGGGGGGGGGGGGGGSIQMTO SPSSLSASVGDRVITTCQASQSIQNRLAWYQQKPGKAPKLLIYAS TLASGVPSRFSRGSGSGTDFLTITSSLQPEDFATYYCQSTYYGNDGN AEGTGTKVTVLGGGGSEVQLVESGGGLVQPGGSRLSCAASGF SFNSDYWIYWVRQAPGKLEWIASIYGGSSGNTQYASWAQGRFT ISRDNSKNTVYLOQMSLRAEDTAVYYFCARGYVDYGGATDLWGQ GTLTVSSGGGGGGGGSIQMTOQSPSSLSASVGDRVITTCQSSSEV YSNQLSWYQQKPCOPPKLIYDASDLASGVPSRFSGSGSGTDF LTISLQPEDFATYYCAGGGSSSSDTAFGGGKTLTVLGGGGGG GGGGGGGGGGSEVQLVESGGGLVQPGGSRLSCAASGFSLSS NAMGWRQAPGKGLEYIGIISVGGFTYYASWAKGRFTISRDNSK NTVYLOQMSLRAEDTATYPCARDRHGGDSSGAFYLWGQGTLVT VSS	257
ACE-05 YBL-013 Bispecific tetrameric antibody-like cell engager (ALICE) comprising two identical light chains (LC) consisting of antigen binding domains (ABDs) targeting programmed cell death-ligand 1 (PD-L1), and two different heavy chain (HC)-like chains (ACE-05-VL) each consisting of an anti-PD-L1 ABD and an anti-CD3 ABD; wherein	PD-L1	QMQLVQSGAEVKKPGSSVKVSCKASGGTFSSYAIswvrQAPGQG LEWGRRIPILGIANYAQKFQGRVTITADKSTSTAYMELSSLRS TAVYYCAKPRDGYNLVAFDIWGQTMVTVSSASTKGPSPVPLAP SSSKTSGGTAALGCLVKDVFPEPVTVSWNSGALTSGVHTFP QSSGLYSLSVVTVTPSSSLGTQTYICNVNHKPSNTKVDKKVEPK CDKHTCPGCPAPEELLGGPGGGSEVQLQQSGPELVKPGPSMKIS CKASGYSFTGYTMNWVKQSHGKNLLEMGLINPYKGVSTYNQK FKDKATLTVDKSSSTAYMELLSTSLEDASVYYCARSGYYGDSDW YFDVGQGTTLVFSQMLVQSGAEVKKPGSSVKVSCKASGGT FSSYAIswvrQAPQGLEWMGRIPILGIANYAQKFQGRVTITAD KSTSTAYMELSSLRSEDTAVYYCAKPRDGYNLVAFDIWGQGTM VTVSSASTKGPSPVFLAPSSKSTGGTAALGCLVKDVFPEPVTV WNSGALTSGVHTFPAVLQSSGLYLSLSSVTVPPSSSLGTQTYICNV NHKPSNTKVDKKVEPKSCDKTHTCPGCPAPEELLGGPGGGSDIQ MTQTSSLSASLGDRVTISCRASQDIRNYLNWYQQKPDGTVKLLI YYTSRLHSGVPSKESFGSGSGTDYSLTISNLQEDIAFYFCQQGNTL PWTFAAGGKLEIKRQLVLTQPPSVSGAPGQRVTISCTGSSSNIGAG YDVHWWYQQLPGAAPKLLIYGDINRPSGVDRFSGSKSGISASLAI GLQAEDEADYYCQSYDSSLSGGVFGGGKLTVLRTVAAPSVFIFP PSDEQLKSGTASVVCCLNNNFYPREAKVQWKVDNALQSGNSQESV TEQDSKDSTYSLSTTLSKADYEHKVYACEVTHQGLSSPVTKS FNRGEC	258

TABLE 1-continued

Exemplary Antibodies for Immune Cell Associated Antigens				
Antibody or Ag-binding fragment	Antigen Bound	Sequence	SEQ ID NO	
each HC comprises a G4S linker between the hinge region and the second ABD				
			15	
Modification to Fc Region				
Disclosed herein are antibodies or antigen binding fragments thereof that comprise an Fc region, wherein the Fc region comprises at least one covalently linked linker. In some embodiments, the linker is a chemical linker. In some embodiments, the chemical linker is covalently attached to a tyrosine, aspartic acid, glutamic acid, arginine, histidine, or lysine residue. In some embodiments, the chemical linker is covalently attached to a lysine, cysteine, or tyrosine residue. In some embodiments, the chemical linker is covalently attached to a cysteine residue. In some embodiments, the chemical linker is covalently attached to a lysine residue. In some embodiments, the chemical linker is covalently attached to a constant region of the antibody.				
In some embodiments, the antibody comprises an Fc region. In some embodiments, the Fc region is an IgG Fc region, an IgA Fc region, an IgD Fc region, an IgM Fc region, or an IgE Fc region. In some embodiments, the Fc region is an IgG Fc region, an IgA Fc region, or an IgD Fc region. In some embodiments, the Fc region is a human Fc region. In some embodiments, the Fc region is a humanized Fc region. In some embodiments, the Fc region is an IgG Fc region. In some instances, an IgG Fc region is an IgG1 Fc region, an IgG2a Fc region, or an IgG4 Fc region.				
One or more mutations may be introduced in an Fc region to reduce Fc-mediated effector functions of an antibody or antigen-binding fragment such as, for example, antibody-dependent cellular cytotoxicity (ADCC) and/or complement function. In some instances, a modified Fc comprises a humanized IgG4 kappa isotype that contains a S229P Fc mutation. In some instances, a modified Fc comprises a human IgG1 kappa where the heavy chain CH2 domain is engineered with a triple mutation such as, for example: (a) L238P, L239E, and P335S; or (2) K248; K288; and K317.				
In some embodiments, the Fc region has an amino acid sequence at least 80%, at least 85%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% identical to a sequence as set forth in SEQ ID NO: 260 (Pro Lys Ser Cys Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Prol Glu Xaa Xaa Gly Xaa Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Asp Val Ser His Glu Asp Pro Glu Val			15	
			Lys Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asp Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Xaa Glu Xaa Thr Lys Asn Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Xaa Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly, where Xaa can be any naturally occurring amino acid). In some embodiments, the Fc region comprises one or more mutations which make the Fc region susceptible to modification or conjugation at a particular residue, such as by incorporation of a cysteine residue at a position which does not contain a cysteine in SEQ ID NO: 260. Alternatively, the Fc region could be modified to incorporate a modified natural amino acid or an unnatural amino acid which comprises a conjugation handle, such as one connected to the modified natural amino acid or unnatural amino acid through a linker. In some embodiments, the Fc region does not comprise any mutations which facilitate the attachment of a linker to an additional cytokine (e.g., an IL-18 polypeptide). In some embodiments, the chemical linker is attached to a native residue as set forth in SEQ ID NO: 260. In some embodiments, the chemical linker is attached to a native lysine residue of SEQ ID NO: 260.	35
			In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at one of positions 10-90 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at one of positions 10-20, 10-30, 10-40, 10-50, 10-60, 10-70, 1-80, 10-90, 10-100, 10-110, 10-120, 10-130, 10-140, 10-150, 10-160, 10-170, 10-180, 10-190, or 10-200 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at one of positions 10-30, 50-70, or 80-100 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at any one of positions 20-40, 65-85, or 90-110 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc	40
				60

**59**

region at an amino acid residue at one of positions 15-26, 55-65, or 85-90 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at any one of positions 25-35, 70-80, or 95-105 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at any one of positions 30, 32, 72, 74, or 101 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at an amino acid residue at any one of positions K30, K32, K72, K74, or K101 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at amino acid residue 30 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at amino acid residue 32 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at amino acid residue 72 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at amino acid residue 74 of SEQ ID NO: 260. In some embodiments, the chemical linker is attached to the Fc region at amino acid residue 101 of SEQ ID NO: 260.

The chemical linker can be covalently attached to one amino acid residue of an Fc region of the antibody. In some embodiments, the chemical linker is covalently attached to a non-terminal residue of the Fc region. In some embodiments, the non-terminal residue is in the CH1, CH2, or CH3 region of the antibody. In some embodiments, the non-terminal residue is in the CH2 region of the antibody.

In some embodiments, the chemical linker is covalently attached at an amino acid residue of the antibody or antigen binding fragment which selectively binds a immune associated antigen (e.g., an anti-PD-1 antibody) such that the function of the antibody or antigen binding fragment is maintained (e.g., without denaturing the polypeptide). For example, when the antibody or antigen binding fragment is a human IgG (e.g., human IgG1), exposed lysine residues and exposed tyrosine residues are present at the following positions (refer to web site [www.imgt.org/IMGTScientific-Chart/Numbering/HuIGHGnber.html](http://www.imgt.org/IMGTScientific-Chart/Numbering/HuIGHGnber.html) by EU numbering). Exemplary exposed Lysine Residues: CH2 domain (position 246, position 248, position 274, position 288, position 290, position 317, position 320, position 322, and position 338) CH3 domain (position 360, position 414, and position 439). Exemplary exposed Tyrosine Residues: CH2 domain (position 278, position 296, and position 300) CH3 domain (position 436).

The human IgG, such as human IgG1, may also be modified with a lysine or tyrosine residue at any one of the positions listed above in order provide a residue which is ideally surface exposed for subsequent modification.

In some embodiments, the chemical linker is covalently attached at an amino acid residue in the constant region of an antibody. In some embodiments, the chemical linker is covalently attached at an amino acid residue in the CH1, CH2, or CH3 region. In some embodiments, the chemical linker is covalently attached at an amino acid residue in the CH2 region. In some embodiments, the chemical linker may

**60**

be covalently attached to one amino acid residue in the following groups of residues following EU numbering in human IgG Fc: amino acid residues 1-478, amino acid residues 2-478, amino acid residues 1-477, amino acid residues 2-477, amino acid residues 10-467, amino acid residues 30-447, amino acid residues 50-427, amino acid residues 100-377, amino acid residues 150-327, amino acid residues 200-327, amino acid residues 240-327, and amino acid residues 240-320.

In some embodiments, the chemical linker is covalently attached to one lysine residue of a human IgG Fc region. In some embodiments, the chemical linker is covalently attached at Lys 246 of an Fc region of the antibody, wherein amino acid residue position number is based on Eu numbering. In some embodiments, the chemical linker is covalently attached at Lys 248 of an Fc region of the antibody, wherein amino acid residue position number is based on Eu numbering. In some embodiments, the chemical linker is covalently attached at Lys 288 of an Fc region of the antibody, wherein amino acid residue position number is based on Eu numbering. In some embodiments, the chemical linker is covalently attached at Lys 290 of an Fc region of the antibody, wherein amino acid residue position number is based on Eu numbering. In some embodiments, the chemical linker is covalently attached at Lys 317 of the antibody, wherein amino acid residue position number is based on Eu numbering.

The chemical linker can be covalently attached to an amino acid residue selected from a subset of amino acid residues. In some embodiments, the subset comprises two three, four, five, six, seven, eight, nine, or ten amino acid residues of an Fc region of the antibody. The chemical linker can be covalently attached to one of two lysine residues of an Fc region of the antibody.

In some embodiments, the antibody will comprise two linkers covalently attached to the Fc region of the antibody. In some embodiments, each of the two linkers will be covalently attached to a different heavy chain of the antibody. In some embodiments, each of the two linkers will be covalently attached to a different heavy chain of the antibody at a residue position which is the same. In some embodiments, each of the two linkers will be covalently attached to a different heavy chain of antibody at a residue position which is different. When the two linkers are covalently attached to residue positions which differ, any combination of the residue positions provided herein may be used in combination.

In some embodiments, a first chemical linker is covalently attached at Lys 248 of a first Fc region of the antibody, and a second chemical linker is covalently attached at Lys 288 of a second Fc region of the antibody, wherein amino acid residue position number is based on Eu numbering. In some embodiments, a first chemical linker is covalently attached at Lys 248 of a first Fc region of the antibody, and a second chemical linker is covalently attached at Lys 317 of a second Fc region of the antibody, wherein residue position number

**61**

is based on Eu numbering. In some embodiments, a first chemical linker is covalently attached at Lys 288 of a first Fc region of the antibody, and a second chemical linker is covalently attached at Lys 317 of a second Fc region of the antibody, wherein amino acid residue position number is based on Eu numbering.

#### Method of Modifying an Fc Region

Also provided herein are method of preparing a modified Fc region of an antibody or antigen binding fragment, such as for the attachment of a linker, a conjugation handle, the IL-18 polypeptide, or any combination thereof to the antibody or antigen binding fragment. A variety of methods for site-specific modification of Fc regions of antibodies are known in the art.

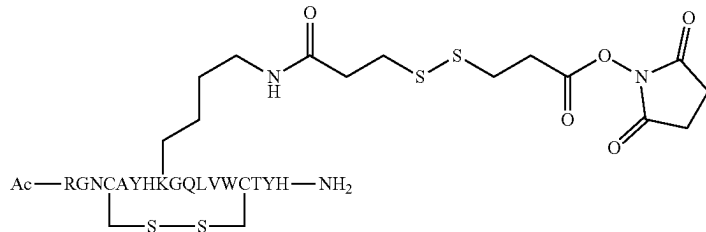
#### Modification with an Affinity Peptide Configured to Site-Specifically Attach Linker to the Antibody

In some embodiments, an Fc region is modified to incorporate a linker, a conjugation handle, or a combination thereof. In some embodiments, the modification is performed by contacting the Fc region with an affinity peptide bearing a payload configured to attach a linker or other group to the Fc region, such as at a specific residue of the Fc region. In some embodiments, the linker is attached using a reactive group which forms a bond with a residue of the Fc region. In some embodiments, the affinity peptide comprises a cleavable linker. The cleavable linker is configured on the affinity peptide such that after the linker or other group is

**62**

selected from (1) QETNPTENLYFQQKQNMQCQRRF-YEALHDPNLNEEQRNARIRSIRDDDC (SEQ ID NO: 261); (2) QTADNQKQNMQCQRRFYEAHDPNLNEEQRNARIRSIRDDCSQSANLLAEAQQLNDAQA PQA (SEQ ID NO: 262); (3) QETKNMQCQRRFYEAHDPNLNEEQRNARIRSIRDDDC (SEQ ID NO: 263); (4) QETFNKQCQRRFYEAHDPNLNEEQRNARIRSIRDDDC (SEQ ID NO: 264); (5) QETFNMQCQRRF-YEALHDPNLNEEQRNARIRSIRDDDC (SEQ ID NO: 265); (6) QETFNMQCQRRFYEAHDPNLNEEQRNARIRSIRDKDC (SEQ ID NO: 266); (7) QETMQCQRRF-YEALHDPNLNEEQRNARIRSIRDKDC (SEQ ID NO: 267); (8) QETQCQRRFYEAHDPNLNEEQRNARIRSIRDKDC (SEQ ID NO: 268); (9) QETCQRRF-YEALHDPNLNEEQRNARIRSIRDKDC (SEQ ID NO: 269); (10) QETRGNCAHKGQLVWCTYH (SEQ ID NO: 270); and (11) QETRGNCAHKGQIIWCTYH (SEQ ID NO: 271), or a corresponding peptide which has been truncated at the N-terminus by one, two, three, four, or five residues.

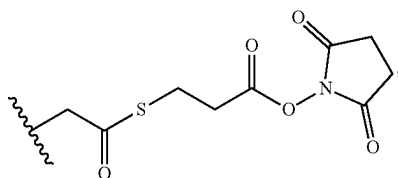
An exemplary affinity peptide with cleavable linker and conjugation handle payload capable of attaching the payload to residue K248 of an antibody as provided herein is shown below (as reported in Matsuda et al., "Chemical Site-Specific Conjugation Platform to Improve the Pharmacokinetics and Therapeutic Index of Antibody-Drug Conjugates," *Mol. Pharmaceutics* 2021, 18, 11, 4058-4066).



attached to the Fc region, the affinity peptide can be removed, leaving behind only the desired linker or other group attached to the Fc region. The linker or other group can then be used further to add attach additional groups, such as a cytokine or a linker attached to a cytokine, to the Fc region.

Non-limiting examples of such affinity peptides can be found at least in PCT Publication No. WO2018199337A1, PCT Publication No. WO2019240288A1, PCT Publication No. WO2019240287A1, and PCT Publication No. WO2020090979A1, each of which is incorporated by reference as if set forth herein in its entirety. In some embodiments, the affinity peptide is a peptide which has been modified to deliver the linker/conjugation handle payload one or more specific residues of the Fc region of the antibody. In some embodiments, the affinity peptide has at least 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or 100% sequence identity to a peptide

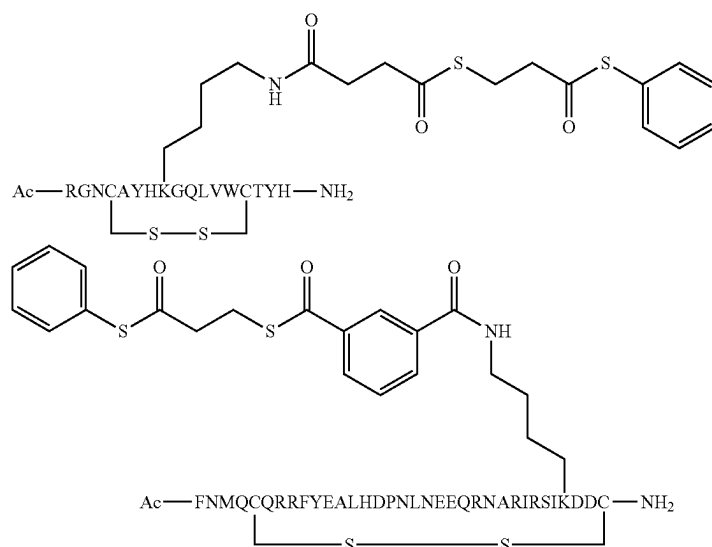
Alternative affinity peptides targeting alternative residues of the Fc region are described in the references cited above for AJICAP™ technology, and such affinity peptides can be used to attach the desired functionality to an alternative residue of the Fc region (e.g., K246, K288, etc.). For example, the disulfide group of the above affinity peptide could instead be replaced with a thioester to provide a sulfhydryl protecting group as a cleavable portion of the linking group (e.g., the relevant portion of the affinity peptide would have a structure of



or another of the cleavable linkers discussed below). Such alternative affinity peptides include those described in, for

63

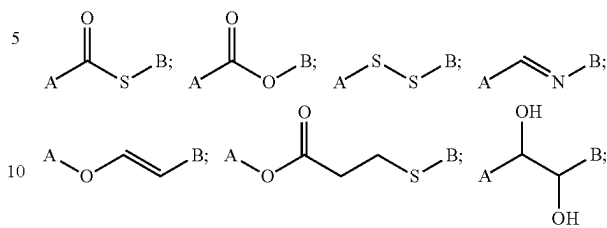
example “AJICAP Second Generation: Improved Chemical Site-Specific Conjugation Technology for Antibody-Drug Conjugation Technology for Antibody-Drug Conjugate Production” (Working Paper, Fujii et al., DOI: 10.26434/chemrxiv-2023-9p5p7, chemrxiv.org/engage/chemrxiv/article-details/63d5f7131125965a9e7df8a5 (Accessed 20 Feb. 2023, Version 1 published 30 Jan. 2023)). Exemplary affinity peptides provided therein include those shown below, wherein the left structure targets K248 of the Fc region and the right structure targets K288 of the Fc region (EU numbering).



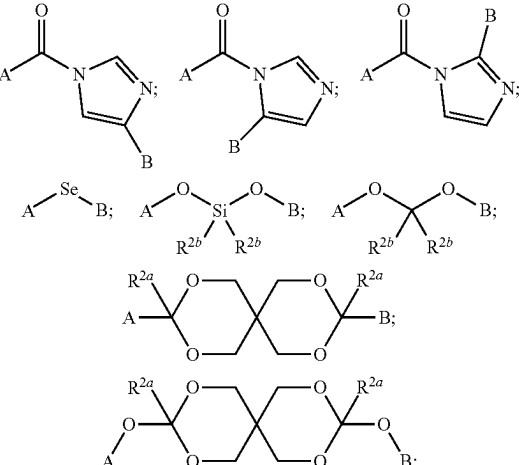
The affinity peptide of the disclosure can comprise a cleavable linker. In some embodiments, the cleavable linker of the affinity peptide connects the affinity peptide to the group which is to be attached to the Fc region and is configured such that the peptide can be cleaved after the group comprising the linker or conjugation handle has been attached. In some embodiments, the cleavable linker is a 40  
divalent group. In some embodiments, the cleavable linker can comprise a thioester group, an ester group, a sulfane group; a methanimine group; an oxyvinyl group; a thiopropanoate group; an ethane-1,2-diol group; an (imidazole-1-yl)methan-1-one group; a seleno ether group; a silylether group; a di-oxysilane group; an ether group; a di-oxymethane group; a tetraoxospiro[5.5]undecane group; an acetamidoethyl phosphoramidite group; a bis(methylthio)-pyrazolopyrazole-dione group; a 2-oxo-2-phenylethyl formate group; a 4-oxybenzylcarbamate group; a 2-(4-hydroxy-oxy-phenyl)diazinyl)benzoic acid group; a 4-amino-2-(2-amino-2-oxoethyl)-4-oxobut-2-enic acid group; a 2-(2-methylenehydrazinyl)pyridine group; an N<sup>1</sup>-methyleneformohydrazide group; or an isopropylcarbamate group, any of which is unsubstituted or substituted. 45  
Composition and points of attachment of the cleavable linker to the affinity peptide, as well as related methods of use, are described in, at least, PCT Publication No. WO2018199337A1, PCT Publication No. 50  
WO2019240288A1, PCT Publication No. 55  
WO2019240287A1, and PCT Publication No. 60  
WO2020090979A1.

64

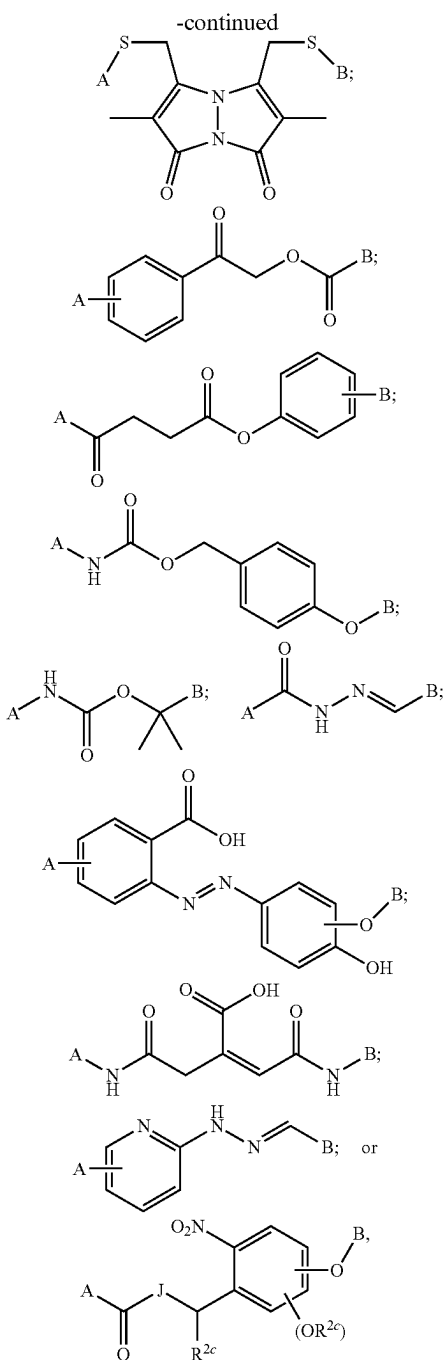
In some embodiments, the cleavable linker is:



-continued

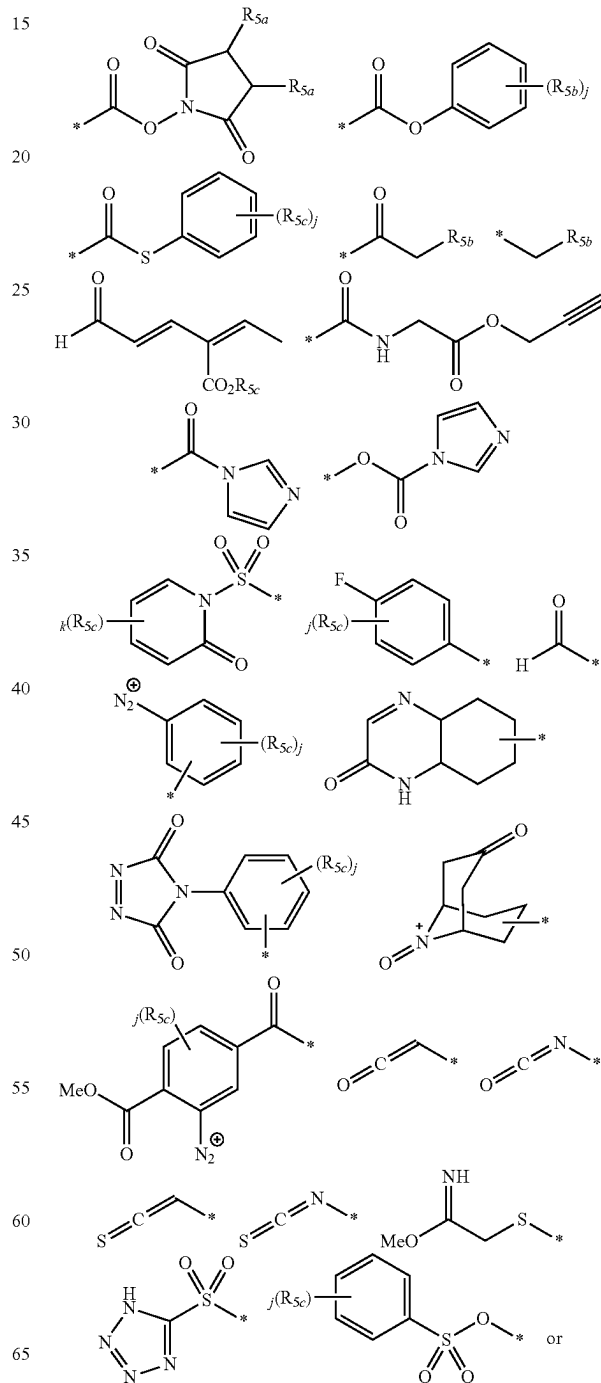


65



66

the reactive group is selective for a functional group of a specific amino acid residue, such as a lysine residue, tyrosine residue, serine residue, cysteine residue, or an unnatural amino acid residue of the Fc region incorporated to facilitate the attachment of the linker. The reactive group may be any suitable functional group, such as an activated ester for reaction with a lysine (e.g., N-hydroxysuccinimide ester or a derivate thereof, a pentafluorophenyl ester, etc.) or a sulphydryl reactive group for reaction with a cysteine (e.g., a Michael acceptor, such as an alpha-beta unsaturated carbonyl or a maleimide). In some embodiments, the reactive group is:



wherein:

one of A or B is a point of attachment the linker and the other of A or B is a point of attachment to the affinity peptide;

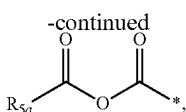
each R<sup>2a</sup> is independently H or optionally substituted alkyl;

each R<sup>2b</sup> is independently H or optionally substituted alkyl;

R<sup>20</sup> is a H or optionally substituted alkyl;

J is a methyl, a N, a S, a Si, or an O atom; and r is 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10.

The affinity peptide comprises a reactive group which is configured to enable the covalent attachment of the linker/conjugation handle to the Fc region. In some embodiments,

**67**

wherein:

- each  $\text{R}_{5a}$ ,  $\text{R}_{5b}$ , and  $\text{R}_{5c}$  is independently H, halogen, or optionally substituted alkyl;
- each j is 1, 2, 3, 4, or 5; and
- each k is 1, 2, 3, 4, or 5.

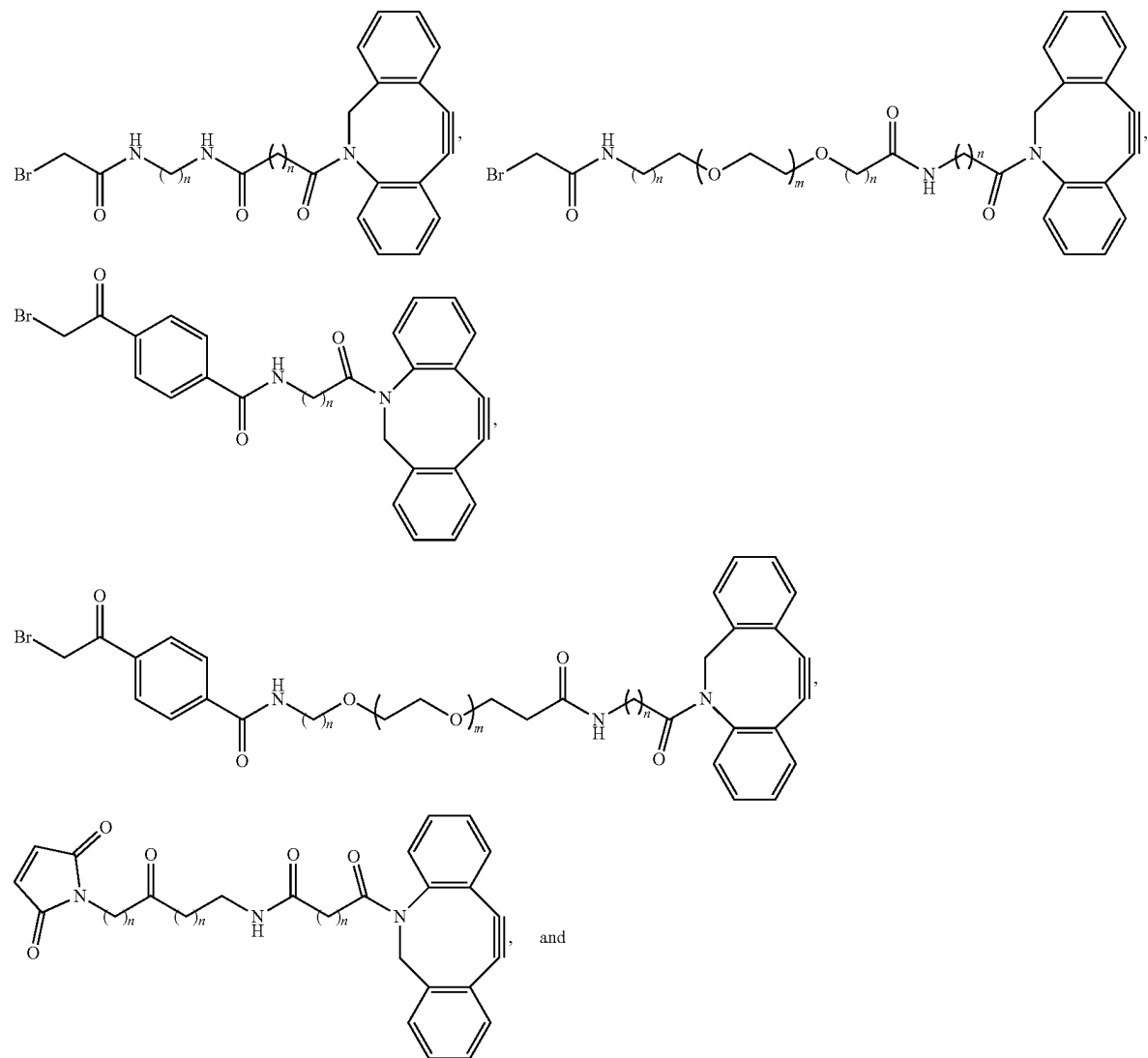
In some embodiments, the affinity peptide is used to deliver a reactive moiety to the desired amino acid residue such that the reactive moiety is exposed upon cleavage of the cleavable linker. By way of non-limiting example, the reactive group forms a covalent bond with a desired residue of the Fc region of the antibody or antigen binding fragment due to an interaction between the affinity peptide and the Fc region. Following this covalent bond formation, the cleavable linker is cleaved under appropriate conditions to reveal a reactive moiety (e.g., if the cleavable linker comprises a thioester, a free sulphydryl group is attached to the Fc region following cleavage of the cleavable linker). This new reac-

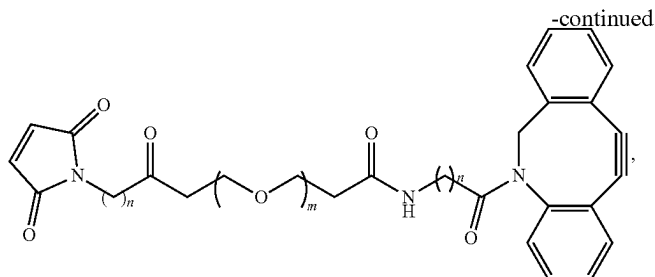
**68**

tive moiety can then be used to subsequently add an additional moiety, such as a conjugation handle, by way of reagent comprising the conjugation handle tethered to a sulphydryl reactive group (e.g., alpha-halogenated carbonyl group, alpha-beta unsaturated carbonyl group, maleimide group, etc.).

In some embodiments, an affinity peptide is used to deliver a free sulphydryl group to a lysine of the Fc region. In some embodiments, the free sulphydryl group is then reacted with a bifunctional linking reagent to attach a new conjugation handle to the Fc region. In some embodiments, the new conjugation handle is then used to form the linker to the attached cytokine. In some embodiments, the new conjugation handle is an alkyne functional group. In some embodiments, the new conjugation handle is a DBCO functional group.

Exemplary bifunctional linking reagents useful for this purpose are of a formula A-B-C, wherein A is the sulphydryl reactive conjugation handle (e.g., maleimide,  $\alpha,\beta$ -unsaturated carbonyl,  $\alpha$ -halogenated carbonyl), B is a linking group, and C is the new conjugation handle (e.g., an alkyne such as DBCO). Specific non-limiting examples of bifunctional linking reagents include





wherein each n is independently an integer from 1-6 and each m is independently an integer from 1-30, and related molecules (e.g., isomers).

Alternatively, the affinity peptide can be configured such that a conjugation handle is added to the Fc region (such as by a linker group) immediately after covalent bond formation between the reactive group and a residue of the Fc region. In such cases, the affinity peptide is cleaved and the conjugation handle is immediately ready for subsequent conjugation to the IL-18 polypeptide.

#### Alternative Methods of Modifying Fc Region

While the affinity peptide mediated modification of an Fc region of an antibody provide supra possesses many advantages over other methods which can be used to site-specifically modify the Fc region (e.g., ease of use, ability to rapidly generate many different antibody conjugates, ability to use many “off-the-shelf” commercial antibodies without the need to do time consuming protein engineering, etc.), other methods of performing the modification are also contemplated as being within the scope of the present disclosure.

In some embodiments, the present disclosure relates generally to transglutaminase-mediated site-specific antibody-drug conjugates (ADCs) comprising: 1) glutamine-containing tags, endogenous glutamines (e.g., native glutamines without engineering, such as glutamines in variable domains, CDRs, etc.), and/or endogenous glutamines made reactive by antibody engineering or an engineered transglutaminase; and 2) amine donor agents comprising amine donor units, linkers, and agent moieties. Non-limiting examples of such transglutaminase mediated site-specific modifications can be found at least in publications PCT Publication No. WO2020188061, US Patent Publication No. US2019194641, US Patent Publication No. US2021128743, U.S. Pat. No. 9,764,038, and U.S. patent Ser. No. 10/434,180, which are incorporated by reference as if set forth herein in their entirety.

In another aspect, the disclosure provides an engineered Fc-containing polypeptide conjugate comprising the formula: (Fc-containing polypeptide-T-A), wherein T is an acyl donor glutamine-containing tag engineered at a specific site, wherein A is an amine donor agent, wherein the amine donor agent is site-specifically conjugated to the acyl donor glutamine-containing tag at a carboxyl terminus, an amino terminus, or at an another site in the Fc-containing polypeptide, wherein the acyl donor glutamine-containing tag comprises an amino acid sequence XXQX, wherein X is any amino acid (e.g., X can be the same or different amino acid), and wherein the engineered Fc-containing polypeptide conjugate comprises an amino acid substitution from glutamine to asparagine at position 295 (Q295N; EU numbering scheme).

In some embodiments, the acyl donor glutamine-containing tag is not spatially adjacent to a reactive Lys (e.g., the ability to form a covalent bond as an amine donor in the presence of an acyl donor and a transglutaminase) in the polypeptide or the Fc-containing polypeptide. In some embodiments, the polypeptide or the Fc-containing polypeptide comprises an amino acid modification at the last amino acid position in the carboxyl terminus relative to a wild-type polypeptide at the same position. The amino acid modification can be an amino acid deletion, insertion, substitution, mutation, or any combination thereof.

In some embodiments, the immunocytokine composition comprises a full length antibody heavy chain and an antibody light chain, wherein the acyl donor glutamine-containing tag is located at the carboxyl terminus of a heavy chain, a light chain, or both the heavy chain and the light chain.

In some embodiments, the immunocytokine composition comprises an antibody, wherein the antibody is a monoclonal antibody, a polyclonal antibody, a human antibody, a humanized antibody, a chimeric antibody, a bispecific antibody, a minibody, a diabody, or an antibody fragment. In some embodiments, the antibody is an IgG.

In another aspect, provided herein is a method for preparing an engineered Fc-containing polypeptide conjugate comprising the formula: (Fc-containing polypeptide-T-A), wherein T is an acyl donor glutamine-containing tag engineered at a specific site, wherein A is an amine donor agent, wherein the amine donor agent is site-specifically conjugated to the acyl donor glutamine-containing tag at a carboxyl terminus, an amino terminus, or at an another site in the Fc-containing polypeptide, wherein the acyl donor glutamine-containing tag comprises an amino acid sequence XXQX, wherein X is any amino acid (e.g., X can be the same or a different amino acid), and wherein the engineered Fc-containing polypeptide conjugate comprises an amino acid substitution from glutamine to asparagine at position 295 (Q295N; EU numbering scheme), comprising the steps of: a) providing an engineered (Fc-containing polypeptide)-T molecule comprising the Fc-containing polypeptide and the acyl donor glutamine-containing tag; b) contacting the amine donor agent with the engineered (Fc-containing polypeptide)-T molecule in the presence of a transglutaminase; and c) allowing the engineered (Fc-containing polypeptide)-T to covalently link to the amine donor agent to form the engineered Fc-containing polypeptide conjugate.

In another aspect, provided herein is a method for preparing an engineered polypeptide conjugate comprising the formula: polypeptide-T-A, wherein T is an acyl donor glutamine-containing tag engineered at a specific site, wherein A is an amine donor agent, wherein the amine donor agent is site-specifically conjugated to the acyl donor glutamine-containing tag at a carboxyl terminus, an amino terminus, or at an another site in the polypeptide, and wherein the acyl

donor glutamine-containing tag comprises an amino acid sequence LLQGPX, wherein X is A or P (SEQ ID NO: 272), or GGLLQGPP (SEQ ID NO: 273), comprising the steps of: a) providing an engineered polypeptide-T molecule comprising the polypeptide and the acyl donor glutamine-containing tag; b) contacting the amine donor agent with the engineered polypeptide-T molecule in the presence of a transglutaminase; and c) allowing the engineered polypeptide-T to covalently link to the amine donor agent to form the engineered Fc-containing polypeptide conjugate.

In some embodiments, the engineered polypeptide conjugate (e.g., the engineered Fc-containing polypeptide conjugate, the engineered Fab-containing polypeptide conjugate, or the engineered antibody conjugate) as described herein has conjugation efficiency of at least about 51%. In another aspect, the invention provides a pharmaceutical composition comprising the engineered polypeptide conjugate as described herein (e.g., the engineered Fc-containing polypeptide conjugate, the engineered Fab-containing polypeptide conjugate, or the engineered antibody conjugate) and a pharmaceutically acceptable excipient.

In some embodiments, provided herein is a method for conjugating a moiety of interest (Z) to an antibody, comprising the steps of: (a) providing an antibody having (e.g., within the primary sequence of a constant region) at least one acceptor amino acid residue (e.g., a naturally occurring amino acid) that is reactive with a linking reagent (linker) in the presence of a coupling enzyme, e.g., a transamidase; and (b) reacting said antibody with a linking reagent (e.g., a linker comprising a primary amine) comprising a reactive group (R), optionally a protected reactive group or optionally an unprotected reactive group, in the presence of an enzyme capable of causing the formation of a covalent bond between the acceptor amino acid residue and the linking reagent (other than at the R moiety), under conditions sufficient to obtain an antibody comprising an acceptor amino acid residue linked (covalently) to a reactive group (R) via the linking reagent. Optionally, said acceptor residue of the antibody or antibody fragment is flanked at the +2 position by a non-aspartic acid residue. Optionally, the residue at the +2 position is a non-aspartic acid residue. In one embodiment, the residue at the +2 position is a non-aspartic acid, non-glutamine residue. In one embodiment, the residue at the +2 position is a non-aspartic acid, non-asparagine residue. In one embodiment, the residue at the +2 position is a non-negatively charged amino acid (an amino acid other than an aspartic acid or a glutamic acid). Optionally, the acceptor glutamine is in an Fc domain of an antibody heavy chain, optionally further-within the CH2 domain. Optionally, the antibody is free of heavy chain N297-linked glycosylation. Optionally, the acceptor glutamine is at position 295 and the residue at the +2 position is the residue at position 297 (EU index numbering) of an antibody heavy chain.

In one aspect, provided herein is a method for conjugating a moiety of interest (Z) to an antibody, comprising the steps of: (a) providing an antibody having at least one acceptor glutamine residue; and (b) reacting said antibody with a linker comprising a primary amine (a lysine-based linker) comprising a reactive group (R), preferably a protected reactive group, in the presence of a transglutaminase (TGase), under conditions sufficient to obtain an antibody comprising an acceptor glutamine linked (covalently) to a reactive group (R) via said linker. Optionally, said acceptor glutamine residue of the antibody or antibody fragment is flanked at the +2 position by a non-aspartic acid residue. Optionally, the residue at the +2 position is a non-aspartic

acid residue. In one embodiment, the residue at the +2 position is a non-aspartic acid, non-glutamine residue. In one embodiment, the residue at the +2 position is a non-aspartic acid, non-asparagine residue. In one embodiment, the residue at the +2 position is a non-negatively charged amino acid (an amino acid other than an aspartic acid or a glutamic acid). Optionally, the acceptor glutamine is in an Fc domain of an antibody heavy chain, optionally further-within the CH2 domain. Optionally, the antibody is free of heavy chain N297-linked glycosylation. Optionally, the acceptor glutamine is at position 295 and the residue at the +2 position is the residue at position 297 (EU index numbering) of an antibody heavy chain. The antibody comprising an acceptor residue or acceptor glutamine residue linked to a reactive group (R) via a linker comprising a primary amine (a lysine-based linker) can thereafter be reacted with a reaction partner comprising a moiety of interest (Z) to generate an antibody comprising an acceptor residue or acceptor glutamine residue linked to a moiety of interest (Z) via the linker. Thus, in one embodiment, the method further comprises a step (c): reacting (i) an antibody of step b) comprising an acceptor glutamine linked to a reactive group (R) via a linker comprising a primary amine (a lysine-based linker), optionally immobilized on a solid support, with (ii) a compound comprising a moiety of interest (Z) and a reactive group (R') capable of reacting with reactive group R, under conditions sufficient to obtain an antibody comprising an acceptor glutamine linked to a moiety of interest (Z) via a linker comprising a primary amine (a lysine-based linker). Preferably, said compound comprising a moiety of interest (Z) and a reactive group (R') capable of reacting with reactive group R is provided at a less than 80 times, 40 times, 20 times, 10 times, 5 times or 4 molar equivalents to the antibody. In one embodiment, the antibody comprises two acceptor glutamines and the compound comprising a moiety of interest (Z) and a reactive group (R') is provided at 10 or less molar equivalents to the antibody. In one embodiment, the antibody comprises two acceptor glutamines and the compound comprising a moiety of interest (Z) and a reactive group (R') is provided at 5 or less molar equivalents to the antibody. In one embodiment, the antibody comprises four acceptor glutamines and the compound comprising a moiety of interest (Z) and a reactive group (R') is provided at 20 or less molar equivalents to the antibody. In one embodiment, the antibody comprises four acceptor glutamines and the compound comprising a moiety of interest (Z) and a reactive group (R') is provided at 10 or less molar equivalents to the antibody. In one embodiment, steps (b) and/or (c) are carried out in aqueous conditions. Optionally, step (c) comprises: immobilizing a sample of an antibody comprising a functionalized acceptor glutamine residue of Formula II on a solid support to provide a sample comprising immobilized antibodies, reacting the sample comprising immobilized antibodies, optionally recovering any unreacted compound and re-introducing such recovered compound to the solid support for reaction with immobilized antibodies, and eluting the antibody conjugates to provide an antibody composition comprising a Z moiety.

In an alternative embodiment, an amino acid residue comprising a conjugation handle can be incorporated into the Fc region of the antibody (e.g., during expression of the antibody) at a desired location (e.g., any of the locations provided herein). In some embodiments, the amino acid residue comprising the conjugation handle is an unnatural amino acid.

## Conjugation Handle Chemistry

In some embodiments, the appropriately modified Fc region of the antibody or antigen binding fragment will comprise a conjugation handle which is used to conjugate the antibody or antigen binding fragment to an IL-18 polypeptide to produce an immunocytokine composition provided herein.

Any suitable reactive group capable of reacting with a complementary reactive group attached to the IL-18 polypeptide can be used as the conjugation handle. In some embodiments, the conjugation handle comprises a reagent for a Cu(I)-catalyzed or “copper-free” alkyne-azide triazole-forming reaction (e.g., strain promoted cycloadditions), the Staudinger ligation, inverse-electron-demand Diels-Alder (IEDDA) reaction, “photo-click” chemistry, tetrazine cycloadditions with trans-cyclooctenes, potassium acyl trifluoroborate (KAT) ligation or a metal-mediated process such as olefin metathesis and Suzuki-Miyaura or Sonogashira cross-coupling.

In some embodiments, the conjugation handle comprises a reagent for a “copper-free” alkyne azide triazole-forming reaction. Non-limiting examples of alkynes for said alkyne, azide triazole forming reaction include cyclooctyne reagents (e.g., (1R,8S,9s)-Bicyclo[6.1.0]non-4-yn-9-ylmethanol containing reagents, dibenzocyclooctyne-amine reagents, difluorocyclooctynes, or derivatives thereof). In some embodiments, the alkyne functional group is attached to the Fc region. In some embodiments, the azide functional group is attached to the Fc region.

In some embodiments, the conjugation handle comprises a reactive group selected from azide, alkyne, tetrazine, halide, sulphydryl, disulfide, maleimide, activated ester, alkene, aldehyde, ketone, imine, hydrazine, potassium acyl trifluoroborate, hydroxylamine (e.g., 0-substituted hydroxylamine) and hydrazide. In some embodiments, the IL-18 polypeptide comprises a reactive group complementary to the conjugation handle of the Fc region. In some embodiments, the conjugation handle and the complementary conjugation handle comprise “CLICK” chemistry reagents. Exemplary groups of click chemistry residue are shown in Hein et al., “Click Chemistry, A Powerful Tool for Pharmaceutical Sciences,” *Pharmaceutical Research*, volume 25, pages 2216-2230 (2008); Thirumurugan et al., “Click Chemistry for Drug Development and Diverse Chemical-Biology Applications,” *Chem. Rev.* 2013, 113, 7, 4905-4979; US20160107999A1; U.S. Ser. No. 10/266,502B2; and US20190204330A1, each of which is incorporated by reference in its entirety.

## Linker Structure

In some embodiments, the linker used to attach the antibody or antigen binding fragment and the IL-18 polypeptide comprises points of attachment at both moieties. The points of attachment can be any of the residues for facilitating the attachment as provided herein. The linker structure can be any suitable structure for creating the spatial attachment between the two moieties. In some embodiments, the linker provides covalent attachment of both moieties. In some embodiments, the linker is a chemical linker (e.g., not an expressed polypeptide as in a fusion protein). In some embodiments, the linker is a peptide linker. In some embodiments, the linker is a non-peptide linker (e.g., does not consist of amino acid residues).

## Chemical Linkers

In some embodiments, the linker is a chemical linker. In some embodiments, the chemical linker comprises at least one portion which is not comprised of amino acid residues. In some embodiments, the linker comprises a polymer. In

some embodiments, the linker comprises a water soluble polymer. In some embodiments, the linker comprises poly(alkylene oxide), polysaccharide, poly(vinyl pyrrolidone), poly(vinyl alcohol), polyoxazoline, poly(acryloylmorpholine), or a combination thereof. In some embodiments, the linker comprises poly(alkylene oxide). In some embodiments, the poly(alkylene oxide) is polyethylene glycol or polypropylene glycol, or a combination thereof. In some embodiments, the poly(alkylene oxide) is polyethylene glycol.

In some embodiments, the linker is a bifunctional linker. In some embodiments, the bifunctional linker comprises an amide group, an ester group, an ether group, a thioether group, or a carbonyl group. In some embodiments, the linker comprises a non-polymer linker. In some embodiments, the linker comprises a non-polymer, bifunctional linker. In some embodiments, the non-polymer, bifunctional linker comprises succinimidyl 4-(N-maleimidomethyl)cyclohexane-1-carboxylate; Maleimidocaproyl; Valine-citrulline; Allyl(4-methoxyphenyl)dimethylsilane; 6-(Allyloxycarbonylamino)-1-hexanol; 4-Aminobutyraldehyde diethyl acetal; or (E)-N-(2-Aminoethyl)-4-{2-[4-(3-azidopropoxy)phenyl]diazenyl}benzamide hydrochloride.

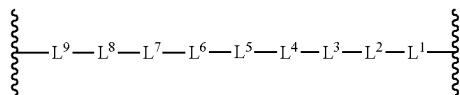
The linker can be branched or linear. In some embodiments, the linker is linear. In some embodiments, the linker is branched. In some embodiments, the linker comprises a linear portion (e.g., between the first point of attachment and the second point of attachment) of a chain of at least 10, 20, 50, 100, 500, 1000, 2000, 3000, or 5000 atoms. In some embodiments, the linker comprises a linear portion of a chain of at least 10, 20, 30, 40, or 50 atoms. In some embodiments, the linker comprises a linear portion of at least 10 atoms. In some embodiments, the linker is branched and comprises a linear portion of a chain of at least 10, 20, 50, 100, 500, 1000, 2000, 3000, or 5000 atoms. In some embodiments, the linker comprises a linear portion of at least 1 to 1000 atoms, 1 to 900 atoms, 1 to 800 atoms, 1 to 500 atoms, 1 to 400 atoms, 1 to 300 atoms, 1 to 200 atoms, 1 to 100 atoms, 1 to 50 atoms, 10 to 1000 atoms, 10 to 900 atoms, 10 to 800 atoms, 10 to 500 atoms, 10 to 400 atoms, 10 to 300 atoms, 10 to 200 atoms, 10 to 100 atoms, 10 to 50 atoms, 25 to 1000 atoms, 25 to 900 atoms, 25 to 800 atoms, 25 to 500 atoms, 25 to 400 atoms, 25 to 300 atoms, 25 to 200 atoms, 25 to 100 atoms, 25 to 50 atoms, 50 to 1000 atoms, 50 to 900 atoms, 50 to 800 atoms, 50 to 500 atoms, 50 to 400 atoms, 50 to 300 atoms, 50 to 200 atoms, or 50 to 100 atoms. In some embodiments, the linker has a linear length of from about 10 angstroms to about 200 angstroms. In some embodiments, the linker has a linear length of from about 10 to 500, 10 to 200, 10 to 150, 10 to 125, 10 to 100, 10 to 75, 10 to 50, 25 to 200, 25 to 150, 25 to 125, 25 to 100, 25 to 75, 25 to 50, 50 to 200, 50 to 150, 50 to 100, or 50 to 75 angstroms.

In some embodiments, the linker has a molecular weight of about 200 Daltons to about 2000 Daltons. In some embodiments, the linker has a molecular weight of about 200 Daltons to about 5000 Daltons. In some embodiments, the linker has a molecular weight of 200 Daltons to 100,000 Daltons. In some embodiments, the linker has a molecular weight of at least about 500 Daltons, at least about 1,000 Daltons, at least about 5,000 Daltons, at least about 10,000 Daltons, at least about 15,000 Daltons, at least about 20,000 Daltons, at least about 25,000 Daltons, or at least about 30,000 Daltons. In some embodiments, the linker has a molecular weight of at most about 100,000 Daltons, at most about 50,000 Daltons, at most about 40,000 Daltons, at most about 30,000 Daltons, at most about 25,000 Daltons, at most

about 20,000 Daltons at most about 15,000 Daltons, at most about 10,000 Daltons, or at most about 5,000 Daltons.

In some embodiments, the linker comprises a reaction product of one or more pairs of conjugation handles and a complementary conjugation handle thereof. In some embodiments, the reaction product comprises a triazole, a hydrazone, pyridazine, a sulfide, a disulfide, an amide, an ester, an ether, an oxime, an alkene, or any combination thereof. In some embodiments, the reaction product comprises a triazole. The reaction product can be separated from the first point of attachment and the second point of attachment by any portion of the linker. In some embodiments, the reaction product is substantially in the center of the linker. In some embodiments, the reaction product is substantially closer to one point of attachment than the other is.

In some embodiments, the linker comprises a structure of Formula (X)



wherein each of L<sup>1</sup>, L<sup>2</sup>, L<sup>3</sup>, L<sup>4</sup>, L<sup>5</sup>, L<sup>6</sup>, L<sup>7</sup>, L<sup>8</sup>, and L<sup>9</sup> is independently —O—, —NR<sup>L</sup>—, —(C<sub>1</sub>-C<sub>6</sub> alkylene) NR<sup>L</sup>—, —NR<sup>L</sup>(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—, —N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —OP(=O)(OR<sup>L</sup>)O—, —S—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)S—, —S(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —S(=O)—, —S(=O)<sub>2</sub>—, —C(=O)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)C(=O)—, —C(=O)(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —C(=O)O—, —OC(=O)—, —OC(=O)O—, —C(=O)NR<sup>L</sup>—, —C(=O)NR<sup>L</sup>(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)C(=O)NR<sup>L</sup>—, —NR<sup>L</sup>C(=O)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)NR<sup>L</sup>C(=O)—, —NR<sup>L</sup>C(=O)(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —OC(=O)NR<sup>L</sup>—, —NR<sup>L</sup>C(=O)O—, —NR<sup>L</sup>—, —NR<sup>L</sup>C(S)NR<sup>L</sup>—, —CR<sup>L</sup>=N—, —N=CR<sup>L</sup>—, —NR<sup>L</sup>S(=O)<sub>2</sub>—, —S(=O)<sub>2</sub>NR<sup>L</sup>—, —C(=O)NR<sup>L</sup>S(=O)<sub>2</sub>—, —S(=O)<sub>2</sub>NR<sup>L</sup>C(=O)—, substituted or unsubstituted C<sub>1</sub>-C<sub>6</sub> alkylene, substituted or unsubstituted C<sub>1</sub>-C<sub>6</sub> heteroalkylene, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkenylene, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkynylene, substituted or unsubstituted C<sub>6</sub>-C<sub>20</sub> arylene, substituted or unsubstituted C<sub>2</sub>-C<sub>20</sub> heteroarylene, —(CH<sub>2</sub>-CH<sub>2</sub>-O)<sub>qa</sub>—, —(O-CH<sub>2</sub>-CH<sub>2</sub>)<sub>qb</sub>—, —(CH<sub>2</sub>-CH(CH<sub>3</sub>)-O)<sub>qc</sub>—, —(O-CH(CH<sub>3</sub>)-CH<sub>2</sub>)<sub>qd</sub>—, a reaction product of a conjugation handle and a complementary conjugation handle, or absent;

each R<sup>L</sup> is independently hydrogen, substituted or unsubstituted C<sub>1</sub>-C<sub>4</sub> alkyl, substituted or unsubstituted C<sub>1</sub>-C<sub>4</sub> heteroalkyl, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkenyl, substituted or unsubstituted C<sub>2</sub>-C<sub>5</sub> alkynyl, substituted or unsubstituted C<sub>3</sub>-C<sub>8</sub> cycloalkyl, substituted or unsubstituted C<sub>2</sub>-C<sub>7</sub> heterocycloalkyl, substituted or unsubstituted aryl, or substituted or unsubstituted heteroaryl; and

each of qa, qb, qc and qd is independently an integer from 1-100,

wherein each



is a point of attachment to the antibody or antigen binding fragment or the IL-18 polypeptide.

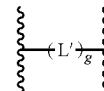
In some embodiments, the linker consists of a plurality of structures of Formula (X) to form the linkage between the antibody or antigen binding fragment and the IL-18 polypeptide (e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, or more structures of Formula (X) appended from end to end, where



10

only the terminal denote points of attachment to the antibody or antigen binding fragment or the IL-18 polypeptide).

In some embodiments, the polymer comprises a linker comprising a structure of Formula (X')



wherein each L' is independently —O—, —NR<sup>L</sup>—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)NR<sup>L</sup>—, —NR<sup>L</sup>(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—, —N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —OP(=O)(OR<sup>L</sup>)O—, —S—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)S—, —S(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —S(=O)—, —S(=O)<sub>2</sub>—, —C(=O)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)C(=O)—, —C(=O)(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —C(=O)O—, —OC(=O)—, —OC(=O)O—, —C(=O)NR<sup>L</sup>—, —C(=O)NR<sup>L</sup>(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)C(=O)NR<sup>L</sup>—, —NR<sup>L</sup>C(=O)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)NR<sup>L</sup>C(=O)—, —NR<sup>L</sup>—, —NR<sup>L</sup>C(S)NR<sup>L</sup>—, —CR<sup>L</sup>=N—, —N=CR<sup>L</sup>—, —NR<sup>L</sup>S(=O)<sub>2</sub>—, —S(=O)<sub>2</sub>NR<sup>L</sup>—, —C(=O)NR<sup>L</sup>S(=O)<sub>2</sub>—, —S(=O)<sub>2</sub>NR<sup>L</sup>C(=O)—, substituted or unsubstituted C<sub>1</sub>-C<sub>6</sub> alkylene, substituted or unsubstituted C<sub>1</sub>-C<sub>6</sub> heteroalkylene, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkenylene, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkynylene, substituted or unsubstituted C<sub>6</sub>-C<sub>20</sub> arylene, substituted or unsubstituted C<sub>2</sub>-C<sub>20</sub> heteroarylene, —(CH<sub>2</sub>-CH<sub>2</sub>-O)<sub>qa</sub>—, —(O-CH<sub>2</sub>-CH<sub>2</sub>)<sub>qb</sub>—, —(CH<sub>2</sub>-CH(CH<sub>3</sub>)-O)<sub>qc</sub>—, —(O-CH(CH<sub>3</sub>)-CH<sub>2</sub>)<sub>qd</sub>—, a reaction product of a conjugation handle and a complementary conjugation handle, or absent; (C<sub>1</sub>-C<sub>6</sub> alkylene);

each R<sup>L</sup> is independently hydrogen, substituted or unsubstituted C<sub>1</sub>-C<sub>4</sub> alkyl, substituted or unsubstituted C<sub>1</sub>-C<sub>4</sub> heteroalkyl, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkenyl, substituted or unsubstituted C<sub>2</sub>-C<sub>5</sub> alkynyl, substituted or unsubstituted C<sub>3</sub>-C<sub>8</sub> cycloalkyl, substituted or unsubstituted C<sub>2</sub>-C<sub>7</sub> heterocycloalkyl, substituted or unsubstituted aryl, or substituted or unsubstituted heteroaryl; and

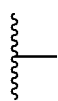
each of qa, qb, qc and qd is independently an integer from 1-100,

g is an integer from 1-100,

60

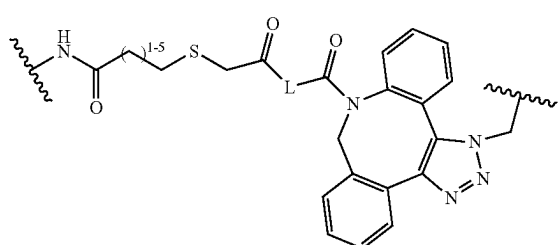
65

wherein each

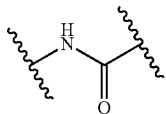


is a point of attachment to the modified IL-18 polypeptide or the antibody or antigen binding fragment.

In some embodiments, the linker of Formula (X) or Formula (X') comprises the structure:

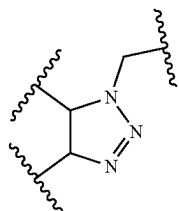


wherein



is the first point of attachment to a lysine residue of the antibody or antigen binding fragment;

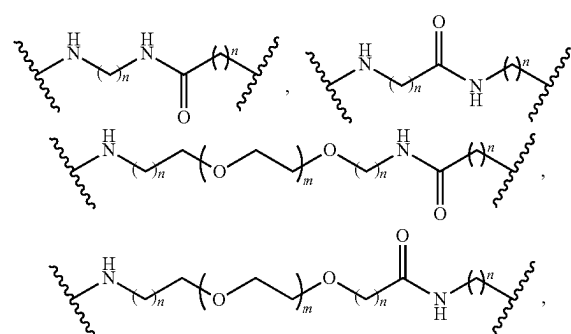
L is a linking group; and



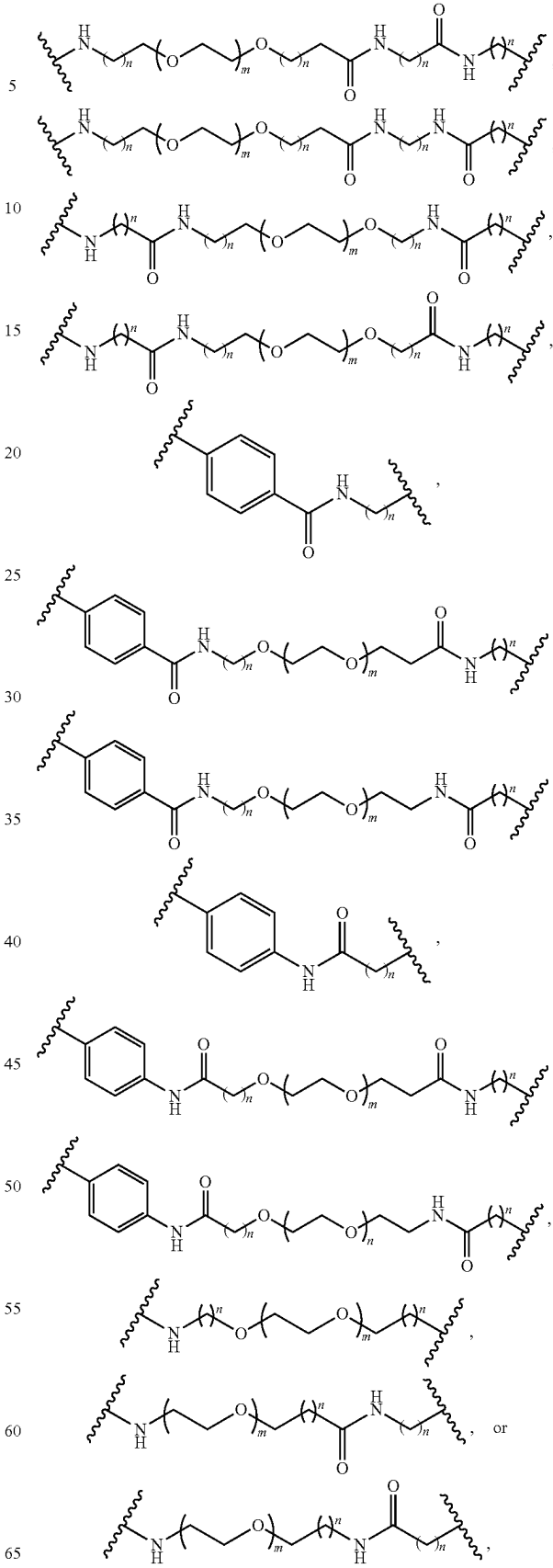
is a point of attachment to a linking group which connects to the first point of attachment,

or a regioisomer thereof.

In some embodiments, L has a structure 0 n H

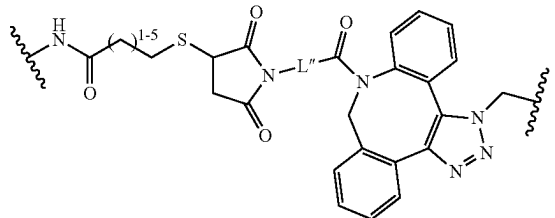


-continued

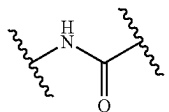


79

In some embodiments, the linker of Formula (X) or of Formula (X<sup>a</sup>) or of Formula (X') comprises the structure:

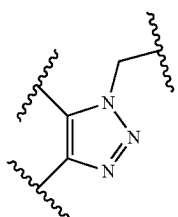


wherein



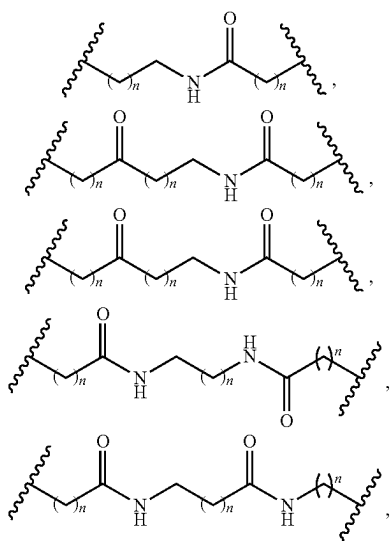
is the first point of attachment to a lysine residue of the polypeptide which selectively binds to PD-1;

L is a linking group; and



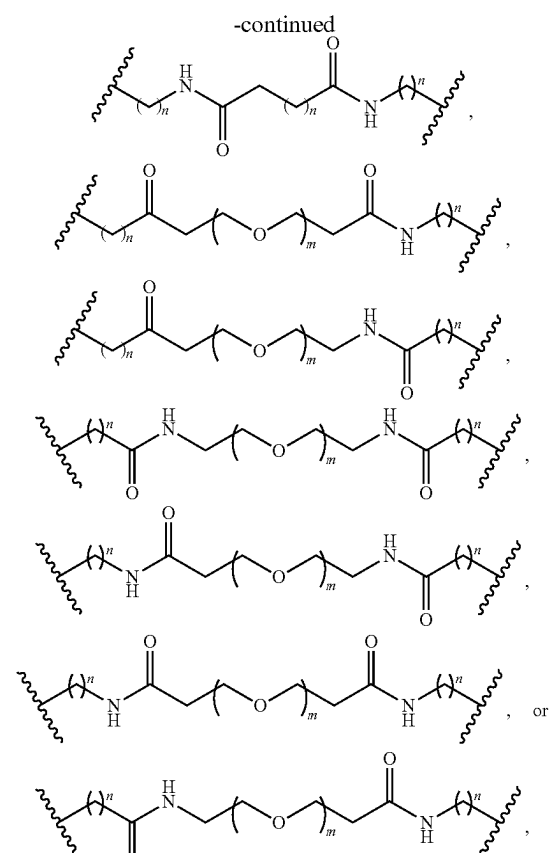
is a point of attachment to a linking group which connects to the first point of attachment, or a regioisomer thereof.

In some embodiments, L<sup>a</sup> has a structure



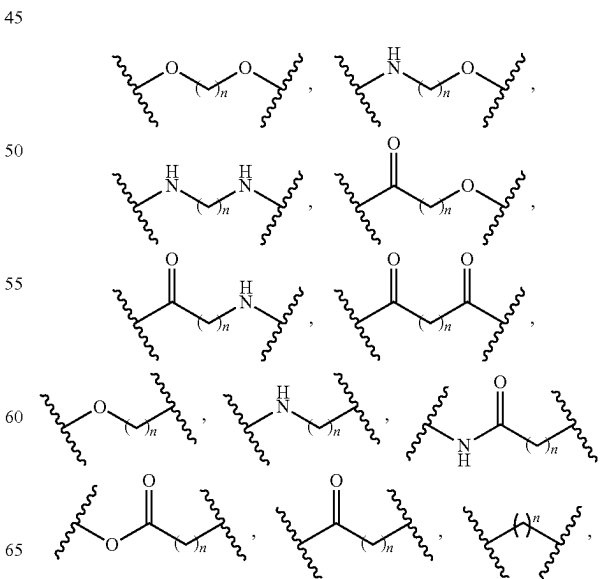
80

-continued

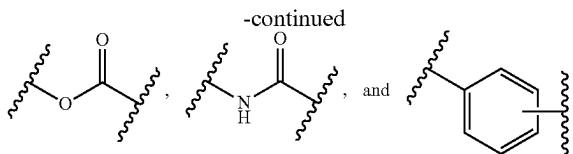


35 wherein each n is independently an integer from 1-6 and each m is independently an integer from 1-30. In some embodiments, each m is independently 2 or 3. In some embodiments, each m is an integer from 1-24, from 1-18, from 1-12, or from 1-6.

In some embodiments, L or L<sup>a</sup> comprises 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 or more subunits each independently selected from

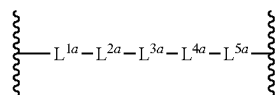


81



wherein each n is independently an integer from 1-30. In some embodiments, each n is independently an integer from 1-6. In some embodiments, L or L" comprise 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 of the subunits.

In some embodiments, L or L" is a structure of Formula (X")



—O—, —NR<sup>La</sup>—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)NR<sup>La</sup>—, —NR<sup>La</sup>(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)N(R<sup>La</sup>)<sub>2</sub><sup>+</sup>—(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —N(R<sup>L</sup>)<sub>2</sub><sup>+</sup>—, —OP(=O)(OR<sup>La</sup>)O—, —S—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)S—, —S(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —S(=O)—, —S(=O)<sub>2</sub>—, —C(=O)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)C(=O)—, —C(=O)(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —C(=O)O—, —OC(=O)—, —OC(=O)O—, —C(=O)NR<sup>La</sup>—, —C(=O)NR<sup>La</sup>—(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)C(=O)NR<sup>La</sup>—, —NR<sup>La</sup>C(=O)—, —(C<sub>1</sub>-C<sub>6</sub> alkylene)NR<sup>La</sup>C(=O)—, —NR<sup>La</sup>C(=O)(C<sub>1</sub>-C<sub>6</sub> alkylene)—, —OC(=O)NR<sup>La</sup>—, —NR<sup>La</sup>C(=O)O—, —NR<sup>La</sup>C(=O)NR<sup>La</sup>—, —NR<sup>La</sup>C(=S)NR<sup>La</sup>—, —CR<sup>La</sup>=N—, —N=CR<sup>La</sup>—, —NR<sup>La</sup>S(=O)<sub>2</sub>—, —S(=O)<sub>2</sub>NR<sup>La</sup>—, —C(=O)NR<sup>La</sup>S(=O)<sub>2</sub>—, —S(=O)<sub>2</sub>NR<sup>La</sup>C(=O)—, substituted or unsubstituted C<sub>1</sub>-C<sub>6</sub> alkylene, substituted or unsubstituted C<sub>1</sub>-C<sub>6</sub> heteroalkylene, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkenylene, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkynylene, substituted or unsubstituted C<sub>6</sub>-C<sub>20</sub> arylene, substituted or unsubstituted C<sub>2</sub>-C<sub>20</sub> heteroarylene, —(CH<sub>2</sub>—CH<sub>2</sub>—O)<sub>qe</sub>—, —(O—CH<sub>2</sub>—CH<sub>2</sub>)<sub>qf</sub>—, —(CH<sub>2</sub>—CH(CH<sub>3</sub>)—O)<sub>qg</sub>—, —(O—CH(CH<sub>3</sub>)—CH<sub>2</sub>)<sub>qh</sub>—, a reaction product of a conjugation handle and a complementary conjugation handle, or absent; (C<sub>1</sub>-C<sub>6</sub> alkylene)

each R<sup>La</sup> is independently hydrogen, substituted or unsubstituted C<sub>1</sub>-C<sub>4</sub> alkyl, substituted or unsubstituted C<sub>1</sub>-C<sub>4</sub> heteroalkyl, substituted or unsubstituted C<sub>2</sub>-C<sub>6</sub> alkenyl, substituted or unsubstituted C<sub>2</sub>-C<sub>5</sub> alkynyl, substituted or unsubstituted C<sub>3</sub>-C<sub>8</sub> cycloalkyl, substituted or unsubstituted C<sub>2</sub>-C<sub>7</sub> heterocycloalkyl, substituted or unsubstituted aryl, or substituted or unsubstituted heteroaryl; and

each of qe, qf, qg and qh is independently an integer from 1-100.

In some embodiments, L or L" comprises a linear chain of 2 to 10, 2 to 15, 2 to 20, 2 to 25, or 2 to 30 atoms. In some embodiments, the linear chain comprises one or more alkyl groups (e.g., lower alkyl (C<sub>1</sub>-C<sub>4</sub>)), one or more aromatic groups (e.g., phenyl), one or more amide groups, one or more ether groups, one or more ester groups, or any combination thereof.

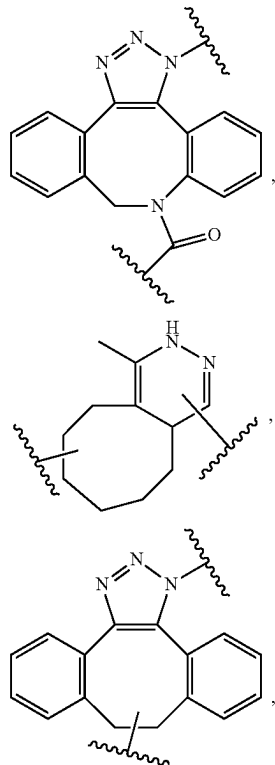
In some embodiments, the linking group which connects to the first point of attachment (e.g., the point of attachment to the IL-18 polypeptide) comprises poly(ethylene glycol).

82

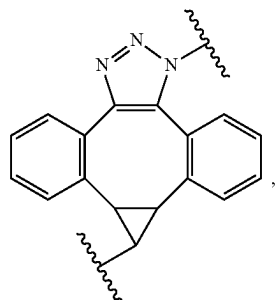
In some embodiments, the linking group comprises about 2 to about 30 poly(ethylene glycol) units. In some embodiments, the linking group which connects to the first point of attachment (e.g., the point of attachment to the IL-18 polypeptide) is a functionality attached to a cytokine provided herein which comprises an azide (e.g., the triazole is the reaction product of the azide).

In some embodiments, each reaction product of a conjugation handle and a complementary conjugation handle independently comprises a triazole, a hydrazone, pyridazine, a sulfide, a disulfide, an amide, an ester, an ether, an oxime, or an alkene. In some embodiments, each reaction product of a conjugation handle and a complementary conjugation handle comprises a triazole. In some embodiments, each reaction product of a conjugation handle and a complementary conjugation handle comprise a structure of

20



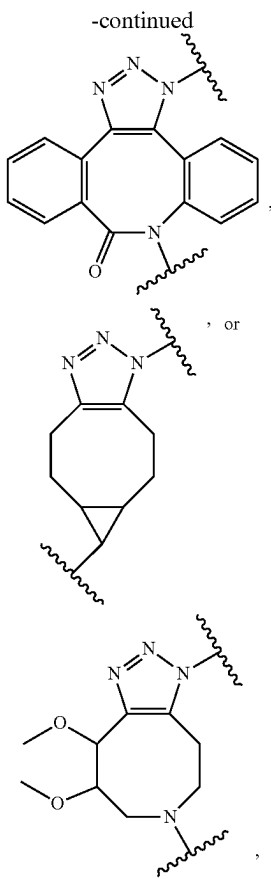
55



60

65

83



or a regioisomer or derivative thereof.

#### Peptide Linkers

In some embodiments, the antibody or antigen binding fragment is linked to the IL-18 polypeptide through a peptide linker. In some embodiments, the antibody or antigen binding fragment is linked to the IL-18 polypeptide as a fusion protein. In such instances, the linker comprises one or more peptide bonds between the antibody or antigen binding fragment and the IL-18 polypeptide. In some embodiments, the linker between the fusion protein of the antibody or antigen binding fragment and the IL-18 polypeptide is a bond. In some embodiments, the linker between the fusion protein of the antibody or antigen binding fragment and the IL-18 polypeptide is a linking peptide. Non-limiting examples of linking peptides include, but are not limited to (GS)<sub>n</sub> (SEQ ID NO: 224), (GGS)<sub>n</sub> (SEQ ID NO: 225), (GGGS)<sub>n</sub> (SEQ ID NO: 226), (GGSG)<sub>n</sub> (SEQ ID NO: 227), or (GGSGG)<sub>n</sub> (SEQ ID NO: 228), (GGGGS)<sub>n</sub> (SEQ ID NO: 229), wherein n is 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10. For example, a linking peptide can be (GGGGS)<sub>3</sub> (SEQ ID NO: 230) or (GGGGS)<sub>4</sub> (SEQ ID NO: 231). In some embodiments, the IL-18 polypeptide is fused to the C-terminal end of the antibody or antigen binding fragment (optionally through a linking peptide). In some embodiments, the IL-18 polypeptide is fused to the N-terminal end of the antibody or antigen binding fragment (optionally through a linking peptide).

#### Cleavable Linkers

In some embodiments, the linker (e.g., a chemical or peptide linker as provided herein) is a cleavable linker. In some embodiments, the cleavable linker is cleaved at, near, or in a tumor microenvironment. In some embodiments, the

84

tumor is mechanically or physically cleaved at, near, or in the tumor microenvironment. In some embodiments, the tumor is chemically cleaved at, near, or in a tumor microenvironment. In some embodiments, the cleavable linker is a reduction sensitive linker. In some embodiments, the cleavable linker is an oxidation sensitive linker. In some embodiments, the cleavable linker is cleaved as a result of pH at, near, or in the tumor microenvironment. In some embodiments, the cleavable linker is cleaved by a tumor metabolite at, near, or in the tumor microenvironment. In some embodiments, the cleavable linker is cleaved by a protease at, near, or in the tumor microenvironment.

#### IL-18 Polypeptides

The present disclosure describes antibodies or antigen binding fragments linked to interleukin-18 (IL-18) polypeptides as immunocytokine compositions and their use as therapeutic agents. IL-18 is a pro-inflammatory cytokine that elicits biological activities that initiate or promote host defense and inflammation following infection or injury. IL-18 has been implicated in autoimmune diseases, myocardial function, emphysema, metabolic syndromes, psoriasis, inflammatory bowel disease, hemophagocytic syndromes, macrophage activation syndrome, sepsis, and acute kidney injury. In some models of disease, IL-18 plays a protective role.

IL-18 also plays a major role in the production of IFN $\gamma$  from T-cells and natural killer cells. IFN $\gamma$  is a T helper type 1 cytokine mainly produced by T cells, NK cells, and macrophages and is critical for innate and adaptive immunity against viral, some bacterial, and protozoal infections. IFN $\gamma$  is also an important activator of macrophages and inducer of Class II major histocompatibility complex (MHC) molecule expression.

IL-18 forms a signaling complex by binding to the IL-18 alpha chain (IL-18R $\alpha$ ), which is the ligand binding chain for mature IL-18. However, the binding affinity of IL-18 to IL-18R $\alpha$  is low. In cells that express the co-receptor, IL-18 receptor beta chain (IL-18R $\beta$ ), a high affinity heterodimer complex is formed, which then activates cell signaling.

The activity of IL-18 is balanced by the presence of a high affinity, naturally occurring IL-18 binding protein (IL-18BP). IL-18BP binds IL-18 and neutralizes the biological activity of IL-18. Cell surface IL-18R $\alpha$  competes with IL-18BP for IL-18 binding. Increased disease severity can be associated with an imbalance of IL-18 to IL-18BP such that levels of free IL-18 are elevated in the circulation. FIG. 4 illustrates the mechanism of action of IL-18, IFN $\gamma$  production, IL-18BP production, and inhibition of IL-18 activity by IL-18BP. IL-18 induces IFN $\gamma$  production, which in turn induces IL-18BP production. IL-18BP then competes with IL-18R $\alpha$  to inhibit IL-18 activity.

In some embodiments, the IL-18 polypeptides of the immunocytokines provided herein display reduced binding to IL-18BP and retain binding to the IL-18 receptor. The IL-18 polypeptides with this property provided herein are able to retain IL-18 receptor signaling activity (including inducing production of IFN $\gamma$ ) even in the presence of IL-18BP. This allows the immunocytokines provided herein to retain IL-18 signaling activity well beyond a short period of time after administration, or upon repeat administrations. In some embodiments, the modified IL-18 polypeptides with this property comprise a modification (e.g., substitution, polymer attachment, or deletion) at one or more amino acid residues which convey this property to the IL-18 polypeptide. Examples of IL-18 polypeptides with this property are provided herein, as well as those otherwise known, such as those described in Patent Cooperation Treaty Publication

85

No. WO2019051015A1, which is hereby incorporated by reference as if set forth herein in its entirety. In addition to IL-18 polypeptides provided herein, these otherwise known IL-18 polypeptides or their analogs may similarly be modified with points of attachment to the linker as provided herein.

#### Points of Attachment of Chemical Linkers to IL-18 Polypeptides

The immunocytokines provided herein comprise linkers which have a point of attachment to the IL-18 polypeptide. In some embodiments, the linker is a chemical linker. As discussed supra, the linker has another point of attachment to the antibody or antigen binding fragment at any residue as provided herein. The point of attachment to the IL-18 polypeptide is to a residue as provided herein.

In some embodiments, the linker is attached to an amino acid residue of the IL-18 polypeptide. In some embodiments, the linker is attached to any amino acid residue of the IL-18 polypeptide (e.g., at a position corresponding to any one of positions 1-157 of SEQ ID NO: 1). In some embodiments, the linker is attached at a non-terminal residue of the IL-18 polypeptide (e.g., a residue at position corresponding to any one of positions 2-156 of SEQ ID NO: 1). In some embodiments, the linker is attached at a non-terminal residue of the IL-18 polypeptide, wherein the IL-18 polypeptide has been extended or truncated by one or more amino acids relative to SEQ ID NO: 1.

In some embodiments, the linker is attached to the IL-18 polypeptide at a residue in a region comprising residues 2-156, wherein residue position numbering is based on SEQ ID NO: 1 as a reference sequence. In some embodiments, the linker is attached to the IL-18 polypeptide at a residue in a region comprising residues 30-150. In some embodiments, the linker is attached to the IL-18 polypeptide at a residue in a region comprising residues 33-43, residues 60-100, residues 65-75, residues 80-90, residues 85-100, residues 90-110, residues 115-130, residues 120-130, or residues 140-150. In some embodiments, the linker is attached to the IL-18 polypeptide at a residue selected from residue 38, 68, 69, 70, 76, 78, 85, 86, 95, 98, 121, 127, and 144. In some embodiments, the linker is attached to the IL-18 polypeptide at a residue selected from 68, 69, 70, 85, 86, and 98. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 68, 69, or 70. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 85, 86, 95, or 98. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 68. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 69. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 70. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 85. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 86. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 95. In some embodiments, the linker is attached to the IL-18 polypeptide at residue 98.

In some embodiments, the linker is attached to the IL-18 polypeptide at a residue which is known in the art to be compatible with attachment of a polymer to the IL-18 polypeptide without having a profound impact on the bioactivity of the IL-18 polypeptide. Examples of these residues include residues 38, 76, 78, 121, 127, and 144, as described in PCT Pub. No. WO2004091517A2, which is hereby incorporated by reference as if set forth in its entirety.

In some embodiments, the residue to which the linker is attached is a natural amino acid residue. In some embodiments, the residue to which the linker is covalently attached

86

is selected from cysteine, aspartate, asparagine, glutamate, glutamine, serine, threonine, lysine, and tyrosine. In some embodiments, the residue to which the linker is covalently attached is selected from asparagine, aspartic acid, cysteine, glutamic acid, glutamine, lysine, and tyrosine. In some embodiments, the linker is covalently attached to a cysteine. In some embodiments, the linker is covalently attached to a lysine. In some embodiments, the linker is covalently attached to a glutamine. In some embodiments, the linker is covalently attached to an asparagine. In some embodiments, the residue to which the linker is attached is a tyrosine. In some embodiments, the residue to which the linker is attached is the natural amino acid in that position in SEQ ID NO: 1.

In some embodiments, the linker is attached to a different natural amino acid which is substituted at the relevant position. The substitution can be for a naturally occurring amino acid which is more amenable to attachment of additional functional groups (e.g., aspartic acid, cysteine, glutamic acid, lysine, serine, threonine, or tyrosine), a derivative of modified version of any naturally occurring amino acid, or any unnatural amino acid (e.g., an amino acid containing a desired conjugation handle, such as a CLICK chemistry reagent such as an azide, alkyne, etc.). In some embodiments, the linker is covalently attached to site-specifically to a natural amino acid.

In some embodiments, the linker is attached at an unnatural amino acid residue. In some embodiments, the unnatural amino acid residue comprises a conjugation handle. In some embodiments, the conjugation handle facilitates the addition of the linker to the modified IL-18 polypeptide. The conjugation handle can be any of the conjugation handles provided herein. In some embodiments, the linker is covalently attached site-specifically to the unnatural amino acid. Non-limiting examples of amino acid residues comprising conjugation handles can be found, for example, in PCT Pub. Nos. WO2015054658A1, WO2014036492A1, WO2021133839A1 WO2006069246A2, and WO2007079130A2, each of which is incorporated by reference as if set forth in its entirety.

In some embodiments, the linker is covalently attached at residue 68. In some embodiments, the linker is covalently attached at residue C68, C68E, C68D, C68Q, C68K, C68N, or C68Y. In some embodiments, the linker is covalently attached at residue C68. In some embodiments, the linker is covalently attached to an unnatural amino acid at residue 68.

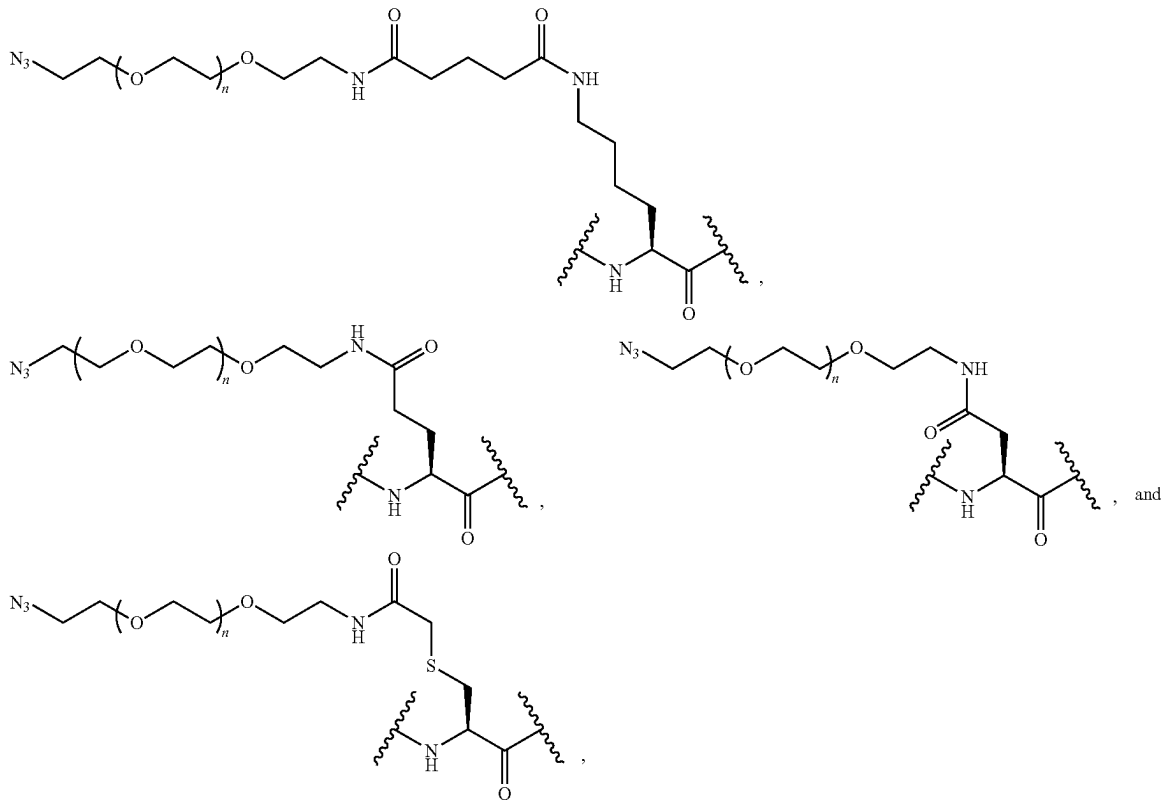
In some embodiments, the linker is covalently attached at residue 69. In some embodiments, the linker is covalently attached at residue E69, E69C, E69D, E69Q, E69K, E69N, or E69Y. In some embodiments, the linker is covalently attached at residue E69. In some embodiments, the linker is covalently attached residue E69C. In some embodiments, the linker is covalently attached to an unnatural amino acid at residue 69.

In some embodiments, the linker is covalently attached at residue 70. In some embodiments, the linker is covalently attached at residue K70, K70C, K70D, K70Q, K70E, K70N, or K70Y. In some embodiments, the linker is covalently attached at residue K70. In some embodiments, the linker is covalently attached residue K70C. In some embodiments, the linker is covalently attached to an unnatural amino acid at residue 70.

In some embodiments, the linker is covalently attached at residue 85. In some embodiments, the linker is covalently attached at residue E85, E85C, E85D, E85Q, E85K, E85N, or E85Y. In some embodiments, the linker is covalently attached at residue E85. In some embodiments, the linker is covalently attached residue E85C. In some embodiments, the linker is covalently attached to an unnatural amino acid at residue 85.

In some embodiments, the linker is covalently attached at residue 86. In some embodiments, the linker is covalently

ments, the modified natural amino acid comprises a conjugation handle. In some embodiments, the linker is covalently attached through a modified amino acid a. In some embodiments, the modified amino acid a is an amino-acid-PEG-azide group. In some embodiments, the modified amino acid a is a glutamate, aspartate, lysine, cysteine, or tyrosine modified to incorporate an azide group linked to the amino acid through a PEG spacer. In some embodiments, the modified amino acid a has a structure selected from:



attached at residue M86C, M86D, M86Q, M86K, M86N, M86E, or M86Y. In some embodiments, the linker is covalently attached M86C. In some embodiments, the linker is covalently attached to an unnatural amino acid at residue 86.

In some embodiments, the polymer is covalently attached at residue 95. In some embodiments, the polymer is covalently attached at residue T95, T95C, T95D, T95Q, T95K, T95N, T95E, or T95Y. In some embodiments, the polymer is covalently attached at residue T95C, T95D, T95Q, T95K, T95N, T95E, or T95Y. In some embodiments, the polymer is covalently attached at residue T95C. In some embodiments, the polymer is covalently attached to an unnatural amino acid at residue 95.

In some embodiments, the linker is covalently attached at residue 98. In some embodiments, the linker is covalently attached at residue D98, D98C, D98Q, D98K, D98N, D98E, or D98Y. In some embodiments, the linker is covalently attached at residue D98C. In some embodiments, the linker is covalently attached to an unnatural amino acid at residue 98.

In some embodiments, the linker is covalently attached through a modified natural amino acid. In some embodi-

5 where each n is independently an integer from 1-30. In some embodiments, n is an integer from 1-20, 1-10, 2-30, 2-20, 2-10, 5-30, 5-20, or 5-10. In some embodiments, n is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, or 30. In some embodiments, n is 10. In some embodiments, n is 8. In some embodiments, n is 6. In some embodiments, n is 12. The modified amino acid a can be incorporated at any point of attachment of the IL-18 polypeptide as provided herein. In some embodiments, the modified amino acid a is located at a position on the modified IL-18 polypeptide selected from residue 68, residue 69, residue 70, residue 85, residue 86, residue 95, or residue 98.

50 Where IL-18 polypeptides contain unnatural amino acids or modified natural amino acids (e.g., those provided herein for purposes of conjugation), these amino acids may be incorporated into the IL-18 polypeptides using many techniques known in the art for introduction such modifications. For example, recombinant proteins with unnatural amino acids can be made using methods as described in Patent Cooperation Treaty Publication Nos. WO2016115168, 55 WO2002085923, WO2005019415, and WO2005003294.

Alternatively or in combination, unnatural or modified natural amino acids can be incorporated into chemically synthesized proteins during synthesis.

#### Modifications to IL-18 Polypeptides

In some embodiments, the IL-18 polypeptide of the immunocytokine comprises one or more modifications to that of SEQ ID NO: 1. The modifications provided herein are in addition to any modification at the point of attachment as discussed supra. In some embodiments, the residue position numbering of the modified IL-18 polypeptide is based on SEQ ID NO: 1 as a reference sequence.

Modifications to the IL-18 polypeptide described herein encompass mutations, addition of various functionalities, deletion of amino acids, addition of amino acids, or any other alteration of the wild-type version of the protein or protein fragment. Functionalities which may be added to polypeptides include polymers, linkers, alkyl groups, detectable molecules such as chromophores or fluorophores, reactive functional groups, or any combination thereof. In some embodiments, functionalities are added to individual amino acids of the polypeptides. In some embodiments, functionalities are added site-specifically to the polypeptides.

In some embodiments, the IL-18 polypeptide of the immunocytokine comprise one or more modifications in addition to a modification needed to attach the linker to the relevant residue of the IL-18 polypeptide (e.g., an amino acid substitution at a residue to which the linker is not attached). In some embodiments, the modification is in the range of amino acid residues 1-127, based on the sequence of human IL-18<sup>37-193</sup> (SEQ ID NO: 1). SEQ ID NO: 1 reflects the bioactive form of IL-18. Endogenously, IL-18 is initially expressed with an additional 36 amino acid segment at the N-terminus which is cleaved by caspases to mediate biologic activity.

In some embodiments, the IL-18 polypeptide of the immunocytokine described herein contain 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, or more modified amino acid residues.

In some embodiments, the IL-18 polypeptide of the immunocytokine comprises an amino acid sequence having at least 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% sequence identity to the sequence set forth in SEQ ID NO: 1.

In some embodiments, the IL-18 polypeptide of the immunocytokine provided herein comprises an amino acid sequence of any one of SEQ ID NOs: 2-203 provided herein. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence at least 85% identical to the sequence of any one of SEQ ID NOs: 2-203. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence of SEQ ID NO: 30. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence at least 85% identical to the sequence of SEQ ID NO: 30. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence of SEQ ID NO: 59. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence at least 85% identical to the sequence of SEQ ID NO: 59.

In some embodiments, the IL-18 polypeptide of the immunocytokine described herein comprises at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, or at least 9 amino acid substitutions, wherein the amino acid substitutions are relative to SEQ ID NO: 1. In some embodiments, the IL-18 polypeptide comprises 1 to 9 amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 1 or 2 amino acid substitutions, 1 to 3 amino acid substitutions, 1 to 4 amino acid substitutions, 1 to 5 amino acid substitutions, 1 to 6 amino acid substitutions, 1 to 7

amino acid substitutions, 1 to 8 amino acid substitutions, 2 to 3 amino acid substitutions, 2 to 4 amino acid substitutions, 2 to 5 amino acid substitutions, 2 to 6 amino acid substitutions, 2 to 7 amino acid substitutions, 2 to 8 amino acid substitutions, 2 to 9 amino acid substitutions 3 or 4 amino acid substitutions, 3 to 5 amino acid substitutions, 3 to 6 amino acid substitutions, 3 to 7 amino acid substitutions, 3 to 9 amino acid substitutions, 4 or 5 amino acid substitutions, 4 to 6 amino acid substitutions, 4 to 7 amino acid substitutions, 4 to 8 amino acid substitutions, 4 to 9 amino acid substitutions, 5 or 6 amino acid substitutions, 5 to 7 amino acid substitutions, 5 to 9 amino acid substitutions, 6 or 7 amino acid substitutions, 6 to 9 amino acid substitutions, or 7 to 9 amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 3 amino acid substitutions, 4 amino acid substitutions, 5 amino acid substitutions, 6 amino acid substitutions, 7 amino acid substitutions, or 9 amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises at most 4 amino acid substitutions, 5 amino acid substitutions, 6 amino acid substitutions, 7 amino acid substitutions, or 9 amino acid substitutions.

In some embodiments, the IL-18 polypeptide comprising of the immunocytokine described herein comprises at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, or at least 9 natural amino acid substitutions, wherein the natural amino acid substitutions are relative to SEQ ID NO: 1. In some embodiments, the IL-18 polypeptide comprises 1 to 9 natural amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 1 or 2 natural amino acid substitutions, 1 to 3 natural amino acid substitutions, 1 to 4 natural amino acid substitutions, 1 to 5 natural amino acid substitutions, 1 to 6 natural amino acid substitutions, 1 to 7 natural amino acid substitutions, 1 to 8 natural amino acid substitutions, 2 to 3 natural amino acid substitutions, 2 to 4 natural amino acid substitutions, 2 to 5 natural amino acid substitutions, 2 to 6 natural amino acid substitutions, 2 to 7 natural amino acid substitutions, 2 to 8 natural amino acid substitutions, 2 to 9 natural amino acid substitutions, 3 or 4 natural amino acid substitutions, 3 to 5 natural amino acid substitutions, 3 to 6 natural amino acid substitutions, 3 to 7 natural amino acid substitutions, 3 to 9 natural amino acid substitutions, 4 or 5 natural amino acid substitutions, 4 to 6 natural amino acid substitutions, 4 to 7 amino acid substitutions, 4 to 9 natural amino acid substitutions, 5 or 6 natural amino acid substitutions, 5 to 7 amino acid substitutions, 5 to 9 natural amino acid substitutions, 6 or 7 natural amino acid substitutions, 6 to 9 natural amino acid substitutions, or 7 to 9 natural amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 3 natural amino acid substitutions, 4 natural amino acid substitutions, 5 amino acid substitutions, 6 natural amino acid substitutions, 7 natural amino acid substitutions, or 9 natural amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises at most 4 natural amino acid substitutions, 5 natural amino acid substitutions, 6 natural amino acid substitutions, 7 natural amino acid substitutions, or 9 natural amino acid substitutions. In some embodiments, the IL-18 polypeptide further comprises up to 10 non-canonical amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 unnatural amino acid substitutions. In some embodiments, the IL-18 polypeptide further comprises unnatural amino acid substitutions at residues M33, M51, N60, M86, M113, and/or M150. In some embodiments, the unnatural amino acid residues substituted for the methionines are each independently norleucine or O-methyl-homoserine. In some embodiments, the IL-18 polypeptide further unnatural

amino acid substitutions at residues 31, 75, and 116. In some embodiments, the IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 75, and Hse 116. In some embodiments, the IL-18 polypeptide further unnatural amino acid substitutions at residues 31, 63, and 116. In some embodiments, the IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 63, and Hse 116. In some embodiments, the modified IL-18 polypeptide further unnatural amino acid substitutions at residues 31, 63, 75, and 116. In some embodiments, the modified IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 63, Hse 75, and Hse 116. In some embodiments, the modified IL-18 polypeptide further unnatural amino acid substitutions at residues 31, 67, 75, and 116. In some embodiments, the modified IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 67, Hse 75, and Hse 116. In some embodiments, the modified IL-18 polypeptide further unnatural amino acid substitutions at residues 31, 57, 75, and 116. In some embodiments, the modified IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 57, Hse 75, and Hse 116. In some embodiments, the modified IL-18 polypeptide further unnatural amino acid substitutions at residues 31, 50, 75, and 116. In some embodiments, the modified IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 50, Hse 75, and Hse 116. In some embodiments, the modified IL-18 polypeptide further unnatural amino acid substitutions at residues 31, 50, 75, and 121. In some embodiments, the modified IL-18 polypeptide further comprises homoserine (Hse) 31, Hse 50, Hse 75, and Hse 121.

In some embodiments, the IL-18 polypeptide of the immunocytokine described herein comprises at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, or at least 9 amino acid substitutions, wherein the amino acid substitutions are relative to any one of SEQ ID NOs: 68, 92, 116, 140, or 170. In some embodiments, a modified IL-18 polypeptide described herein comprises at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, or at least 9 additional amino acid substitutions, wherein the amino acid substitutions are relative to any one of SEQ ID NOs: 68, 92, 116, 140, or 170. In some embodiments, the IL-18 polypeptide comprises 1 to 9 amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 1 or 2 amino acid substitutions, 1 to 3 amino acid substitutions, 1 to 4 amino acid substitutions, 1 to 5 amino acid substitutions, 1 to 6 amino acid substitutions, 1 to 7 amino acid substitutions, 1 to 8 amino acid substitutions, 2 to 3 amino acid substitutions, 2 to 4 amino acid substitutions, 2 to 5 amino acid substitutions, 2 to 6 amino acid substitutions, 2 to 7 amino acid substitutions, 2 to 8 amino acid substitutions, 2 to 9 amino acid substitutions 3 or 4 amino acid substitutions, 3 to 5 amino acid substitutions, 3 to 6 amino acid substitutions, 3 to 7 amino acid substitutions, 3 to 9 amino acid substitutions, 4 or 5 amino acid substitutions, 4 to 6 amino acid substitutions, 4 to 7 amino acid substitutions, 4 to 9 amino acid substitutions, 5 or 6 amino acid substitutions, 5 to 7 amino acid substitutions, 5 to 9 amino acid substitutions, 6 or 7 amino acid substitutions, 6 to 9 amino acid substitutions, or 7 to 9 amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises 3 amino acid substitutions, 4 amino acid substitutions, 5 amino acid substitutions, 6 amino acid substitutions, 7 amino acid substitutions, or 9 amino acid substitutions. In some embodiments, the IL-18 polypeptide comprises at most 4 amino acid substitutions, 5 amino acid substitutions, 6 amino acid substitutions, 7 amino acid substitutions, or 9 amino acid substitutions.

In some embodiments, one modification is at amino acid residue 6. In some embodiments, one modification is in the range of amino acid residues 53-63. In some embodiments, one modification is at amino acid residue 53. In some embodiments, one modification is at amino acid residue 63.

In some embodiments, the IL-18 polypeptide comprises at least one modification to the amino acid sequence of SEQ ID NO: 1 selected from: Y01X, F02X, E06X, SOX, V11X, D17X, C38X, M51X, K53X, D54X, S55X, T63X, C68X, C76X, AND C127X, wherein each X is independently a natural or non-natural amino acid. In some embodiments, the IL-18 polypeptide further comprises an amino acid substitution at the point of attachment of the linker, such as residue 69, residue 70, residue 85, residue 86, residue 95, or residue 98. In some embodiments, the IL-18 polypeptide comprises at least one modification to the amino acid sequence of SEQ ID NO: 1 selected from: Y01G, F02A, E06K, S10T, V11I, D17N, C38S, C38A, C38Q, M51G, K53A, D54A, S55A, T63A, C68S, C68A, C76S, C76A, C127A, and C127S. In some embodiments, the IL-18 polypeptide further comprises an amino acid substitution at the point of attachment of the linker, such as E69C, K70C, E85C, M86C, T95C, or D98C.

In one aspect, described herein is a modified interleukin-18 (IL-18) polypeptide comprising E06K and K53A, wherein residue position numbering of the IL-18 polypeptide is based on SEQ ID NO: 1 as a reference sequence. In some embodiments, the IL-18 polypeptide further comprises V11I.

In some embodiments, the IL-18 polypeptide further comprises T63A. In some embodiments, the IL-18 polypeptide further comprises at least one of Y01X, S55X, F02X, D54X, C38X, C68X, E69X, K70X, C76X, or C127X, wherein each X is independently an amino acid or an amino acid derivative. In some embodiments, the IL-18 polypeptide further comprises at least one of Y01G, S55A, F02A, D54A, C38S, C38A, C38Q, C68S, C68A, E69C, K70C, C76S, C76A, C127S, or C127A. In some embodiments, the IL-18 polypeptide further comprises an amino acid substitution at the point of attachment of the linker, such as residue 69, residue 70, residue 85, residue 86, residue 95, or residue 98.

In some embodiments, the IL-18 peptide comprises at least one modification to the amino acid sequence of SEQ ID NO: 1, wherein the modification is E06X, V11X, K53X, S55X, or T63X, wherein X is a natural or non-natural amino acid. In some embodiments, the IL-18 peptide comprises at least two modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06X and K53X; E06X and S55X; K53X and S55X; E06X and T63X; or K53X and T63X, wherein X is a natural or non-natural amino acid. In some embodiments, the IL-18 peptide comprises at least three modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06X, K53X, and S55X; or E06X, K53X, and T63X, wherein X is a natural or non-natural amino acid. In some embodiments, the IL-18 peptide comprises at least four modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06X, K53X, S55X, and T63X; E06X, K53X, S55X, and Y01X; E06X, K53X, S55X, and F02X; E06X, K53X, S55X, and D54X; E06X, K53X, S55X, and M51X; or C38X, C68X, C76X, and C127X, wherein X is a natural or non-natural amino acid. In each embodiment wherein a plurality of amino acids residues are replaced with a natural or non-natural amino acid X, each X is independently the same or a different amino acid.

In some embodiments, the IL-18 peptide comprises at least one modification to the amino acid sequence of SEQ ID NO: 1, wherein the modification is E06K, V11, K53A, S55A, or T63A. In some embodiments, the IL-18 peptide comprises at least two modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06K and K53A; E06K and S55A; K53A and S55A; E06K and T63A; or K53A and T63A. In some embodiments, the IL-18 peptide comprises at least three modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06K, K53A, and S55A; E06K, V11I, and K53A; E06K, C38A, and K53A; or E06K, K53A, and T63A. In some embodiments, the IL-18 peptide comprises at least four modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06K, K53A, S55A, and T63A; E06K, K53A, S55A, and Y01G; E06K, K53A, S55A, and F02A; E06K, K53A, S55A, and D54A; E06K, K53A, S55A, and M51G; or C38S, C68S, C76S, and C127S. In some embodiments, the IL-18 peptide comprises at least six modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise E06K, K53A, C38S, C68S, C76S, and C127S; or K53A, T63A, C38S, C68S, C76S, and C127S. In some embodiments, the modified IL-18 polypeptide comprises at least seven modifications to the sequence of SEQ ID NO: 1, wherein the seven modifications comprise E6K, V11, C38A, K53A, T63A, C76A, C127A. In some embodiments, the IL-18 peptide comprises at least eight modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise Y01G, F02A, E06K, M51G, K53A, D54A, S55A, and T63A. In some embodiments, the IL-18 peptide comprises at least eight modifications to the amino acid sequence of SEQ ID NO: 1, wherein the modifications comprise Y01G, F02A, E06K, M51G, K53A, S55A, and T63A.

In one aspect, provided herein, is a modified IL-18 polypeptide as provided herein, further comprising E06K and K53A, wherein residue position numbering of the IL-18 polypeptide is based on SEQ ID NO: 1 as a reference sequence. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence at least 80%, at least 85%, at least 90%, at least 95%, or at least 98% identical to the amino acid sequence of SEQ ID NO: 30. In some embodiments, the IL-18 polypeptide comprises an amino acid sequence at least 80%, at least 85%, at least 90%, at least 95%, or at least 98% identical to the amino acid sequence of SEQ ID NO: 59. In some embodiments, the IL-18 polypeptide further comprises an amino acid substitution at one or more cysteine residues. In some embodiments, the IL-18 polypeptide comprises one or more cysteines substituted with either serine or alanine. In some embodiments, the IL-18 polypeptide comprise amino acid substitutions at each cysteine residue of SEQ ID NO: 1. In some embodiments, each cysteine residue is substituted with serine or alanine. In some embodiments, the IL-18 polypeptide comprises amino acid substitutions at 1, 2, 3, 4, 5, or 6 methionine residues. In some embodiments, each substitution at a methionine residue is for an O-methyl-L-homoserine residue or a norleucine residue. In some embodiments, each methionine residue is substituted with an O-methyl-L-homoserine residue. In some embodiments, the IL-18 polypeptide comprises homoserine residues at positions 31, 116, and one of 63 and 75. In some embodiments, the modified IL-18 polypeptide comprises homoserine residues at positions 31, 116, 75, and one of 50, 57, 63, and 67. In some

embodiments, the modified IL-18 polypeptide comprises homoserine residues at positions 31, 121, 75, and one of 50, 57, 63, and 67.

In some embodiments, the IL-18 polypeptide comprises a polypeptide sequence having at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 98%, at least about 99%, or about 100% sequence identity to SEQ ID NO: 2-12. In some embodiments, the IL-18 polypeptide comprises a polypeptide sequence having at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 98%, at least about 99%, or about 100% sequence identity to SEQ ID NO: 13-23. In some embodiments, the modified IL-18 polypeptide comprises a polypeptide sequence having at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 98%, or at least about 99% sequence identity to SEQ ID NO: 24-33. In some embodiments, the polypeptide sequence is at least about 80% identical to SEQ ID NO: 30 or SEQ ID NO: 59. In some embodiments, the polypeptide sequence is at least about 80% identical to SEQ ID NO: 30. In some embodiments, the polypeptide sequence is at least about 90% identical to SEQ ID NO: 30. In some embodiments, the polypeptide sequence is at least about 95% identical to SEQ ID NO: 30. In some embodiments, the polypeptide sequence is identical to SEQ ID NO: 30. In some embodiments, the polypeptide sequence is at least about 80% identical to SEQ ID NO: 59. In some embodiments, the polypeptide sequence is at least about 90% identical to SEQ ID NO: 59. In some embodiments, the polypeptide sequence is at least about 95% identical to SEQ ID NO: 59. In some embodiments, the IL-18 polypeptide is recombinant. In some embodiments, the IL-18 polypeptide is one provided in Table 2. In some embodiments, the IL-18 polypeptide is one described in Table 3.

#### 35 Biological Activity

In some embodiments, the immunocytokine composition exhibits one or more activities associated with the antibody or antigen binding fragment and/or an IL-18 polypeptide.

In some embodiments, the immunocytokine composition 40 exhibits an ability to bind to the IL-18 receptor. In some embodiments, the immunocytokine composition exhibits an ability to bind to the IL-18 receptor which is comparable to WT IL-18. In some embodiments, immunocytokine composition exhibits an ability to bind to the IL-18 receptor (IL-18R $\alpha\beta$ ) which is reduced by at most 2-fold, at most 5-fold, at most 10-fold, at most 20-fold, at most 50-fold, at most 100-fold, at most 200-fold, at most 300-fold, at most 400-fold, or at most 1000-fold compared to WT IL-18. In some embodiments, the immunocytokine composition exhibits an enhanced ability to bind the IL-18R $\alpha\beta$ . In some embodiments, the immunocytokine composition exhibits an ability to bind to the IL-18R $\alpha\beta$  which is increased by at least 2-fold, at least 3-fold, at least 5-fold, or at least 10-fold compared to WT IL-18.

In some embodiments, immunocytokine composition 55 exhibits an ability to stimulate production of IFN $\gamma$  upon contact with a cell (e.g., an immune cell, such as an NK cell). In some embodiments, the ability of the immunocytokine composition to stimulate IFN $\gamma$  production is somewhat reduced compared to WT IL-18. In some embodiments, a half-maximal effective concentration ( $EC_{50}$ ) of the ability of the immunocytokine composition to stimulate production of IFN $\gamma$  is at most 100-fold higher than, at most 50-fold higher than, at most 20-fold higher than, at most 10-fold higher than, at most 5-fold higher than, or at most 2-fold higher than that of a WT IL-18. In some embodiments, the ability 60 of the immunocytokine composition to stimulate IFN $\gamma$  pro-

duction is enhanced compared to WT IL-18. In some embodiments, a half-maximal effective concentration ( $EC_{50}$ ) of the ability of the immunocytokine composition to stimulate production of IFN $\gamma$  is at least 5-fold lower than, at least 10-fold lower than, at least 20-fold lower than, at least 50-fold lower than, at least 75-fold lower than, or at least 100-fold higher than that of a WT IL-18.

In some embodiments, the immunocytokine composition exhibits an ability to stimulate production of IFN $\gamma$  upon contact with a cell (e.g., an immune cell, such as an NK cell) which is only somewhat reduced as compared to the IL-18 polypeptide not comprised in the immunocytokine composition (e.g., unconjugated IL-18 polypeptide). In some embodiments, the  $EC_{50}$  of IFN $\gamma$  stimulation is at most 5-fold greater than, at most 10-fold greater than, at most 50-fold greater than, or at most 100-fold greater than that of the IL-18 polypeptide not comprised in the immunocytokine composition. In some embodiments, the immunocytokine composition exhibits an ability to induce IFN $\gamma$  production in a cell as measured by half-maximal effective concentration ( $EC_{50}$ ) which is within about 100-fold of the corresponding IL-18 polypeptide not comprised in the immunocytokine composition. In some embodiments, the immunocytokine composition exhibits a lower  $EC_{50}$  than WT IL-18. In some embodiments, the immunocytokine composition exhibits a lower  $EC_{50}$  than WT IL-18 by at least 2-fold, 5-fold, 10-fold, 20-fold, 30-fold, 50-fold, or 100-fold.

In some embodiments, the immunocytokine composition exhibits a reduced ability to bind IL-18 binding protein (IL-18BP). In some embodiments, the ability of immunocytokine composition to bind IL-18BP is reduced by at least 2-fold, at least 5-fold, at least 10-fold, at least 20-fold, at least 30-fold, at least 50-fold, or at least 100-fold compared to WT IL-18. In some embodiments, the immunocytokine composition does not display any substantial ability to bind IL-18 BP.

In some embodiments, the immunocytokine composition exhibits a reduced ability to have its IFN $\gamma$  production stimulatory activity inhibited by IL-18BP. In some embodiments, the ability of the immunocytokine composition to be inhibited by IL-18BP is measured as a half maximal inhibitory concentration ( $IC_{50}$ ). In some embodiments, the immunocytokine composition exhibits an  $IC_{50}$  by IL-18BP that is at least 2-fold higher than, at least 5-fold higher than, at least 10-fold higher than, at least 15-fold higher than, at least 20-fold higher than, at least 25-fold higher than, at least 30-fold higher than, at least 40-fold higher than, or at least 50-fold higher than an  $IC_{50}$  of WT IL-18's inhibition by IL-18BP. In some embodiments, the immunocytokine composition exhibits an  $IC_{50}$  by IL-18BP that is at least 100-fold higher than an  $IC_{50}$  of WT IL-18's inhibition by IL-18BP. In some embodiments, the immunocytokine composition exhibits an  $IC_{50}$  by IL-18BP that is at least 200-fold higher than an  $IC_{50}$  of WT IL-18's inhibition by IL-18BP. In some embodiments, the immunocytokine composition exhibits an  $IC_{50}$  by IL-18BP that is at least 500-fold higher than an  $IC_{50}$  of WT IL-18's inhibition by IL-18BP. In some embodiments, the immunocytokine composition exhibits an  $IC_{50}$  by IL-18BP that is at least 1000-fold higher than an  $IC_{50}$  of WT IL-18's inhibition by IL-18BP.

In some embodiments, the immunocytokine composition retains binding associated with the antibody or antigen binding fragment. In some embodiments, the immunocytokine composition retains binding to the antigen of the antibody or antigen binding fragment. In some embodiments, the immunocytokine composition exhibits binding affinity ( $K_D$ ) to the antigen of the antibody which is within

5-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits binding affinity ( $K_D$ ) to the antigen of the antibody which is within 2.5-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the binding is determined by ELISA. In some embodiments, the binding is determined by BLI.

In some embodiments, the immunocytokine composition retains binding to one or more Fc receptors associated with the antibody or antigen binding fragment. In some embodiments, the Fc receptor is selected from FcRn, CD64, CD32a, CD16, and CD32b, or any combination thereof. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to at least one Fc receptor which is within 10-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to at least one Fc receptor which is less than 10-fold higher, less than 5-fold higher, less than 4-fold higher, less than 3-fold higher, less than 2-fold higher, or less than the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to each of FcRn, CD64, CD32a, CD16, and CD32B which is less than 10-fold higher, less than 5-fold higher, less than 4-fold higher, less than 3-fold higher, less than 2-fold higher, or less than the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to each of FcRn, CD64, CD32a, CD16, and CD32B which is within 10-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to each of FcRn, CD64, CD32a, CD16, and CD32B which is within 20-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to each of FcRn, CD64, CD32a, CD16, and CD32B which is within 50-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide. In some embodiments, the immunocytokine composition exhibits a binding affinity ( $K_D$ ) to each of FcRn, CD64, CD32a, CD16, and CD32B which is within 100-fold of the binding affinity of the antibody not attached to the IL-18 polypeptide.

In some embodiments, the immunocytokine composition exhibits synergistic efficacy owing to the presence of both molecules in one molecule. In some embodiments, the immunocytokine composition exhibits enhanced activity compared to either molecule alone. In some embodiments, the immunocytokine composition exhibits enhanced anti-tumor growth inhibition compared to the antibody alone. In some embodiments, the immunocytokine composition exhibits enhanced anti-tumor growth inhibition compared to the antibody and the IL-18 polypeptide administered in combination. In some embodiments, the IL-18 polypeptide is administered as a half-life extended version (e.g., PEGylated, attached to an Fc region (e.g., an Fc fusion), or attached to a negative control antibody). In some embodiments, the immunocytokine composition exhibits similar or enhanced antitumor activity at the same concentration as the antibody administered alone. In some embodiments, the immunocytokine composition exhibits similar or enhanced antitumor activity when administered at a dose which is less than 0.5-fold, 0.25-fold, or 0.1-fold the dose of the antibody alone.

## Orthogonal Payloads

In one non-limiting instant, the antibody/IL-18 immunocytokines of the disclosure can comprise dual orthogonal payloads. The antibody/IL-18 immunocytokines can comprise an antibody, one IL-18 polypeptide, and one payload linked to the antibody by a chemical orthogonal linking group. The orthogonal payload can be an amino acid, amino acid derivative, peptide, protein, cytokine, alkyl group, aryl or heteroaryl group, therapeutic small molecule drug, polyethylene glycol (PEG) moiety, lipid, sugar, biotin, biotin derivative, deoxyribonucleic acid (DNA), ribonucleic acid (RNA), or peptide nucleic acid (PNA), any of which is substituted, unsubstituted, modified, or unmodified. In some embodiments, the orthogonal payload is a therapeutic small molecule. In some embodiments, the orthogonal payload is a PEG moiety. In some embodiments, the orthogonal payload is an additional cytokine such as, for example, IL-7 or IL-2. In one exemplary instance, human IL-7 has an amino acid sequence of DCDIEGKDGKQYESVLMVSIDQLL-DSMKEIGSNCLNNEFPNFFKRHICDANKEGMFLFRA ARKLRQFLKMNSTGDFDLHLLKVLSEGT-TILLNCTGQVKGRKPAALGEAQPTKSLEENKS LKEQKKLNNDLCFLKRLQQEIKTCWNKILMGTKEH (SEQ ID NO: 210), or is a modified human IL-7 (e.g., an IL-7 having at least about 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% sequence identity to human IL-7). In one exemplary instance, human IL-2 has an amino acid sequence of APTSSSTKKTQLQLEHLLL-LQMILNGINNYKNPKLTRMLTFKFYMPK-KATELKHLQCL EEELKPLEEVNLNAQSKNFHLR-PRDLISNINVIVLELGKSETTFMCEYADETATIVEFLNR WITFCQSIISTLT (SEQ ID NO: 211), or is a modified human IL-2 (e.g., an IL-2 having at least about 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% sequence identity to human IL-2).

## Compositions

In one aspect, provided herein is a pharmaceutical composition comprising an antibody linked to a modified IL-18 polypeptide described herein; and a pharmaceutically acceptable carrier or excipient. In some embodiments, the pharmaceutical composition further comprises one or more excipients, wherein the one or more excipients include, but are not limited to, a carbohydrate, an inorganic salt, an antioxidant, a surfactant, a buffer, or any combination thereof. In some embodiments the pharmaceutical composition further comprises one, two, three, four, five, six, seven, eight, nine, ten, or more excipients, wherein the one or more excipients include, but are not limited to, a carbohydrate, an inorganic salt, an antioxidant, a surfactant, a buffer, or any combination thereof.

In some embodiments, the pharmaceutical composition further comprises a carbohydrate. In certain embodiments, the carbohydrate is selected from the group consisting of fructose, maltose, galactose, glucose, D-mannose, sorbose, lactose, sucrose, trehalose, cellobiose raffinose, melezitose, maltodextrins, dextrans, starches, mannitol, xylitol, maltitol, lactitol, xylitol, sorbitol (glucitol), pyranosyl sorbitol, myoinositol, cyclodextrins, and combinations thereof.

Alternately, or in addition, the pharmaceutical composition further comprises an inorganic salt. In certain embodiments, the inorganic salt is selected from the group consisting of sodium chloride, potassium chloride, magnesium chloride, calcium chloride, sodium phosphate, potassium phosphate, sodium sulfate, or combinations thereof.

Alternately, or in addition, the pharmaceutical composition further comprises an antioxidant. In certain embodiments, the antioxidant is selected from the group consisting

of ascorbyl palmitate, butylated hydroxyanisole, butylated hydroxytoluene, potassium metabisulfite, propyl gallate, sodium metabisulfite, sodium thiosulfate, vitamin E, 3,4-dihydroxybenzoic acid, and combinations thereof.

5 Alternately, or in addition, the pharmaceutical composition further comprises a surfactant. In certain embodiments, the surfactant is selected from the group consisting of polysorbates, sorbitan esters, lipids, phospholipids, phosphatidylethanolamines, fatty acids, fatty acid esters, steroids, EDTA, zinc, and combinations thereof.

10 Alternately, or in addition, the pharmaceutical composition further comprises a buffer. In certain embodiments, the buffer is selected from the group consisting of citric acid, sodium phosphate, potassium phosphate, acetic acid, ethanolamine, histidine, amino acids, tartaric acid, succinic acid, fumaric acid, lactic acid, tris, HEPES, or combinations thereof.

15 In some embodiments, the pharmaceutical composition is formulated for parenteral or enteral administration. In some 20 embodiments, the pharmaceutical composition is formulated for intravenous (IV) or subcutaneous (SQ) administration. In some embodiments, the pharmaceutical composition is in a lyophilized form.

25 In one aspect, described herein is a liquid or lyophilized composition that comprises a described antibody or antigen binding fragment linked to a modified IL-18 polypeptide. In some embodiments, the antibody or antigen binding fragment linked to the IL-18 polypeptide modified IL-18 polypeptide is a lyophilized powder. In some embodiments, the lyophilized powder is resuspended in a buffer solution. In some embodiments, the buffer solution comprises a buffer, a sugar, a salt, a surfactant, or any combination thereof. In some embodiments, the buffer solution comprises a phosphate salt. In some embodiments, the phosphate salt is sodium 30 Na<sub>2</sub>HPO<sub>4</sub>. In some embodiments, the salt is sodium chloride. In some embodiments, the buffer solution comprises phosphate buffered saline. In some embodiments, the buffer solution comprises mannitol. In some embodiments, the lyophilized powder is suspended in a solution comprising about 10 mM Na<sub>2</sub>HPO<sub>4</sub> buffer, about 0.022% SDS, and about 50 mg/mL mannitol, and having a pH of about 7.5.

## Dosage Forms

35 The immunocytokine compositions described herein can be in a variety of dosage forms. In some embodiments, the 40 immunocytokine composition is dosed as a lyophilized powder. In some embodiments, the immunocytokine composition is dosed as a suspension. In some embodiments, the immunocytokine composition is dosed as a solution. In some embodiments, the immunocytokine composition is dosed as an injectable solution. In some embodiments, the immunocytokine composition is dosed as an IV solution.

## Methods of Treatment

45 In one aspect, described herein, is a method of treating cancer in a subject in need thereof, comprising: administering 50 to the subject an effective amount of an immunocytokine composition or a pharmaceutical composition as described herein. In some embodiments, the cancer is a solid cancer. A cancer or tumor can be, for example, a primary cancer or tumor or a metastatic cancer or tumor. In some embodiments, the cancer is a solid cancer. In some embodiments, the solid cancer is adrenal cancer, anal cancer, bile duct cancer, bladder cancer, bone cancer, brain cancer, breast cancer, carcinoid cancer, cervical cancer, colorectal cancer, esophageal cancer, eye cancer, gallbladder cancer, gastrointestinal stromal tumor, germ cell cancer, head and neck cancer, kidney cancer, liver cancer, lung cancer, nasal cavity and paranasal sinus cancer, nasopharyngeal cancer, neuro-

blastoma, neuroendocrine cancer, oral cancer, oropharyngeal cancer, ovarian cancer, pancreatic cancer, pediatric cancer, penile cancer, pituitary cancer, prostate cancer, skin cancer, soft tissue cancer, spinal cord cancer, stomach cancer, testicular cancer, thymus cancer, thyroid cancer, ureteral cancer, uterine cancer, vaginal cancer, or vulvar cancer.

In some embodiments, the cancer is a blood cancer. In some embodiments, the blood cancer is leukemia, non-Hodgkin lymphoma, Hodgkin lymphoma, an AIDS-related lymphoma, multiple myeloma, plasmacytoma, post-transplantation lymphoproliferative disorder, or Waldenstrom macroglobulinemia.

Combination therapies with one or more additional active agents are contemplated herein.

An effective response is achieved when the subject experiences partial or total alleviation or reduction of signs or symptoms of illness, and specifically includes, without limitation, prolongation of survival. The expected progression-free survival times may be measured in months to years, depending on prognostic factors including the number of relapses, stage of disease, and other factors. Prolonging survival includes without limitation times of at least 1 month (mo), about at least 2 mos., about at least 3 mos., about at least 4 mos., about at least 6 mos., about at least 1 year, about at least 2 years, about at least 3 years, about at least 4 years, about at least 5 years, etc. Overall or progression-free survival can be also measured in months to years. Alternatively, an effective response may be that a subject's symptoms or cancer burden remain static and do not worsen. Further treatment of indications are described in more detail below. In some instances, a cancer or tumor is reduced by at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, or 100%.

In some embodiments, the immunocytokine composition is administered in a single dose of the effective amount of immunocytokine composition, including further embodiments in which (i) the immunocytokine composition is administered once a day; or (ii) the immunocytokine composition is administered once a day; or (ii) the immunocytokine composition is administered to the subject multiple times over the span of one day. In some embodiments, the conjugate is administered daily, every other day, twice a week, 3 times a week, once a week, every 2 weeks, every 3 weeks, every 4 weeks, every 5 weeks, every 6 weeks, every 12 weeks, every 3 days, every 4 days, every 5 days, every 6 days, 2 times a week, 3 times a week, 4 times a week, 5 times a week, 6 times a week, once a month, twice a month, 3 times a month, 4 times a month, once every 2 months, once every 3 months, once every 4 months, once every 5 months, or once every 6 months. Administration includes, but is not limited to, injection by any suitable route (e.g., parenteral, enteral, intravenous, subcutaneous, etc.).

#### Methods of Manufacturing

In one aspect, described herein, is a method of making an immunocytokine composition, comprising providing an antibody or antigen binding fragment thereof (e.g., an antibody or antigen binding fragment provided herein), wherein the antibody comprises a reactive group (e.g., a conjugation handle), contacting the reactive group with a complementary reactive group attached to an IL-18 polypeptide, and forming the immunocytokine composition. The resulting composition is any of the compositions provided herein.

In some embodiments, providing the antibody comprising the reactive group comprises attaching the reactive group to the antibody. In some embodiments, the reactive group is added site-specifically. In some embodiments, attaching the

reactive group to the antibody comprises contacting the antibody with an affinity group comprising a reactive functionality which forms a bond with a specific residue of the antibody. In some embodiments, attaching the reactive group to the antibody comprises contacting the antibody with an enzyme. In some embodiments, the enzyme is configured to site-specifically attach the reactive group to a specific residue of the antibody. In some embodiments, the enzyme is a glycosylation enzyme or a transglutaminase enzyme.

In some embodiments, the method further comprises attaching the complementary reactive group to the cytokine. Definitions

All terms are intended to be understood as they would be understood by a person skilled in the art. Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which the disclosure pertains.

The following definitions supplement those in the art and are directed to the current application and are not to be imputed to any related or unrelated case, e.g., to any commonly owned patent or application. Although any methods and materials similar or equivalent to those described herein can be used in the practice for testing of the present disclosure, the preferred materials and methods are described herein. Accordingly, the terminology used herein is for the purpose of describing particular embodiments only and is not intended to be limiting.

The terminology used herein is for the purpose of describing particular cases only and is not intended to be limiting. In this application, the use of the singular includes the plural unless specifically stated otherwise. As used herein, the singular forms "a", "an" and "the" are intended to include the plural forms as well, unless the context clearly indicates otherwise.

In this application, the use of "or" means "and/or" unless stated otherwise. The terms "and/or" and "any combination thereof" and their grammatical equivalents as used herein, can be used interchangeably. These terms can convey that any combination is specifically contemplated. Solely for illustrative purposes, the following phrases "A, B, and/or C" or "A, B, C, or any combination thereof" can mean "A individually; B individually; C individually; A and B; B and C; A and C; and A, B, and C." The term "or" can be used conjunctively or disjunctively, unless the context specifically refers to a disjunctive use.

The term "about" or "approximately" can mean within an acceptable error range for the particular value as determined by one of ordinary skill in the art, which will depend in part on how the value is measured or determined, i.e., the limitations of the measurement system. For example, "about" can mean within 1 or more than 1 standard deviation, per the practice in the art. Alternatively, "about" can mean a range of up to 20%, up to 15%, up to 10%, up to 5%, or up to 1% of a given value. Alternatively, particularly with respect to biological systems or processes, the term can mean within an order of magnitude, within 5-fold, or within 2-fold, of a value. Where particular values are described in the application and claims, unless otherwise stated the term "about" meaning within an acceptable error range for the particular value should be assumed.

As used in this specification and claim(s), the words "comprising" (and any form of comprising, such as "comprise" and "comprises"), "having" (and any form of having, such as "have" and "has"), "including" (and any form of including, such as "includes" and "include") or "containing" (and any form of containing, such as "contains" and "con-

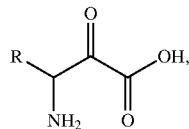
## 101

tain") are inclusive or open-ended and do not exclude additional, unrecited elements or method steps. It is contemplated that any embodiment discussed in this specification can be implemented with respect to any method or composition of the present disclosure, and vice versa. Furthermore, compositions of the present disclosure can be used to achieve methods of the present disclosure.

Reference in the specification to "some embodiments," "an embodiment," "one embodiment" or "other embodiments" means that a particular feature, structure, or characteristic described in connection with the embodiments is included in at least some embodiments, but not necessarily all embodiments, of the present disclosures. To facilitate an understanding of the present disclosure, a number of terms and phrases are defined below.

Referred to herein are groups which are "attached" or "covalently attached" to residues of IL-18 polypeptides or other polypeptides. As used herein, "attached" or "covalently attached" means that the group is tethered to the indicated residue, and such tethering can include a linking group (i.e., a linker). Thus, for a group "attached" or "covalently attached" to a residue, it is expressly contemplated that such linking groups are also encompassed.

As used herein, an "alpha-keto amino acid" or the phrase "alpha-keto" before the name of an amino acid refers to an amino acid or amino acid derivative having a ketone functional group positioned between the carbon bearing the amino group and the carboxylic acid of an amino acid. Alpha-keto amino acids of the instant disclosure have a structure as set forth in the following formula:



wherein R is the side chain of any natural or unnatural amino acid. The R functionality can be in either the L or D orientation in accordance with standard amino acid nomenclature. In preferred embodiments, alpha-keto amino acids are in the L orientation. When the phrase "alpha-keto" is used before the name of a traditional natural amino acid (e.g., alpha-keto leucine, alpha-keto phenylalanine, etc.) or a common unnatural amino acid (e.g., alpha-keto norleucine, alpha-keto O-methyl-homoserine, etc.), it is intended that the alpha-keto amino acid referred to matches the above formula with the side chain of the referred to amino acid. When an alpha-keto amino acid residue is set forth in a peptide or polypeptide sequence herein, it is intended that a protected version of the relevant alpha-keto amino acid is also encompassed (e.g., for a sequence terminating in a C-terminal alpha-keto amino acid, the terminal carboxylic acid group may be appropriately capped with a protecting group such as a tert-butyl group, or the ketone group with an acetal protecting group). Other protecting groups encompassed are well known in the art.

Binding affinity refers to the strength of a binding interaction between a single molecule and its ligand/binding partner. A higher binding affinity refers to a higher strength bond than a lower binding affinity. In some instances, binding affinity is measured by the dissociation constant ( $K_D$ ) between the two relevant molecules. When comparing  $K_D$  values, a binding interaction with a lower value will have a higher binding affinity than a binding interaction with a

## 102

higher value. For a protein-ligand interaction,  $K_D$  is calculated according to the following formula:

$$5 \quad K_D = \frac{[L][P]}{[LP]}$$

where [L] is the concentration of the ligand, [P] is the concentration of the protein, and [LP] is the concentration of the ligand/protein complex.

Referred to herein are certain amino acid sequences (e.g., polypeptide sequences) which have a certain percent sequence identity to a reference sequence or refer to a residue at a position corresponding to a position of a reference sequence. Sequence identity is measured by protein-protein BLAST algorithm using parameters of Matrix BLOSUM62, Gap Costs Existence:11, Extension:1, and Compositional Adjustments Conditional Compositional Score Matrix Adjustment. This alignment algorithm is also used to assess if a residue is at a "corresponding" position through an analysis of the alignment of the two sequences being compared.

Unless otherwise specified, is contemplated that "protected" versions of amino acids (e.g., those containing a chemical protecting group affixed to a functionality of the amino acid, particularly a side chain of the amino acid but also at another point of the amino acid) qualify as the same amino acid as the "unprotected" version for sequence identity purposes, particularly for chemically synthesized polypeptides. It is also contemplated that such protected versions are also encompassed by the SEQ ID NOs provided herein. Non-limiting examples of protecting groups which may be encompassed include fluorenylmethyloxycarbonyl (Fmoc), triphenylmethyl (trityl or trt), tert-Butyloxycarbonyl (Boc), 2,2,4,6,7-pentamethylidihydrobenzofuran-5-sulfonyl (Pbf), acetamidomethyl (AcM), tert-butyl (tBu or OtBu), 2,2-dimethyl-1-(4-methoxyphenyl)propane-1,3-diol ketal or acetal, and 2,2-dimethyl-1-(2-nitrophenyl)propane-1,3-diol ketal or acetal. Other protecting groups well known in the art are also encompassed. Similarly, modified versions of natural amino acids are also intended to qualify as natural version of the amino acid for sequence identity purposes. For example, an amino acid comprising a side chain heteroatom which can be covalently modified (e.g., to add a conjugation handle, optionally through a linker), such as a lysine, glutamine, glutamic acid, asparagine, aspartic acid, cysteine, or tyrosine, which has been covalently modified would be counted as the base amino acid (see, e.g., Structure 2 below, which would be counted as a lysine for sequence identity and SEQ ID purposes). Similarly, an amino acid comprising another group added to the C or N-terminus would be counted as the base amino acid.

The term "pharmaceutically acceptable" refers to approved or approvable by a regulatory agency of the federal or a state government or listed in the U.S. Pharmacopeia (U.S.P.) or other generally recognized pharmacopeia for use in animals, including humans.

A "pharmaceutically acceptable excipient, carrier, or diluent" refers to an excipient, carrier, or diluent that can be administered to a subject, together with an agent, and which does not destroy the pharmacological activity thereof and is nontoxic when administered in doses sufficient to deliver a therapeutic amount of the agent.

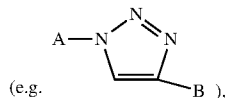
A "pharmaceutically acceptable salt" suitable for the disclosure may be an acid or base salt that is generally considered in the art to be suitable for use in contact with the

103

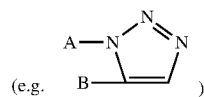
tissues of human beings or animals without excessive toxicity, irritation, allergic response, or other problem or complication. Such salts include mineral and organic acid salts of basic residues such as amines, as well as alkali or organic salts of acidic residues such as carboxylic acids. Specific pharmaceutical salts include, but are not limited to, salts of acids such as hydrochloric, phosphoric, hydrobromic, malic, glycolic, fumaric, sulfuric, sulfamic, sulfanilic, formic, toluenesulfonic, methanesulfonic, benzene sulfonic, ethane disulfonic, 2-hydroxyethyl sulfonic, nitric, benzoic, 2-acetoxybenzoic, citric, tartaric, lactic, stearic, salicylic, glutamic, ascorbic, pamoic, succinic, fumaric, maleic, propionic, hydroxymaleic, hydroiodic, phenylacetic, alkanoic such as acetic, HOOC—(CH<sub>2</sub>)<sub>n</sub>-COOH where n is 0-4, and the like. Similarly, pharmaceutically acceptable cations include, but are not limited to sodium, potassium, calcium, aluminum, lithium and ammonium. Those of ordinary skill in the art will recognize from this disclosure and the knowledge in the art that further pharmaceutically acceptable salts include those listed by Remington's Pharmaceutical Sciences, 17th ed., Mack Publishing Company, Easton, PA, p. 1418 (1985). In general, a pharmaceutically acceptable acid or base salt can be synthesized from a parent compound that contains a basic or acidic moiety by any conventional chemical method. Briefly, such salts can be prepared by reacting the free acid or base forms of these compounds with a stoichiometric amount of the appropriate base or acid in an appropriate solvent.

Ranges provided herein are understood to be shorthand for all of the values within the range. For example, a range of 1 to 50 is understood to include any number, combination of numbers, or sub-range from the group consisting of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, or 50, as well as all intervening decimal values between the aforementioned integers such as, for example, 1.1, 1.2, 1.3, 1.4, 1.5, 1.6, 1.7, 1.8, and 1.9. With respect to sub-ranges, "nested sub-ranges" that extend from either end point of the range are specifically contemplated. For example, a nested sub-range of an exemplary range of 1 to 50 may comprise 1 to 10, 1 to 20, 1 to 30, and 1 to 40 in one direction, or 50 to 40, 50 to 30, 50 to 20, and 50 to 10 in the other direction.

Certain formulas and other illustrations provided herein depict triazole reaction products resulting from azide-alkyne cycloaddition reactions. While such formulas generally depict only a single regioisomer of the resulting triazole formed in the reaction, it is intended that the formulas encompass both resulting regioisomers. Thus, while the formulas depict only a single regioisomer



it is intended that the other regioisomer



is also encompassed.

104

The term "subject" refers to an animal which is the object of treatment, observation, or experiment. By way of example only, a subject includes, but is not limited to, a mammal, including, but not limited to, a human or a non-human mammal, such as a non-human primate, bovine, equine, canine, ovine, or feline.

The term "optional" or "optionally" denotes that a subsequently described event or circumstance can but need not occur, and that the description includes instances where the event or circumstance occurs and instances in which it does not.

The term "moiety" refers to a specific segment or functional group of a molecule. Chemical moieties are often recognized chemical entities embedded in or appended to a molecule.

As used herein, the term "number average molecular weight" (M<sub>n</sub>) means the statistical average molecular weight of all the individual units in a sample, and is defined by Formula (1):

$$M_n = \frac{\sum N_i M_i}{\sum N_i} \quad \text{Formula (1)}$$

where M<sub>i</sub> is the molecular weight of a unit and N<sub>i</sub> is the number of units of that molecular weight.

As used herein, the term "weight average molecular weight" (M<sub>w</sub>) means the number defined by Formula (2):

$$M_w = \frac{\sum N_i M_i^2}{\sum N_i M_i} \quad \text{Formula (2)}$$

where M<sub>i</sub> is the molecular weight of a unit and N<sub>i</sub> is the number of units of that molecular weight.

As used herein, "peak molecular weight" (M<sub>p</sub>) means the molecular weight of the highest peak in a given analytical method (e.g., mass spectrometry, size exclusion chromatography, dynamic light scattering, analytical centrifugation, etc.).

As used herein, "conjugation handle" refers to a reactive group capable of forming a bond upon contacting a complementary reactive group. In some instances, a conjugation handle preferably does not have a substantial reactivity with other molecules which do not comprise the intended complementary reactive group. Non-limiting examples of conjugation handles, their respective complementary conjugation handles, and corresponding reaction products can be found in the table below. While table headings place certain reactive groups under the title "conjugation handle" or "complementary conjugation handle," it is intended that any reference to a conjugation handle can instead encompass the complementary conjugation handles listed in the table (e.g., a trans-cyclooctene can be a conjugation handle, in which case tetrazine would be the complementary conjugation handle). In some instances, amine conjugation handles and conjugation handles complementary to amines are less preferable for use in biological systems owing to the ubiquitous presence of amines in biological systems and the increased likelihood for off-target conjugation.

Table of Conjugation Handles

Conjugation Handle	Complementary Conjugation Handle	Reaction Product
Sulfhydryl	alpha-halo-carbonyl (e.g., bromo-acetamide), alpha-beta unsaturated carbonyl (e.g., maleimide, acrylamide)	thioether
Azide	alkyne (e.g., terminal alkyne, substituted triazole cyclooctyne (e.g., dibenzocyclooctyne (DBCO), difluorocyclooctyne, bicyclo[6.1.0]nonyne, etc.) )	
Phosphine Tetrazine	Azide/ester pair trans-cyclooctene	amide dihydro-pyridazine
Amine	Activated ester (e.g., N-hydroxy-succinimide ester, pentafluorophenyl ester)	amide
isocyanate	amine	urea
epoxide	amine	alkyl-amine
hydroxyl amine	aldehyde, ketone	oxime
hydrazide	aldehyde, ketone	hydrazone
potassium acyl trifluoroborate	O-substituted hydroxylamine (e.g., O-carbamoylhydroxylamine)	amide

Throughout the instant application, prefixes are used before the term “conjugation handle” to denote the functionality to which the conjugation handle is linked. For example, a “protein conjugation handle” is a conjugation handle attached to a protein (either directly or through a linker), an “antibody conjugation handle” is a conjugation handle attached to an antibody (either directly or through a linker), and a “linker conjugation handle” is a conjugation handle attached to a linker group (e.g., a bifunctional linker used to link a synthetic protein and an antibody).

The term “alkyl” refers to a straight or branched hydrocarbon chain radical, having from one to twenty carbon atoms, and which is attached to the rest of the molecule by a single bond. An alkyl comprising up to 10 carbon atoms is referred to as a C<sub>1</sub>-C<sub>10</sub> alkyl, likewise, for example, an alkyl comprising up to 6 carbon atoms is a C<sub>1</sub>-C<sub>6</sub> alkyl. Alkyls (and other moieties defined herein) comprising other numbers of carbon atoms are represented similarly. Alkyl groups include, but are not limited to, C<sub>1</sub>-C<sub>10</sub> alkyl, C<sub>1</sub>-C<sub>9</sub> alkyl, C<sub>1</sub>-C<sub>8</sub> alkyl, C<sub>1</sub>-C<sub>7</sub> alkyl, C<sub>1</sub>-C<sub>6</sub> alkyl, C<sub>1</sub>-C<sub>5</sub> alkyl, C<sub>1</sub>-C<sub>4</sub> alkyl, C<sub>1</sub>-C<sub>3</sub> alkyl, C<sub>1</sub>-C<sub>2</sub> alkyl, C<sub>2</sub>-C<sub>8</sub> alkyl, C<sub>3</sub>-C<sub>8</sub> alkyl and C<sub>4</sub>-C<sub>8</sub> alkyl. Representative alkyl groups include, but are not limited to, methyl, ethyl, n-propyl, 1-methyl ethyl (i-propyl), n-butyl, i-butyl, 5-butyl, n-pentyl, 1,1-dimethyl ethyl (i-butyl), 3-methylhexyl, 2-methylhexyl, 1-ethyl-propyl, and the like. In some embodiments, the alkyl is methyl or ethyl. In some embodiments, the alkyl is —CH(CH<sub>3</sub>)<sub>2</sub> or —C(CH<sub>3</sub>)<sub>3</sub>. Unless stated otherwise specifically in the specification, an alkyl group may be optionally substituted. “Alkylene” or “alkylene chain” refers to a straight or branched divalent hydrocarbon chain linking the rest of the molecule to a radical group. In some embodiments, the alkylene is —CFF—, —CH<sub>2</sub>CH<sub>2</sub>—, or —CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—. In some embodiments, the alkylene is —CH<sub>2</sub>—. In some embodiments, the alkylene is —CH<sub>2</sub>CH<sub>2</sub>—. In some embodiments, the alkylene is —CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—. Unless stated otherwise specifically in the specification, an alkylene group may be optionally substituted.

The term “alkenylene” or “alkenylene chain” refers to a straight or branched divalent hydrocarbon chain in which at least one carbon-carbon double bond is present linking the rest of the molecule to a radical group. In some embodiments, the alkenylene is —CH=CH—, —CH<sub>2</sub>CH=CH—, or —CH=CHCH<sub>2</sub>—. In some embodiments, the alkenylene

is —CH=CH—. In some embodiments, the alkenylene is —CH<sub>2</sub>CH=CH—. In some embodiments, the alkenylene is —CH=CHCH<sub>2</sub>—.

The term “alkynyl” refers to a type of alkyl group in which at least one carbon-carbon triple bond is present. In one embodiment, an alkenyl group has the formula —C≡C-R<sup>X</sup>, wherein R<sup>X</sup> refers to the remaining portions of the alkynyl group. In some embodiments, R<sup>X</sup> is H or an alkyl. In some embodiments, an alkynyl is selected from ethynyl, propynyl, butynyl, pentynyl, hexynyl, and the like. Non-limiting examples of an alkynyl group include —C≡CH, —C≡CCH<sub>3</sub>, —C≡CCH<sub>2</sub>CH, and —CH<sub>2</sub>C≡CH.

The term “aryl” refers to a radical comprising at least one aromatic ring wherein each of the atoms forming the ring is a carbon atom. Aryl groups can be optionally substituted. Examples of aryl groups include, but are not limited to phenyl, and naphthyl. In some embodiments, the aryl is phenyl. Depending on the structure, an aryl group can be a monoradical or a diradical (i.e., an arylene group). Unless stated otherwise specifically in the specification, the term “aryl” or the prefix “ar-” (such as in “aralkyl”) is meant to include aryl radicals that are optionally substituted. In some embodiments, an aryl group comprises a partially reduced cycloalkyl group defined herein (e.g., 1,2-dihydronaphthalene). In some embodiments, an aryl group comprises a fully reduced cycloalkyl group defined herein (e.g., 1,2,3,4-tetrahydronaphthalene). When aryl comprises a cycloalkyl group, the aryl is bonded to the rest of the molecule through an aromatic ring carbon atom. An aryl radical can be a monocyclic or polycyclic (e.g., bicyclic, tricyclic, or tetracyclic) ring system, which may include fused, spiro or bridged ring systems.

The term “cycloalkyl” refers to a monocyclic or polycyclic non-aromatic radical, wherein each of the atoms forming the ring (i.e. skeletal atoms) is a carbon atom. In some embodiments, cycloalkyls are saturated or partially unsaturated. In some embodiments, cycloalkyls are spirocyclic or bridged compounds. In some embodiments, cycloalkyls are fused with an aromatic ring (in which case the cycloalkyl is bonded through a non-aromatic ring carbon atom). Cycloalkyl groups include groups having from 3 to 10 ring atoms. Representative cycloalkyls include, but are not limited to, cycloalkyls having from three to ten carbon atoms, from three to eight carbon atoms, from three to six carbon atoms, or from three to five carbon atoms. Monocyclic cycloalkyl radicals include, for example, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl, and cyclooctyl. In some embodiments, the monocyclic cycloalkyl is cyclopentyl. In some embodiments, the monocyclic cycloalkyl is cyclopentenyl or cyclohexenyl. In some embodiments, the monocyclic cycloalkyl is cyclopentenyl. Polycyclic radicals include, for example, adamantyl, 1,2-dihydronaphthalenyl, 1,4-dihydronaphthalenyl, tetralinyl, decalinyl, 3,4-dihydronaphthalenyl-1(2H)-one, spiro[2.2]pentyl, norbornyl and bicyclic[1.1.1]pentyl. Unless otherwise stated specifically in the specification, a cycloalkyl group may be optionally substituted.

The term “heteroalkylene” or “heteroalkylene chain” refers to a straight or branched divalent heteroalkyl chain linking the rest of the molecule to a radical group. Unless stated otherwise specifically in the specification, the heteroalkyl or heteroalkylene group may be optionally substituted as described below. Representative heteroalkylene groups include, but are not limited to —CH<sub>2</sub>—O—CH<sub>2</sub>—, —CH<sub>2</sub>—N(alkyl)-CH<sub>2</sub>—, —CH<sub>2</sub>—N(aryl)-CH<sub>2</sub>—, —OCH<sub>2</sub>CH<sub>2</sub>O—, —OCH<sub>2</sub>CH<sub>2</sub>OCH<sub>2</sub>CH<sub>2</sub>O—, or —OCH<sub>2</sub>CH<sub>2</sub>OCH<sub>2</sub>CH<sub>2</sub>OCH<sub>2</sub>CH<sub>2</sub>O—.

The term “heterocycloalkyl” refers to a cycloalkyl group that includes at least one heteroatom selected from nitrogen, oxygen, and sulfur. Unless stated otherwise specifically in the specification, the heterocycloalkyl radical may be a monocyclic, or bicyclic ring system, which may include fused (when fused with an aryl or a heteroaryl ring, the heterocycloalkyl is bonded through a non-aromatic ring atom) or bridged ring systems. The nitrogen, carbon or sulfur atoms in the heterocycl radical may be optionally oxidized. The nitrogen atom may be optionally quaternized. The heterocycloalkyl radical is partially or fully saturated. Examples of heterocycloalkyl radicals include, but are not limited to, dioxolanyl, thienyl[1,3]dithianyl, tetrahydroquinolyl, tetrahydroisoquinolyl, decahydroquinolyl, decahydroisoquinolyl, imidazolinyl, imidazolidinyl, isothiazolidinyl, isoxazolidinyl, morpholinyl, octahydroindolyl, octahydroisoindolyl, 2-oxopiperazinyl, 2-oxopiperidinyl, 2-oxopyrrolidinyl, oxazolidinyl, piperidinyl, piperazinyl, 4-piperidonyl, pyrrolidinyl, pyrazolidinyl, quinuclidinyl, thiazolidinyl, tetrahydrofuryl, trithianyl, tetrahydropyranyl, thiomorpholinyl, thiamorpholinyl, 1-oxo-thiomorpholinyl, 1,1-dioxo-thiomorpholinyl. The term heterocycloalkyl also includes all ring forms of carbohydrates, including but not limited to monosaccharides, disaccharides and oligosaccharides. Unless otherwise noted, heterocycloalkyls have from 2 to 12 carbons in the ring. In some embodiments, heterocycloalkyls have from 2 to 10 carbons in the ring. In some embodiments, heterocycloalkyls have from 2 to 10 carbons in the ring and 1 or 2 N atoms. In some embodiments, heterocycloalkyls have from 2 to 10 carbons in the ring and 3 or 4 N atoms. In some embodiments, heterocycloalkyls have from 2 to 12 carbons, 0-2 N atoms, 0-2 O atoms, 0-2 P atoms, and 0-1 S atoms in the ring. In some embodiments, heterocycloalkyls have from 2 to 12 carbons, 1-3 N atoms, 0-1 O atoms, and 0-1 S atoms in the ring. It is understood that when referring to the number of carbon atoms in a heterocycloalkyl, the number of carbon atoms in the heterocycloalkyl is not the same as the total number of atoms (including the heteroatoms) that make up the heterocycloalkyl (i.e. skeletal atoms of the heterocycloalkyl ring). Unless stated otherwise specifically in the specification, a heterocycloalkyl group may be optionally substituted.

The term “heteroaryl” refers to an aryl group that includes one or more ring heteroatoms selected from nitrogen, oxygen, and sulfur. In some embodiments, heteroaryl is monocyclic or bicyclic. Illustrative examples of monocyclic heteroaryls include pyridinyl, imidazolyl, pyrimidinyl, pyrazolyl, triazolyl, pyrazinyl, tetrazolyl, furyl, thienyl, isoxazolyl, thiazolyl, oxazolyl, isothiazolyl, pyrrolyl, pyridazinyl, triazinyl, oxadiazolyl, thiadiazolyl, furazanyl, indolizine, indole, benzofuran, benzothiophene, indazole, benzimidazole, purine, quinolizine, quinoline, isoquinoline, cinnoline, phthalazine, quinazoline, quinoxaline, 1,8-naphthyridine, and pteridine. Illustrative examples of monocyclic heteroaryls include pyridinyl, imidazolyl, pyrimidinyl, pyrazolyl, triazolyl, pyrazinyl, tetrazolyl, furyl, thienyl, isoxazolyl, thiazolyl, oxazolyl, isothiazolyl, pyrrolyl, pyridazinyl, triazinyl, oxadiazolyl, thiadiazolyl, and furazanyl. Illustrative examples of bicyclic heteroaryls include indolizine, indole, benzofuran, benzothiophene, indazole, benzimidazole, purine, quinolizine, quinoline, isoquinoline, cinnoline, phthalazine, quinazoline, quinoxaline, 1,8-naphthyridine, and pteridine. In some embodiments, heteroaryl is pyridinyl, pyrazinyl, pyrimidinyl, thiazolyl, thienyl, thiadiazolyl or furyl. In some embodiments, a heteroaryl contains 0-6 N atoms in the ring. In some embodiments, a heteroaryl contains 1-4 N atoms in the ring. In some embodiments, a

heteroaryl contains 4-6 N atoms in the ring. In some embodiments, a heteroaryl contains 0-4 N atoms, 0-1 O atoms, 0-1 P atoms, and 0-1 S atoms in the ring. In some embodiments, a heteroaryl contains 1-4 N atoms, 0-1 O atoms, and 0-1 S atoms in the ring. In some embodiments, heteroaryl is a C<sub>1</sub>-C<sub>9</sub> heteroaryl. In some embodiments, monocyclic heteroaryl is a C<sub>1</sub>-C<sub>5</sub> heteroaryl. In some embodiments, monocyclic heteroaryl is a 5-membered or 6-membered heteroaryl. In some embodiments, a bicyclic heteroaryl is a C<sub>6</sub>-C<sub>9</sub> heteroaryl. In some embodiments, a heteroaryl group comprises a partially reduced cycloalkyl or heterocycloalkyl group defined herein (e.g., 7,8-dihydroquinoline). In some embodiments, a heteroaryl group comprises a fully reduced cycloalkyl or heterocycloalkyl group defined herein (e.g., 5,6,7, 8-tetrahydroquinoline). When heteroaryl comprises a cycloalkyl or heterocycloalkyl group, the heteroaryl is bonded to the rest of the molecule through a heteroaromatic ring carbon or hetero atom. A heteroaryl radical can be a monocyclic or polycyclic (e.g., bicyclic, tricyclic, or tetracyclic) ring system, which may include fused, spiro or bridged ring systems.

The term “optionally substituted” or “substituted” means that the referenced group is optionally substituted with one or more additional group(s) individually and independently selected from D, halogen, —CN, —NH<sub>2</sub>, —NH(alkyl), —N(alkyl)<sub>2</sub>, —OH, —CO<sub>2</sub>H, —CO<sub>2</sub>alkyl, —C(=O)NH<sub>2</sub>, —C(=O)NH(alkyl), —C(=O)N(alkyl)<sub>2</sub>, —S(=O)<sub>2</sub>NH<sub>2</sub>, —S(=O)<sub>2</sub>NH(alkyl), —S(=O)<sub>2</sub>N(alkyl)<sub>2</sub>, alkyl, cycloalkyl, fluoroalkyl, heteroalkyl, alkoxy, fluoroalkoxy, heterocycloalkyl, aryl, heteroaryl, aryloxy, alkylthio, arylthio, alkylsulfoxide, arylsulfoxide, alkylsulfone, and arylsulfone. In some other embodiments, optional substituents are independently selected from D, halogen, —CN, —NH<sub>2</sub>, —NH(CH<sub>3</sub>), —N(CH<sub>3</sub>)<sub>2</sub>, —OH, —CO<sub>2</sub>H, —CO<sub>2</sub>(C<sub>1</sub>-C<sub>4</sub>alkyl), —C(=O)NH<sub>2</sub>, —C(=O)NH(C<sub>1</sub>-C<sub>4</sub>alkyl), —C(=O)N(C<sub>1</sub>-C<sub>4</sub>alkyl)<sub>2</sub>, —S(=O)<sub>2</sub>NH<sub>2</sub>, —S(=O)<sub>2</sub>NH(C<sub>1</sub>-C<sub>4</sub>alkyl), —S(=O)<sub>2</sub>N(C<sub>1</sub>-C<sub>4</sub>alkyl)<sub>2</sub>, C<sub>1</sub>-C<sub>4</sub>alkyl, C<sub>3</sub>-C<sub>6</sub>cycloalkyl, C<sub>1</sub>-C<sub>4</sub>fluoroalkyl, C<sub>1</sub>-C<sub>4</sub>heteroalkyl, C<sub>1</sub>-C<sub>4</sub>alkoxy, C<sub>1</sub>-C<sub>4</sub>fluoroalkoxy, —SC<sub>1</sub>-C<sub>4</sub>alkyl, —S(=O)C<sub>1</sub>-C<sub>4</sub>alkyl, and —S(=O)<sub>2</sub>C<sub>1</sub>-C<sub>4</sub>alkyl. In some embodiments, optional substituents are independently selected from D, halogen, —CN, —NH<sub>2</sub>, —OH, —NH(CH<sub>3</sub>), —N(CH<sub>3</sub>)<sub>2</sub>, —NH(cyclopropyl), —CH<sub>3</sub>, —CH<sub>2</sub>CH<sub>3</sub>, —CF<sub>3</sub>, —OCH<sub>3</sub>, and —OCF<sub>3</sub>. In some embodiments, substituted groups are substituted with one or two of the preceding groups. In some embodiments, an optional substituent on an aliphatic carbon atom (acyclic or cyclic) includes oxo (=O).

As used herein, “AJICAP™ technology,” “AJICAP™ methods,” and similar terms refer to systems and methods (currently produced by Ajinomoto Bio-Pharma Services (“Ajinomoto”)) for the site specific functionalization of antibodies and related molecules using affinity peptides to deliver the desired functionalization to the desired site. General protocols for the AJICAP™ methodology are found at least in PCT Publication No. WO2018199337A1, PCT Publication No. WO2019240288A1, PCT Publication No. WO2019240287A1, PCT Publication No. WO202009079A1, Matsuda et al., *Mol. Pharmaceutics* 2021, 18, 4058-4066, and Yamada et al., AJICAP: Affinity Peptide Mediated Regiodivergent Functionalization of Native Antibodies. *Angew. Chem., Int. Ed.* 2019, 58, 5592-5597, and in particular Examples 2-4 of US Patent Publication No. US20200190165A1. In some embodiments, such methodologies site specifically incorporate the desired functionalization at lysine residues at a position selected from position 246, position 248, position 288, position 290, and position 317 of an antibody Fc region (e.g., an IgG1 Fc

## US 12,383,631 B2

**109**

region) (EU numbering). In some embodiments, the desired functionalization is incorporated at residue position 248 of an antibody Fc region (EU numbering). In some embodi-

**110**

ments, position 248 corresponds to the 18<sup>th</sup> residue in a human IgG CH<sub>2</sub> region (EU numbering). Sequences (SEQ ID NOS) of IL-18 Polypeptides

TABLE 2

SEQ ID NO:	Modification	Sequence
1	Native sequence	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
2	E6K, C38A, K53A, C68A, E85C	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKCMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
3	E6K, V11I, C38A, K53A, T63A, C68A, C76A, E85C, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKCMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
4	E6K, C38A, K53A, C68A, M86C	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKECNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
5	E6K, V11I, C38A, K53A, T63A, C68A, C76A, M86C, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKECNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
6	E6K, V11I, C38A, K53A, T63A, C68A, C76A, D98C, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSCII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
7	E6K, C38A, K53A, C68A, C76A, M86C, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSAENKI ISFKECNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
8	E6K, V11I, C38A, K53A, T63A, C68A, C76A, T95C, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
9	E6K, C38A, K53A, C68A, D98C	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSCII FFQRSVPGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
10	E6K, C38A, K53A, C68A, C76A, D98C, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSCII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
11	E6K, V11I, C38A, K53A, C76A, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
12	E6K, C38A, K53A, T63A, C76A, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
13	E6K, K53A, T63N	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVNISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
14	E6K, K53A, S50A, T63N	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVNISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSVPGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED

TABLE 2-continued

SEQ		Sequence
ID NO:	Modification	
15	E6K, K53A, S50H, T63N	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVNISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
16	E6K, K53A, T63N, S65A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVNIAVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
17	E6K, K53A, S50H	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
18	E6K, C38A, K53A, C68A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
19	E6K, K53A, K79A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISFKEMNPPD NIKDTKSII FFQRSVPFGHD ISTLSCENAI NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
20	E6K, K53A, R104A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENAI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
21	E6K, K53A, G108A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFAHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
22	E6K, K53A, H109A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGAD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
23	E6K, K53A, K112A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NAMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
24	E6K, C38A, K53A, T63A, C76A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
25	E6K, C38Q, K53A, T63A, C76A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
26	E6K, C38A, K53A, T63A, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
27	E6K, C38Q, K53A, T63A, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
28	E6K, C38A, K53A, T63A, C76A, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
29	E6K, V11I, C38A, K53A, T63A	YFGKLKSKLS ISFKEMNPPD FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSCENKI IIRNLNDQVL NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED

TABLE 2-continued

SEQ ID NO:	Modification	Sequence
30	E6K, V11I, C38A, K53A, T63A, C76A, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
31	C38A, C76A, C127A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYKDSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
32	C38A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYKDSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
33	E6K, C38A, K53A, T63A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
34	E06K, K53A, S55A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
35	Y01G, F02A, E06K, M51G, K53A, D54A, S55A, T63A	GAGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS GYAAAQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
36	K53A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
37	S55A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDAQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
38	E06K	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
39	E06K, K53A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
40	E06K, S55A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDAQPRGM AVTISVKCEK ISFKEMNPPD ISTLSCENKI NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
41	K53A, S55A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
42	E06K, K53A, S55A, T63A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
43	E06K, K53A, S55A, Y01G	GFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM ISTLSCENKI AVTISVKCEK ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
44	E06K, K53A, S55A, F02A	YAGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED

TABLE 2-continued

SEQ ID NO:	Modification	Sequence
45	E06K, K53A, S55A, D54A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYAAQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
46	E06K, K53A, S55A, M51G	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS GYADAQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
47	C38S, C68S, C76S, C127S	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYKDSQPRGM AVTISVKSEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
48	C38S, C68S, C76S, C127S, K70C	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYKDSQPRGM ISTLSENKI AVTISVKSEC ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
49	E06K, K53A, S55A, C38S, C68S, C76S, C127S, K70C	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADAQPRGM AVTISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
50	E06K, K53A, T63A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
51	T63A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
52	E06K, T63A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
53	K53A, T63A	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
54	E06K, K53A, C38S, C68S, C76S, C127S, K70C	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM AVTISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
55	K53A, T63A, C38S, C68S, C76S, C127S, K70C	YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM AVAISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
56	E6K, K53A, C38S, C76S, C127S	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
57	E6K, C38S, K53A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
58	E6K, K53A, C38S, C68S, C76S, C127S, K70C	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM AVTISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED
59	E6K, C38A, K53A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED

TABLE 2-continued

SEQ ID NO:	Modification	Sequence
60	E6K, C38Q, K53A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
61	E6K, C38A, K53A, C76A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
62	E6K, C38A, K53A, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
63	E6K, C38A, K53A, C76A, C127A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
64	E6K, K53A, C38A, S55A, T63A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADAQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
65	E6K, C38Q, K53A, S55A, T63A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADAQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
66	E6K, K53A, K84A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFAEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
67	E6K, K53A, D98A	YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED
68	V11I, C38A, (a.k.a. M51G, K53A, C146) C76A, C127A	YFGKLESKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS GYADSQPRGM AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
69	E6K, V11I, (a.k.a. C38A, M51G, C183) K53A, T63A, C76A, C127A	YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS GYADSQPRGM AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPFGHD NKMQFESSSY EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED
70	N-terminal G, (a.k.a. E6K, V11I, C192) C38A, M51G, K53A, T63A, C76A, C127A	GYFGKLKSKL SIIIRNLNDQV LFIDQGNRPL FEDMTDSRAR DNAPRTIFII SGYADSQPRG MAVAISVKCE KISTLSAENK IISFKEMNPP DN1KDTKSII IFFQRSVPFGH DNKMQFESSSS YEGYFLAAEK ERDLFKLILK KEDELGDRSI MFTVQNED
71	N-terminal G, (a.k.a. E6K, V11I, C141) C38A, K53A, T63A, C76A, C127A	GYFGKLKSKL SIIIRNLNDQV LFIDQGNRPL FEDMTDSAR DNAPRTIFII SMYADSQPRG MAVAISVKCE KISTLSAENK IISFKEMNPP DN1KDTKSII IFFQRSVPFGH DNKMQFESSSS YEGYFLAAEK ERDLFKLILK KEDELGDRSI MFTVQNED
72	N-terminal (a.k.a. 4xG. E6K, C140) V11I, C38A, K53A, T63A, C76A, C127A	GGGGYFGKLK SKLSIIRNLN DQVLFIDQGN RPLFEDMTDS DARDNAPRTI FIISMYADSQ PRGMAVAISVKCEKISTLSA ENK1ISFKEM NPPDNIKDTK SDIIFFQRSV PGHDNKMQFE SSSYEGYFLA AEKERDLFKL ILKKEDELGD RSIMPTVQNE D

## Additional Exemplary II-18 Constructs

Also provided herein are IL-18 polypeptides which comprise the modifications to SEQ ID NO: 1 listed in the table below, each of which is assigned a Composition ID, which can be incorporated into an immunocytokine composition as provided herein. In some embodiments, the IL-18 polypep-

tide of an immunocytokine composition comprises the set of amino acid substitutions shown for any one of the constructs depicted below. In the constructs depicted below, each of the substitutions is listed using SEQ TD NO: 1 as a reference sequence. In some embodiments, the IL-18 polypeptide an immunocytokine composition comprises only the substitu-

119

tions shown for a construct below relative to SEQ TD NO: 1 (i.e., the IL-18 polypeptide has only the indicated set of substitutions and the remaining residues are those set forth in SEQ ID NO: 1).

120

## Soluble His-SUMO-IL18 Variants

*E. coli* BL21 (DE3) harboring a plasmid encoding a N-His-SUMO tagged IL-18 variant fusion is inoculated into 3 L LB culture medium and induced with 0.4 mM IPTG at

TABLE 3

Additional IL-18 Polypeptide					
	Composition ID/ Substitutions to SEQ ID NO: 1	Composition ID/ Substitutions to SEQ ID NO: 1	Composition ID/ Substitutions to SEQ ID NO: 1		
C143	V11I, C38A, K53A, C76A, C127A	C156 V11I, C38A, N41A, C168 K53A, C76A, C127A	V11I, C38A, C76A, S105K, C127A		
C144	V11I, C38A, K53A, T63A, C76A, C127A	C157 V11I, C38A, K53A, C174 C76A, C127A, D132A	K8L, E6K, V11I, C38A, K53A, T63A, C76A, C127A		
C145	V11I, C38A, K53A, S55A, C76A, C127A	C158 V11I, C38A, K53A, C175 C76A, G108A, C127A	E6K, V11I, C38A, I49E, K53A, T63A, C76A, C127A		
C147	V11I, C38A, K53A, D54A, C76A, C127A	C159 V11I, C38A, K53A, C176 C76A, H109A, C127A	E6K, V11I, C38A, I49M, K53A, T63A, C76A, C127A		
C148	F2A, V11I, C38A, K53A, C76A, C127A	C160 V11I, C38A, K53A, C177 C76A, D110A, C127A	E6K, V11I, C38A, I49R, K53A, T63A, C76A, C127A		
C149	V11I, E31A, C38A, K53A, C76A, C127A	C161 K8R, V11I, C38A, C178 C76A, Q103E, C127A	E6K, V11I, C38A, K53A, T63A, C76A, Q103R, C127A		
C150	V11I, T34A, C38A, K53A, C76A, C127A	C162 K8E, V11I, C38A, C179 C76A, Q103R, C127A	E6K, K8E, V11I, C38A, K53A, T63A, C76A, Q103R, C127A		
C151	V11I, D35A, C38A, K53A, C76A, C127A	C163 V11I, C38A, C76A, C180 Q103K, C127A	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153R		
C152	V11I, S36A, C38A, K53A, C76A, C127A	C164 V11I, C38A, S55H, C181 C76A, C127A	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153E		
C153	V11I, D37A, C38A, K53A, C76A, C127A	C165 V11I, C38A, S55R, C182 C76A, C127A	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153Y		
C154	V11I, E31A, D37A, C38A, K53A, C76A, C127A	C166 V11I, C38A, S55T, C184 C76A, C127A	E6R, V11I, C38A, K53A, T63A, C76A, C127A		
C155	V11I, C38A, D40A, K53A, C76A, C127A	C167 V11I, C38A, C76A, C142 S105I, C127A	Y1M, E6K, V11I, C38A, K53A, T63A, C76A, C127A		

45

Although the present disclosure and its advantages have been described in detail, it should be understood that various changes, substitutions and alterations can be made herein without departing from the spirit and scope of the disclosure as defined in the appended claims.

The present disclosure is further illustrated in the following Examples which are given for illustration purposes only and are not intended to limit the disclosure in any way.

## EXAMPLES

## Example 1A—Recombinant IL18 Expression and Purification

Recombinant TL-18 variants suitable for linking to an antibody or antigen binding fragment as provided herein can be prepared according to the protocols provided below. In some instances, the recombinant IL-18 will contain a cysteine residue at the desired point of attachment of the linker, or may include an unnatural amino acid (e.g., azidolysine) suitable for attachment of the linker at the desired point of attachment.

30° C. for 6 h. Cells are pelleted and cell lysis is done by sonication in lysis buffer: PBS, pH 7.4. Soluble protein is purified via Ni-NTA beads 6FF (wash 1 with: PBS, 20 mM imidazole, pH7.4; wash 2 with PBS, 50 mM Imidazole, pH7.4; elution with PBS, 500 mM imidazole, pH7.4).

50 Fractions containing the protein are pooled, dialyzed into PBS pH 7.4 and followed by SUMO digestion. Then the protein is two-step purified with Ni-NTA beads (continue with flow through sample) and gel filtration. Fractions 55 containing the protein are pooled and QC is performed using analytical techniques, such as SDS-PAGE and analytical SEC.

## Insoluble his-SUMO-IL18 Variants

56 *E. coli* BL21 (DE3) harboring a plasmid encoding a N-His-SUMO tagged IL-18 variant fusion are inoculated into 10 L LB culture medium and induced with 0.4 mM IPTG at 30° C. for 6 h. Cells are pelleted and cell lysis is done by sonication in lysis buffer: PBS, 8 M urea, pH 7.4. Protein is purified via Ni-NTA beads 6FF (wash 1 with: PBS, 8 M urea, 20 mM imidazole, pH7.4; wash 2 with PBS, 8 M urea, 50 mM Imidazole, pH7.4; elution with PBS, 8 M urea, 500 mM imidazole, pH7.4).

121

Fractions containing the protein are pooled, dialyzed into PBS pH 7.4 and followed by SUMO digestion. Then the protein is purified with Ni-NTA beads (equilibrate column with PBS, 8 M urea, pH 7.4, wash with PBS, 8 M urea, pH 7.4, elution with PBS, 8 M urea, pH 7.4). Fractions containing the protein are pooled, dialyzed into PBS pH 7.4 and QC is performed using analytical techniques, such as SDS-PAGE and analytical SEC.

#### Insoluble Tagless IL18 Variants

*E. coli* BL21 (DE3) harboring a plasmid encoding mIL-18 is inoculated into 2 L LB culture medium and induced with 0.4 mM IPTG at 30° C. for 6 h. Cells are pelleted and cell lysis was done by sonication in lysis buffer: 110 mM Tris, 1.1 M guanidine HCl, 5 mM DTT, pH 8.9. Protein is purified via Q Sepharose FF (balance buffer 20 mM MES, pH 7.0, elution with an increasing gradient from 0 to 1 M NaCl).

#### Example 1B—Additional Methods for Recombinant IL18 Expression and Purification

The following protocols were also used to prepare certain IL-18 polypeptides provided herein which were subsequently used either in assays for conversion into immunocytokine compositions as provided herein.

#### Expression of IL-18 Polypeptides

IL-18 polypeptides were produced as an N-terminal fusion to N-His-SUMO-IL18. The gene was synthesized and cloned by a commercial vendor. Plasmids were transformed into *E. coli* BL21 (DE3). Expression was performed in shake flasks with TB medium. The cells were grown at 37° C. until an OD<sub>600</sub> of approximately 1.2 was reached, after which they were induced by 0.1 mM IPTG and cultured for another 20 hours at 18° C. Cells were harvested by centrifugation.

#### Purification of IL18 Polypeptides

Cell lysis—Cells were resuspended in lysis buffer (20 mM Tris/HCl, pH 8.0, 0.15 M NaCl, 10 mM Imidazole, 1 tablet of EDTA-free complete protease inhibitor (Roche, COEDTAF-RO) per liter production) at 100 mL buffer/L culture and disrupted twice with a homogenizer at 1000 bar. The lysate was cleared of debris by centrifugation at 40'000 g for 2×45 minutes, changing flask in between, and subsequent filtration through a 0.22 µm filter.

Affinity Purification and Endotoxin Removal—The lysate was loaded on Ni NTA resin (Cytiva, 17524802) pre-equilibrated with 20 mM Tris/HCl, pH 8.0, 0.15 M NaCl, 10 mM Imidazole, at 5 mL/min and washed with the same buffer for 5 CV. To remove endotoxins, the column was washed with 20 mM Tris/HCl, pH 8.0, 0.15 M NaCl, 10 mM Imidazole, 0.1% Tryton X-114 at 10 mL/min for 30 CV. The column was washed with 20 mM Tris/HCl, pH 8.0, 0.15 M NaCl, 10 mM Imidazole, for 5 CV at 5 mL/min and the protein of interest eluted by linear increase of imidazole concentration. The column was then regenerated by 0.5M NaOH.

SUMO digestion and dialysis—To cleave the SUMO tag, SUMO protease was added to the elution pool at a w/w ratio of 1:250 (protein:SUMO enzyme) and incubated for 18 hours at 4° C. At the same time, the protein was dialysed (20 mM Tris, pH 8.0, 150 mM NaCl), to reduce the imidazole concentration.

Purification by reverse IMAC—In order to remove the cleaved tag and the SUMO protease, the digested protein

122

was flown through a Ni NTA resin column pre-equilibrated with 20 mM Tris/HCl, pH 8.0, 0.15 M NaCl, 10 mM Imidazole, at 5 mL/min. The flow-through was collected.

Buffer Exchange—The flow-through was concentrated to 2.6 mg/mL and buffer exchanged into either 20 mM HEPES, 150 mM NaCl, 0.5 mM TCEP, 10% glycerol, pH 7.5 or PBS, 10% glycerol, pH 7.4. Proteins were stored at -70° C. until further quality controls.

#### Example 2—Conjugation of IL-18 Polypeptide with Bifunctional Linking Group

An IL-18 polypeptide as provided herein can conjugated to a bifunctional linking group prior to forming the full linker of the immunocytokine composition. In some cases, the bifunctional linking group first attaches to a desired residue of the IL-18 polypeptide at the point of attachment of the linker. Once attached to the IL-18 polypeptide, the second functionality of the bifunctional linking group is used to attach to a second portion. An exemplary schematic of such a process is shown in FIG. 6. The process shown in FIG. 6 is not needed for all IL-18 polypeptides, as some IL-18 polypeptides provided herein (e.g., an IL-18 with the desired conjugation handle for final linkage with the antibody already attached). An exemplary protocol on an IL-18 polypeptide with a cysteine residue point of attachment provided herein is described below.

Conjugation—The IL-18 polypeptide is stored at a concentration of 2.4 mg/mL at -80° C. in potassium phosphate buffer (pH 7.0) containing 50 mM KCl and 1 mM DTT. The sample is thawed on ice yielding a clear solution. The protein solution is diluted in PBS, pH 7.4. A clear solution is obtained at a concentration of ~0.4 mg/mL.

The protein solution is dialyzed against PBS, pH 7.4 (twice against 600 mL for 2 h and once against 800 mL for 18 h). After dialysis, a clear solution is obtained with no sign of precipitation. Protein concentration is obtained using UV absorbance at 280 nm and by BCA protein assay.

A stock solution of bi-functional linking group (e.g., bromoacetylamo-PEG5-azide, CAS: 1415800-37-1) in water is prepared at a concentration of 20 mM. 500 µL of the protein solution are mixed with 25 µL of linking group solution. pH was adjusted to 7.5 and it was let to react for 3 h at 20° C.

The progress of the synthesis is monitored by reverse-phase HPLC using a gradient of 5 to 30% (2.5 min) and 30 to 75% (7.5 min) CH<sub>3</sub>CN with 0.1% TFA (v/v) on a *Aeris* 50 WIDEPOR<sub>E</sub> C18 200 Å column (3.6 m, 150×4.6 mm) at a flow rate of 1 mL/min at 40° C. and by MALDI-TOF MS.

Purification—In some cases, ion-exchange chromatography is used to purify the conjugated protein. To remove the excess of probe, the reaction mixture (volume is around 500 µL) is flowed through a Hi-Trap-G-FF-1 mL column using 25 mM Tris (pH 7.4) as the buffer. The column is eluted with a linear gradient of 0-0.35 M NaCl in the same buffer. The fractions containing the target protein are gathered, buffer exchanged (25 mM Tris, pH 7.4, 75 mM NaCl, 5% glycerol) and concentrated at 0.4 mg/mL. The concentration of purified protein is determined by UV absorbance at 280 nm and by BCA protein assay. The protein solution is kept at -80° C.

Characterization—The purity and identity of the recombinant protein from commercial source and the conjugated protein is confirmed by aSEC, HPLC and MALDI-TOF MS.

123

Example 3—Conjugation of IL-18 Polypeptide to Antibody

Preparation of a Conjugatable Antibody

A modified antibody (e.g., an anti-PD-1 antibody such as nivolumab or Lzm-009) comprising a DBCO conjugation handle is prepared using a protocol modified from Examples 2-4 of US Patent Publication No. US2020019165A1. An exemplary illustration of this process resulting in the attachment of one DBCO conjugation handle is shown in FIG. 3. Briefly, the CD20 antibody with a free sulphydryl group attached to a lysine residue side chain in the Fc region is prepared by reacting the antibody with an affinity peptide configured to deliver a protected version of the sulphydryl group (e.g., a thioester) to the lysine residue. The protecting group is then removed to reveal the free sulphydryl. The free sulphydryl is then reacted with a bifunctional reagent comprising a bromoacetamide group connected to the DBCO conjugation handle through a linking group (e.g., bromoacetamido-dPEG<sup>®</sup><sub>4</sub>-amido-DBCO). The method can be used to produce an antibody with one DBCO group present (DAR1) and/or two DBCO groups attached to the antibody (DAR2, one DBCO group linked to each Fc of the antibody). Conjugation of Antibody to IL-18 Polypeptide

The DBCO modified antibody is then conjugated to a IL-18 polypeptide comprising an azide moiety at a desired point of attachment (e.g., an IL-18 polypeptide which contains an amino acid with an azide side chain or an IL-18 linked to an azide using a bifunctional linking group as in Example 4). DBCO modified antibody with one (DAR1) or two (DAR2) reactive handles are reacted with 2-10 equivalents of azide containing IL-18 (pH 5.2 buffer, 5% trehalose, rt, 24 h). In an alternative embodiment, antibody comprising two DBCO conjugation handles is reacted either as an excess reagent (e.g., 5-10 equivalents) with 1 equivalent of IL-18 comprising an azide functionality to produce a DAR1 antibody or the antibody comprising two DBCO conjugation handles is reacted with 1 equivalent of antibody with excess reagent of IL-18 comprising an azide (e.g., 5-10 equivalents) to produce a DAR2 antibody. An illustration of this protocol is shown in FIG. 7.

Purification and Characterization of Antibody-IL-18 Immunocytokine

The resulting immunocytokine is purified by cation-exchange chromatography and/or size exclusion chromatography to obtain purified immunocytokine. Antibody-IL-18 polypeptide immunocytokine is purified from unreacted IL-18 and aggregates using a desalting column, CIEX and SEC (GE Healthcare Life Sciences AKTA pure, mobile phase: Histidine 5.2/150 mM NaCl/5% Trehalose, column: GE Healthcare Life Sciences SUPERDEX™ 200 increase 3.2/300, flow rate: 0.5 mL/min).

The purity and identity of the antibody-IL-18 polypeptide immunocytokine is confirmed by RP-HPLC (HPLC: ThermoFisher Scientific UHPLC Ultimate 3000, column: Waters BEH C-4 300A, 3.0 μm, 4.6 mm, 250 mm, mobile phase A: 0.05% TFA in Water, mobile phase B: 0.05% TFA in mixture of ACN:IPA:ETOH:H<sub>2</sub>O (5:1.5:2:1.5), flow rate: 0.5 mL/min, injection amount: μg (10 μL Injection of 1 mg/mL), gradient: 0% to 20% mobile phase B in 50 min) and SDS-PAGE.

Exemplary chromatograms and analytical characterization of DAR1 immunocytokines of an IL-18 of SEQ ID NO: 60 conjugated via residue C68 and an anti-PD-1 antibody (Lzm-009) prepared according to the described methods are shown in FIGS. 8-10.

124

Additional exemplary chromatograms and analytical characterization of a DAR1 immunocytokine of an IL-18 of SEQ ID NO: 30 conjugated via residue C68 to Fc residue K248 of the anti-PD-1 antibody Lzm-009 are shown in FIG. 11 (RP HPLC), FIG. 12 (SEC-HPLC), and FIG. 13 (Q-TOF Mass Spectrometry). The immunocytokine shown is Composition A.

Example 4—Characterization of Immunocytokine IL-18 Activity

The ability of the immunocytokine to perform various IL-18 activities is measured as provided below, as well as relevant comparisons to non-conjugated IL-18 polypeptides. Surface Plasmon Resonance

The interaction of immunocytokines, wild type IL-18, and of modified IL-18 polypeptides with human IL-18 receptor subunits are measured with Surface Plasmon Resonance (SPR) technology. Anti-human IgG antibodies are bound by amine coupling onto a CM5 chip to capture 6 g/mL of Fc fused human IL-18R $\alpha$ , 6 g/mL of Fc fused human IL-18R $\beta$ , or 2 g/mL of Fc fused human IL-18BP isoform a (IL-18BPa) for 30 min before capture. In other settings, 6 g/mL of alpha and beta IL-18 receptors are mixed and pre-incubated for 30 min before capture of the alpha/beta heterodimer IL-18 receptor.

The kinetic binding of the IL-18 analytes and immunocytokines are measured with a Biacore 8K instrument in two-fold serial dilutions starting at 1 μM down to 0.98 nM. Regeneration of the surface back to amine coupled anti IgG antibody is done after every concentration of analyte. To measure the protein association to the receptors, the samples are injected with a flow rate of 50 L/min for 60 s, followed by 300 s buffer only to detect the dissociation. The used running buffer is 1×PBS with 0.05% Tween20. The relative response units (RU, Y-axis) are plotted against time (s, X-axis) and analyzed in a kinetic 1:1 binding model for the monomer receptor binding and for the binding to the IL-18BP. A kinetic heterogenous ligand fit model is applied for the alpha/beta heterodimer binding.

IL-18BP Binding alphaLISA Assay

A human IL-18BP AlphaLISA Assay Kit is used to determine the binding affinity of each immunocytokine and IL-18 variant for IL-18BP, which detected the presence of free form IL-18BP.

Sixteen three-fold serial dilutions of IL-18 analytes are prepared in aMEM medium supplemented with 20% FCS, Glutamax, and 25 μM β-mercaptoethanol in the presence of 5 ng/mL of His-tagged human IL-18BP. Final IL-18 analytes concentration range from 2778 nM to 0.2 pM.

After 1 hr incubation at room temperature, free IL-18BP levels are measured using a Human IFN $\gamma$  AlphaLISA Assay Kit. In a 384 well OPTIplate, 5 μL of 5x Anti-IL-18BP acceptor beads are added to 7.5 μL of an IL-18/IL-18BP mix. After 30 min incubation at room temperature with shaking, 5 μL of biotinylated Anti-IL-18BP antibodies are added to each well. The plate is incubated further for 1 hr at room temperature. Under subdued light, 12.5 μL of 2x streptavidin (SA) donor beads are pipetted into each well, and the wells are incubated with shaking for an additional 30 min at room temperature. The AlphaLISA signal is then measured on an Enspire plate reader with 680 and 615 nm as excitation and emission wavelengths, respectively. The dissociation constant ( $K_D$ ) is calculated based on a variable slope, four parameter analysis using GraphPad PRISM software.

125

IFN $\gamma$  Induction Cellular Assay

The ability of immunocytokines and IL-18 polypeptides provided herein are assessed for ability to induce IFN $\gamma$  in a cellular assay according to the protocol below.

The NK cell line NK-92 derived from a patient with lymphoma (ATCC® CRL-2407™) is cultured in aMEM medium supplemented with 20% FCS, Glutamax, 25  $\mu$ M  $\beta$ -mercaptoethanol, and 100 IU/mL of recombinant human IL-2.

On the day of experiment, cells are harvested and washed with aMEM medium without IL-2 and containing 1 ng/mL of recombinant human IL-12. After counting, cells are seeded at 100,000 cells/well in a 384 well titer plate and incubated at 37° C./5% CO<sub>2</sub>. Sixteen 4-fold serial dilutions of IL-18 analytes are prepared in aMEM medium, and 1 ng/mL of IL-12 were added to the NK-92 cells. Final IL-18 analyte concentrations range from 56 nM to 5 $\times$ 10<sup>-5</sup> pM.

After incubating the cells for 16-20 hr at 37° C./5% CO<sub>2</sub>, 5  $\mu$ L of supernatant is carefully transferred to a 384 microwell OptiPlate. IFN $\gamma$  levels are measured using a human IFN $\gamma$  AlphaLISA Assay Kit. Briefly, 10  $\mu$ L of 2.5 $\times$  AlphaLISA Anti-IFN $\gamma$  acceptor beads and biotinylated antibody anti-IFN $\gamma$  mix are added to the 5  $\mu$ L of NK-92 supernatants. The mixtures are incubated for 1 hr at room temperature with shaking. Under subdued light, 2.5  $\mu$ L of 2 $\times$  streptavidin (SA) donor beads are pipetted into each well, and the wells are incubated for 30 min at room temperature with shaking. AlphaLISA signals are then measured on an EnSpire™ plate reader using 680 nm and 615 nm as excitation and emission wavelengths, respectively. Half maximal effective concentrations (EC<sub>50</sub>) are calculated based on a variable slope and four parameter analysis using GraphPad PRISM software.

## IL-18 Binding Protein Inhibition Cellular Assay

The NK cell line NK-92 derived from a patient with lymphoma (ATCC® CRL-2407™) is cultured in aMEM medium supplemented with 20% FCS-Glutamax, 25  $\mu$ M  $\beta$ -mercaptoethanol, and 100 IU/mL of recombinant human IL-2.

On the day of experiment, cells are harvested and washed with aMEM medium without IL-2 and containing 1 ng/mL of recombinant human IL-12. After counting, the cells are seeded at 100,000 cells/well in a 384 well titer plate and incubated at 37° C./5% CO<sub>2</sub>. Sixteen 2-fold serial dilutions of Fc-fused human IL-18 binding protein isoform a (IL-18BPa) are prepared in aMEM medium. 1 ng/mL of IL-12 containing 2 nM of each modified IL-18 polypeptide variant is added to the NK-92 cells. The final IL-18 analyte concentration is 1 nM, and the final IL-18BPa concentration ranged from 566 nM to 17 pM.

After incubating the cells for 16-20 hr at 37° C./5% CO<sub>2</sub>, 5  $\mu$ L of the supernatant is carefully transferred to a 384 microwell OptiPlate. IFN $\gamma$  levels are measured using a human IFN $\gamma$  AlphaLISA Assay Kit. Briefly, 10  $\mu$ L of 2.5 $\times$  AlphaLISA anti-IFN $\gamma$  acceptor beads and biotinylated antibody anti-IFN $\gamma$  mix are added to 5  $\mu$ L of NK-92 supernatants. The mixtures are incubated for 1 hr at room temperature with shaking. Under subdued light, 2.5  $\mu$ L of 2 $\times$ SA donor beads are pipetted in each well and incubated for 30 min at room temperature with shaking. AlphaLISA signals are then measured on an EnSpire™ plate reader using 680 nm and 615 nm as excitation and emission wavelengths, respectively. Half maximal inhibitory concentrations (IC<sub>50</sub>) are calculated based on a variable slope and four parameter analysis using GraphPad PRISM software.

126

IFN $\gamma$  Induction on Primary Human Cells

Ability of IL-18 variants to stimulate Human peripheral blood mononuclear cells (PBMCs) was assessed according to the following protocol.

Isolation of lymphocytes: Blood from Buffy Coats of healthy volunteers was diluted with equal volume of PBS and slowly poured on top of SepMate tube prefilled with 15 mL Histopaque-1077. Tubes were centrifuged for 10 minutes at 1200 g, the top layer was collected and washed 3 times with PBS containing 2% of Fetal Bovine Serum. PBMCs were counted and cryopreserved as aliquots of 20 $\times$ 10<sup>6</sup> cells.

Cryopreserved PBMCs were thawed and seeded at 150 000 cells/well in a 96w round bottom 96 well plate. PBMCs were stimulated with a gradient of human IL-18 variants ranging from 0.2 pg/mL to 3600 ng/mL. All stimulations were performed in the presence of hIL-12 (1 ng/ml, Sino Biological, #CT011-H08H) for 24 hrs in RPMI containing 10% Fetal Bovine Serum.

Cytokine production after 24 hr stimulation were measured using Legendplex bead-based cytokine assay (Biologen #740930) according to manufacturer protocol. Half maximal effective concentrations (EC<sub>50</sub>) of IFNg released in culture supernatant were calculated based on a variable slope and four parameter analysis using GraphPad PRISM software.

IFN $\gamma$  Induction on Primary Mouse Cells

Ability of IL-18 variants to stimulate murine splenocytes was assessed according to the following protocol.

Cryopreserved splenocytes isolated from BALB/c and C57BL6 mice were purchased from IQ Biosciences (Berkeley, CA, USA).

Cryopreserved splenocytes were thawed, treated with DNaseI, and seeded at 200 000 cells/well in a 96w round bottom 96 well plate. Splenocytes were stimulated with a gradient of human IL-18 variants ranging from 0.2 pg/mL to 3600 ng/mL. All stimulations were performed in the presence of mIL-12 (1 ng/ml, Peprotech, cat #210-12) for 24 hrs in RPMI containing 10% Fetal Bovine Serum.

Cytokine production after 24 hr stimulation were measured using Legendplex bead-based cytokine assay (Biologen #740622) according to manufacturer protocol. Half maximal effective concentrations (EC<sub>50</sub>) of IFNg released in culture supernatant were calculated based on a variable slope and four parameter analysis using GraphPad PRISM software.

## Example 5—Immune Cell Associated Antigen Binding ELISA Assay (FIG. 14A, FIG. 14B, FIG. 15A, and FIG. 15B)

The interaction of the unmodified antibodies and corresponding IL-18 immunocytokines with relevant immune cell associated antigen are measured by ELISA assay. For these studies, Corning high-binding half-area plates (Fisher Scientific, Reinach, Switzerland) are coated overnight at 4° C. with 25  $\mu$ L of unmodified antibodies corresponding IL-18 immunocytokines at 5  $\mu$ g/ml in PBS. Plates are then washed four times with 100  $\mu$ L of PBS-0.02% Tween20. Plate surfaces are blocked with 25  $\mu$ L of PBS-0.02% Tween20-1% BSA at 37° C. during 1 h. Plates are then washed four times with 100  $\mu$ L of PBS-0.02% Tween20. Twenty-five microliters (25  $\mu$ L) of recombinant biotinylated human PD-1 (Biotinylated Recombinant Human PD-1/CD279-Fc Chimera, carrier-free, Biologen #789406) or PD-L1 (Biotinylated Human PD-L1/B7-H1, ACROBiosystems, PD1-H82E5-25UG) protein are added in seven-fold serial dilutions

## 127

starting at 12 nM down to 0.15 pM into PBS-0.02% Tween20-0.1% BSA and incubated at 37° C. during 2 h. Plates are then washed four times with 100 µl of PBS-0.02% Tween20. Twenty-five microliters of Streptavidin-Horseradish peroxidase (#RABHRP3, Merck, Buchs, Switzerland) diluted at 1:500 into PBS-0.02% Tween20-0.1% BSA are added to each well and incubated at Room Temperature during 30 min. Plates are then washed four times with 100 µl of PBS-0.02% Tween20. Fifty microliters of TMB substrate reagent (#CL07, Merck, Buchs, Switzerland) are added to each well and incubated at 37° C. during 5 min. After 5 min at 37° C., Horseradish peroxidase reaction is stopped by adding 50 µl/well of 0.5M H<sub>2</sub>SO<sub>4</sub> stop solution. ELISA signal is then measured at 450 nm on an ENSPTRE® plate reader from Perkin Elmer (Schwerzenbach, Switzerland). Results from this experiment are shown in the table below.

KD Values of the Interaction of Immunocytokines with PD-1 and PD-L1 as Measured by ELISA

Composition	Antibody	IL-18 polypeptide	PD-1 KD (pM)	PD-L1 KD (pM)
—	Pembrolizumab/Keytruda	—	>100000	
—	LZM-009	—	40.7 >100000	
—	Nivolumab/Opdivo	—	91.9 >100000	
—	Durvalumab/Imfinzi	—	>100000	146
—	Atezolizumab/Tecentriq	—	>100000	443
—	Avelumab/Bavencio	—	>100000	36
A	LZM-009	SEQ ID NO: 30	46.7 >100000	
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	>100000	473

FIG. 14A and FIG. 14B show plots measuring ability of the unmodified and of conjugated anti-PD1 antibodies to

## 128

mination of kinetics and affinity of a ligands to its receptor. Here anti-human IgG FC Capture (AHC) sensors are loaded with the test items (ICs). Sensors are first dipped into a kinetic buffer for baseline measurement, then into an analyte solution, here human PD1, to allow association and again into a buffer solution where the analyte is allowed to come off the ligand (dissociation). Several concentrations of analyte are run in parallel and enable the calculation of affinity parameters: Ka, Kd, KD.

Typically, first, the sensors are regenerated by 3 cycles of dipping into 10 mM glycine solution at pH=2 for 20 seconds, followed by 20 second kinetics buffer and a final 60 seconds in kinetics buffer to establish the initial signal (baseline). Second, the loading column will contain the ligand, here the unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody, at a fixed concentration determined in the loading scout experiment (20 ug/mL). Then another wash/baseline step allows non immobilized proteins to be washed away. The association column will contain the 2-fold dilution series of the analyte (His-tagged human PD1, R&D #8986-PD) including a no analyte control. The highest concentration should be ~10-fold the KD. The dissociation designates the sensors to return to previous baseline column with kinetics buffer. After acquisition, the data is analyzed with Data Analysis Studio software (Sartorius). Data sets are first preprocessed by subtracting references samples and aligning curves on the baseline step. Group fitting is then applied to the data series and kinetics parameters are calculated. Results from this experiment are shown in the table below.

Binding Kinetics of the Interaction of Reference Antibodies and Immunocytokines with PD-1 as Measured by Bio-Layer Interferometry (BLI)

Composition	Antibody	IL-18 polypeptide	ka (1/Ms)	k <sub>d</sub> (1/s)	KD (nM)
—	Pembrolizumab/Keytruda	—	4.00E+05	2.72E-03	6.77
—	LZM-009	—	3.42E+05	7.70E-03	23.50
A	LZM-009	SEQ ID NO: 4.54E+05	4.79E-03	11.50	

bind with human PD1/CD279 ligand, with the figure showing ELISA signal on the y-axis and dosage of the biotinylated PD-1 protein on the x-axis. The unconjugated reference antibodies are Pembrolizumab, LZM-009, Nivolumab, Atezolizumab, Durvalumab, and Avelumab. The conjugated antibodies tested in this figure are compositions A and composition B.

FIG. 15A and FIG. 15B show plots measuring ability of the unmodified and of conjugated antibodies to bind with human PD-L1/B7-H1 ligand, with the figure showing ELISA signal on the y-axis and dosage of the biotinylated PD-L1 protein on the x-axis. The unconjugated reference antibodies are Pembrolizumab, LZM-009, Nivolumab, Atezolizumab, Durvalumab, and Avelumab. The conjugated antibodies tested in this figure are compositions A and composition B.

Example 6—Kinetic Analysis of Binding of Reference Antibodies and Immunocytokines to Immune Cell Associated Antigens (FIG. 16)

Based on Bio-Layer Interferometry (BLI), Octet® BLI systems enable real-time, label-free analysis for the deter-

FIG. 16 shows plots measuring ability of the unmodified and of conjugated antibodies to bind to human PD-L1/B7-H1 ligand, with the figure showing net BioLayer interferometry shift in nanometer on the y-axis and time of incubation dosage of the biotinylated PD-L1 protein on the x-axis. The figure shows mean ELISA signal on the y-axis and dosage of the human Fc gamma receptors on the x-axis. The unconjugated reference antibodies are Pembrolizumab and LZM-009. The conjugated antibodies tested is Compositions A.

Example 7—PD-1/PD-L1 Blockade Assay (FIGS. 17A and 17B)

For immunocytokine compositions which comprise PD-1 or PD-L1 antibodies or antigen binding fragments, the experiment outlined below is performed to assess the ability of the immunocytokines to interfere with the PD-1/PD-L1 pathway. The assay is the PD-1/PD-L1 Blockade Bioassay from Promega (Cat #J1250, Madison, WI, USA). PD-1/PD-L1 Blockade Bioassay is a bioluminescent cell-based assay based on the co-culture of effector cells with target cells mimicking an immunological synapse. Jurkat T cells

**129**

expressing human PD-1 and a luciferase reporter driven by a NFAT response element (NFAT-RE) are activated by CHO-K1 cells expressing human PD-L1 and an engineered cell surface protein designed to activate Jurkat cells cognate TCRs. Concurrent interaction PD-1/PD-L1 inhibits TCR signaling and represses NFAT-RE-mediated luminescence. Addition of either an anti-PD-1 or anti-PD-L1 antibody that blocks the PD-1/PD-L1 interaction releases the inhibitory signal, restoring TCR activation and resulting in a gain of signal of NFAT-RE luminescent reporter.

Briefly, PD-L1 aAPC/CHO-K1 Target cells were plated in white tissue culture 96-wells plates and cultured overnight at 37° C./5% CO<sub>2</sub>. Test molecules were measured in four-fold serial dilutions starting at 1 μM down to 0.002 nM and pre-incubated on target cells for 10 min before the addition of freshly thawed PD-1 Jurkat effector cells. After 6 h at 37° C./5% CO<sub>2</sub>, activity NFAT-RE luminescent reporter was evaluated by the addition of Bio-Glo reagent and measured on an ENSPTRE® plate reader (1 sec/well) from Perkin Elmer (Schwerzenbach, Switzerland). Results from this experiment are shown in the table below.

**Activity of Unconjugated IL-18 Variants and Corresponding IL-18 Immunocytokines in the PD-1/PD-L1 Blockade Cellular Assay**

Composition	Antibody	IL-18 polypeptide	KD FcRn (nM)
—	Pembrolizumab/Keytruda	—	0.757
—	LZM-009	—	5.799
—	Durvalumab/Imfinzi	—	0.330
—	Atezolizumab/Tecentriq	—	0.664
—	Avelumab/Bavencio	—	0.676
—	Trastuzumab/Herceptin	>10000	
A	LZM-009	SEQ ID NO: 30	4.075
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	0.573
C	Trastuzumab/Herceptin	SEQ ID NO: 30	NT

FIGS. 17A and 17B shows plots measuring ability of the unmodified and of conjugated anti-PD1 antibodies to interfere with PD1/PDL1 pathway, with the figure showing normalized luminescence intensity of effector cells NFAT-Lucia reporter on the y-axis and dosage of the unmodified and of conjugated anti-PD1 antibodies on the x-axis. The unconjugated reference antibodies are Pembrolizumab, LZM-009, Nivolumab, Atezolizumab, Durvalumab,

and Avelumab. The conjugated antibodies tested in this figure are compositions A and composition B.

**Example 8—Human FcγR Binding Assay (FIG. 18A and FIG. 18B)**

The interaction of the unmodified and of conjugated antibodies with human Fc gamma receptors I (FcγRI/CD64), with human Fc gamma receptors IIa (FcγRIIa/CD32a), with inhibitory human Fc gamma receptors IIb (FcγRIIb/CD32b), and with human Fc gamma receptors III FcγRIIIa/CD16 were measured by ELISA.

Briefly, Corning high-binding half-area plates (Fisher Scientific, Reinach, Switzerland) were coated overnight at 4° C. with 25 μl of unmodified and of conjugated anti-PD1 antibodies at 2.5 μg/ml in PBS. Plates were then washed four times with 100 μl of PBS-0.02% Tween20. Plates surfaces were blocked with 25 μl of PBS-0.02% Tween20-1% BSA at 37° C. during 1 h. Plates were then washed four times with 100 μl of PBS-0.02% Tween20. Then twenty-five microliters of either recombinant Human Fc gamma RI/CD64 Protein (R&D systems, 1257-FC-050, CF), recombinant Human Fc gamma RIIA/CD32a (H167) Protein (R&D systems, 9595-CD-050, CF), recombinant Human Fc gamma RIIB/CD32b Avi-tag Protein (R&D systems, AVI1875-050, CF), or recombinant Human Fc gamma RIIIA/CD16a Protein (R&D systems, 4325-FC-050; CF) were added in five-fold serial dilutions ranging from 1000 nM to 0.001 nM into PBS-0.02% Tween20-0.1% BSA and incubated at 37° C. during 2 h. Plates were then washed four times with 100 μl of PBS-0.02% Tween20. Twenty-five microliters of a 1/500 HRP-anti-His antibody in PBS—0.02% Tween20-0.1% BSA (R&D systems, anti-HIS—HRP Ab, #MAB050H) were added to each well and plates were incubated at Room Temperature during 1 h. Plates were then washed four times with 100 μl of PBS-0.02% Tween20. Fifty microliters of TMB substrate reagent (#CL07, Merck, Buchs, Switzerland) were added to each well and incubated at 37° C. during 5 min. After 5 min at 37° C., Horseradish peroxidase reaction was stopped by adding 50 μl/well of 0.5M H<sub>2</sub>SO<sub>4</sub> stop solution. ELISA signal was then measured at 450 nm on an EnSpire plate reader from Perkin Elmer (Schwerzenbach, Switzerland). Results from this experiment are shown in the table below.

**Binding Affinity of Reference Antibodies and Immunocytokines with Human Fc Gamma Receptors as Measured by ELISA**

Composition	Antibody	IL-18 polypeptide	FcγRI/	FcγRIIa	FcγRIIIa	FcγRIIb
			(CD64) nM	(CD32a) nM	(CD16) nM	(CD32b) nM
—	Pembrolizumab/Keytruda	—	0.5932	3358	1660	>10000
—	LZM-009	—	0.3150	1348	1707	2627
—	Durvalumab/Imfinzi	—	2.34	160	234	233
—	Atezolizumab/Tecentriq	—	9.17	1370	356	>10000
—	Trastuzumab/Herceptin	—	0.0785	350	807	360
A	LZM-009	SEQ ID NO: 30	0.25	1892	598	436
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	10.85	1740	492	>10000
C	Trastuzumab/Herceptin	SEQ ID NO: 30			3299	

NT: Not Tested

## 131

FIG. 18A and FIG. 18B show plots measuring ability of the unmodified and of conjugated antibodies to bind to human Fe gamma receptor I (CD64) on top panels, and to human Fe gamma receptor IIIa (CD16) on lower panels. The figure shows mean ELISA signal on the y-axis and dosage of the human Fe gamma receptors on the x-axis. The unconjugated reference antibodies are LZM-009 and Atezolizumab. The conjugated antibodies tested are Compositions A and B.

## Example 9—Human FcRn Binding Assay (FIG. 19)

The interaction of the unmodified and of conjugated anti-PD1 antibodies with the human neonatal Fc receptor (FcRn) at pH 6 was measured using the AlphaLISA® Human FcRn Binding Kit (AL3095C) from Perkin Elmer (Schwerzenbach, Switzerland). The AlphaLISA® detection of FcRn and IgG binding uses IgG coated AlphaLISA® acceptor beads to interact with biotinylated human FcRn captured on Streptavidin-coated donor beads. When reference IgG binds to FcRn, donor and acceptor beads come into proximity enabling the transfer of singlet oxygen that trigger a cascade of energy transfer reactions in the acceptor beads, resulting in a sharp peak of light emission at 615 nm. Addition of a free IgG antibodies into the AlphaLISA® mixture creates a competition for the binding of FcRn to the reference antibody resulting in a loss of signal.

Briefly, test molecules were measured in serial dilutions starting at 5 uM down to 64 pM and incubated with AlphaLISA® reaction mixture consisting of 800 nM of recombinant biotinylated human FcRn, 40 µg/ml of human IgG conjugated Acceptor beads, and 40 µg/ml of Streptavidin coated Donor beads in pH 6 MES buffer. After 90 min at 23° C. in the dark, AlphaLISA® signal was measured on an EnSpire plate reader (Excitation at 680 nm, Emission at 615 nm) from Perkin Elmer (Schwerzenbach, Switzerland). Results from this experiment are shown in the table below.

## Binding Affinity of Reference Antibodies and Immunocytokines with the Human Neonatal Fc Receptor as Measured by AlphaLISA

Composition	Antibody	IL-18 polypeptide	KD FcRn (nM)
—	Pembrolizumab/Keytruda	—	7.45
—	LZM-009	—	6.00
—	Durvalumab/Imfinzi	—	8.31
—	Atezolizumab/Tecentriq	—	6.40
—	Avelumab/Bavencio	—	5.36
—	Trastuzumab/Herceptin	—	6.55
A	LZM-009	SEQ ID NO: 30	33.45
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	36.68
C	Trastuzumab/Herceptin	SEQ ID NO: 30	25.12

FIG. 19 shows plots measuring ability of the unmodified and of conjugate antibodies to bind to human Fe neonatal receptor. The figure shows mean AlphaLISA signal on the y-axis and dosage of the human Fe neonatal receptor (FcRn) on the x-axis. The unconjugated reference antibodies are LZM-009 and Atezolizumab. The conjugated antibodies tested are Compositions A and B.

## Example 10—IFN Gamma Secretion Assay in NK92 Cells (FIGS. 20 &amp; 21)

The IFNgamma secretion stimulating activity of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines was evaluated on NK92 cell line. The NIK cell line

## 132

NK-92 derived from a patient with lymphoma (ATCC, Cat #CRL-2407) was cultured in aMIEM medium supplemented with 12.5% FCS, 12.5% horse serum (HS), 50 uM B-mercaptoEthanol, and 2 ng/ml of recombinant Human Interleukin-2 (IL-2).

On the day of experiment, cells were harvested and washed with aNMEM medium without IL-2 and resuspended in medium (w/o IL-2) containing 1 ng/ml of recombinant human Interleukin-12 (Sinopharmaceuticals, Cat #CT011-H08H). After counting, cells were seeded at 100 000 cells/well in a 384 well titer plate and incubated at 37° C./5% CO<sub>2</sub>. Sixteen 4-fold serial dilutions of IL-18 analytes were prepared in aMIEM medium-1 ng/ml IL-12 and were added to the NK-92 cells. Final IL-18 analytes concentration ranged from to 200 nM down to 0.01 pM.

After 16-20 h incubation at 37° C./5% CO<sub>2</sub>, 5 µl of supernatant were carefully transferred to a 384 microwells OPTIPlate (Perkin Elmer; Cat #6007270) and Interferon-gamma (IFNgamma) levels measured using the Human IFNgamma AlphaLISA Assay Kit (Perkin Elmer, Cat #AL217C). Briefly, 10 µl of 2.5x AlphaLISA Anti-IFNgamma acceptor beads and biotinylated Antibody Anti-IFNgamma mix were added to the 5 µl of NK-92 supernatants and incubated for 1 h at room temperature under shaking. Under subdued light, 2.5 µl of 2x streptavidin (SA) donor beads were pipetted in each well and incubated for 30 min at room temperature under shaking. AlphaLISA signal was then measured on an Enspire plate reader (Perkin Elmer) using 680 and 615 nm as excitation and emission wavelengths respectively. Half maximal effective concentration (EC50) was calculated based on a variable slope, four parameter analysis using GraphPad PRISM software. Results from this experiment are shown in the table below.

## Activity of Unconjugated IL-18 Variants and Corresponding IL-18 Immunocytokines in the IFNgamma Secretion NK92 Assay

Composition	Antibody	IL-18 polypeptide	EC <sub>50</sub> (nM)
—	—	SEQ ID NO: 1	0.374
—	—	SEQ ID NO: 30	0.002
—	—	SEQ ID NO: 31	0.103
—	—	SEQ ID NO: 60	0.008
A	LZM-009	SEQ ID NO: 30	0.073
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	0.001
C	Trastuzumab/Herceptin	SEQ ID NO: 30	0.027
D	LZM-009	SEQ ID NO: 31	0.471
E	LZM-009	SEQ ID NO: 60	0.056
F	Trastuzumab/Herceptin	SEQ ID NO: 60	0.064

FIG. 20 shows plots measuring the levels PD-1 and PD-L1 surface expression on NK92 cells

FIG. 21 shows plots measuring ability of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines to stimulate the secretion of IFNgamma by NK92 cells. The figure shows mean IFNgamma alphaLISA signal on the y-axis and dosage of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO:1), SEQ ID NO: 30, and SEQ ID NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

## Example 11—IFN Gamma Secretion Assay in KG-1 Cells (FIG. 22-24)

The IL-18 responsive AML cell line KG-1 shows high expression of IL-18Rα and moderate levels of IL18Rβ,

## 133

respectively. The KG-1 cell line was used to generate a PD-1 expressing cell line and furthermore, to measure IFN $\gamma$  release upon incubation with TL-18 variants and corresponding TL-18 immunocytokines.

PD-1 expressing cell line generation: Briefly, KG-1 cells were transduced using lentiviral particles carrying the human PD-1 gene (PDCD1 NM\_005018; Origene, CAT #: RC210364L3V) at a MOI (Multiplicity of Infection) of 30. Spinfection was performed at 1260 g during 90 min at 37° C. in the presence of 5  $\mu$ g/ml of Polybrene and 10 mM of HEPES in complete culture media (RPMI, 10% FBS, 1% L-Glutamine). Five days after transduction, puromycin at a final concentration of 1  $\mu$ g/ml was added to select for PD-1 positive cells. For culture maintenance, puromycin concentration was decreased to 0.5  $\mu$ g/ml. Stable and homogenous expression of PD-1 was verified by surface staining (BD Pharmingen, #557860).

IFN $\gamma$  release was assessed in PD-1 positive (transduced) KG-1 cells, as well as in the parental PD-1 negative cells.  $0.5 \times 10^5$  cells were seeded into a 96-well U-bottom plate in culture media (RPMI, 10% FBS, 100 L-Glutamine) and stimulated with IL-18 variants/ICs for 20-24 h. The test items were diluted to 100 nM in culture medium, followed by 7 10-fold serial dilutions. The lowest concentration assessed was 0.05 fM. After incubation, IFN $\gamma$  release was measured using the LEGENDplex custom human mix and match KIT (Biologen LEGENDplex™ Human IFN- $\gamma$  Capture Bead B5, 13×#740942, LEGENDplex™ P 1 Essential Immune Response Panel Detection Abs, #740931, LEGENDplex™ Buffer Set A #740368). To this end, cell culture supernatant was collected and diluted 1:1 with Assay Buffer. Fluorescence measurements were done with a Quanteon Flow Cytometer from Acea Biosciences. For analysis, MFI values (median fluorescence intensity) were exported and plotted against concentrations used. The EC<sub>50</sub> values (half maximal effective concentration) were calculated based on a variable slope and four parameter analysis using GraphPad PRISM software version 9. Results from this experiment are shown in the table below.

Stimulation of IFN $\gamma$  Secretion by Unconjugated IL-18 Variants and Corresponding IL-18 Immunocytokines in Parental and PD-1 Transduced KG-1 Cells

Composition	Antibody	IL-18 polypeptide	Parental KG-1 EC <sub>50</sub> (nM)	PD-1 positive KG-1 EC <sub>50</sub> (nM)	PD-1 <sup>pos</sup> /PD-1 <sup>neg</sup> ratio
—	—	SEQ ID NO: 1	0.387	0.297	1.298
—	—	SEQ ID NO: 30	0.012	0.008	1.336
—	—	SEQ ID NO: 31	0.1485	0.1354	1.2
A	LZM-009	SEQ ID NO: 30	0.0110	0.0012	10.9
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	0.0016	0.0036	0.6
C	Trastuzumab/Herceptin	SEQ ID NO: 30	0.0166	0.0096	1.7
D	LZM-009	SEQ ID NO: 31	0.2033	0.0225	9.3

FIG. 22 shows plots measuring the levels PD-1 and PD-L1 surface expression on KG-1 cells

FIG. 23 shows plots measuring the ability of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines to stimulate the secretion of IFN $\gamma$  by parental

## 134

PD-1<sup>negative</sup> and by engineered PD-1<sup>positive</sup> KG-1 cells. The figure shows mean IFN $\gamma$  legendplex signal on the y-axis and dosage of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO:1), SEQ TD NO: 30, and SEQ TD NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

Example 12—IL-18 Binding Protein alphaLISA Assay (FIG. 24)

Wild type or modified IL-18 polypeptides samples were diluted at 5.6  $\mu$ M in a solution of 1× alphaLISA Immunoassay Buffer provided in the alphaLISA IFN $\gamma$  Detection kit and were diluted applying 3-fold serial dilutions down to 1.7 pM in 384 deep well plates. A solution of 10 ng/ml of human IL-18BP-His was prepared with 1× alphaLISA Immunoassay Buffer. IL-18/IL-18BP complex formation was performed incubating 30  $\mu$ l of IL-18BP solution to IL-18 sample titrations for 1 h at 20° C. IL-18BP standard was prediluted from stock solution supplied in alphaLISA IFN $\gamma$  Detection kit at 100 ng/ml with 1× alphaLISA Buffer and titration prepared from applying 2-fold serial dilutions. The following solutions were prepared: a 50  $\mu$ g/ml solution of anti-IL-18BP alphaLISA Acceptor beads, a 5 nM solution of biotinylated anti-IL18BP antibody and a 80  $\mu$ g/ml light-protected solution of Streptavidin Donor beads in 1× alphaLISA Immunoassay Buffer. To detect unbound IL-18BP in IL-18/IL-18P complex samples, 5  $\mu$ l of pre-mixed Acceptor beads solution was transferred on top on 7.5  $\mu$ l of samples in 384-well Optiplates, followed by a short centrifugation step at 150 g, and incubated for 30 minutes at 20° C. under shaking at 750 rpm.

5  $\mu$ l of Biotinylated anti-IL-18BP antibody were added, followed by a short centrifugation step at 150 g, and incubated for 60 minutes at 20° C. under shaking at 750 rpm. Under subdued light, 12.5  $\mu$ l of pre-mixed Donor beads were added, followed by a short centrifugation step at 150 g, and incubated for 30 minutes at 20° C. under shaking at 750 rpm with no light. AlphaLISA signal was then measured on an Enspire plate reader (Perkin Elmer) using 680 and 615 nm as excitation and emission wavelengths respectively. Unbound IL-18BP concentration interpolated from the standard signal-concentration curve using GraphPad Prism. Results from this experiment are shown in the table below. Binding Affinity of Reference Antibodies and Immunocytokines with the Human IL-18 Binding Protein as Measured by AlphaLISA

Composition	Antibody	IL-18 polypeptide	IL-18BP KD (nM)
—	—	SEQ ID NO: 1	0.411
—	—	SEQ ID NO: 30	24.600
—	—	SEQ ID NO: 31	0.103
A	LZM-009	SEQ ID NO: 30	10.930
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	6.382
C	Trastuzumab/Herceptin	SEQ ID NO: 30	10.496
D	LZM-009	SEQ ID NO: 31	15.263

FIG. 24 shows plots measuring the ability of the unconjugated IL-18 variants and corresponding IL-18 immunocytokines to bind to the human IL-18 Binding Protein (IL-18BP). The figure shows mean free IL-18BP AlphaLISA signal on the y-axis and dosage of the unconjugated IL-18

135

variants and corresponding IL-18 immunocytokines on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ ID NO: 1), SEQ ID NO: 30, and SEQ ID NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

**Example 13—Cellular IL-18 Binding protein resistance assay (FIG. 25)**

The NK cell line NK-92 derived from a patient with lymphoma (ATCC, Cat #CRL-2407) was cultured in aMEM medium supplemented with 20% FCS-Glutamax, 25  $\mu$ M B-mercaptoEthanol, and 100 IU/ml of recombinant Human Interleukin-2 (TL-2). On the day of experiment, cells were harvested and washed with aMEM medium without IL-2. After counting, cells were seeded at 100 000 cells/well in a 384 well titer plate and incubated at 37° C./5% CO<sub>2</sub>. Sixteen 2-fold serial dilutions of Fc fused human IL-18 binding protein isoform a (IL-18BP<sub>a</sub>; R&D systems, Cat #119-BP) were prepared in aMEM medium-1 ng/ml IL-12 containing 2 nM of each IL-18 variants and were added to the NK-92 cells. Final IL-18 analytes concentration was 1 nM and final IL-18BP<sub>a</sub> concentrations ranged from 566 nM down to 17 pM.

After 16-20 h incubation at 37° C./5% CO<sub>2</sub>, 5  $\mu$ l of supernatant were carefully transferred to a 384 microwells OPTIplate (Perkin Elmer, Cat #6007270) and Interferon-gamma (IFN $\gamma$ ) levels measured using the Human IFN $\gamma$  AlphaLISA Assay Kit (Perkin Elmer, Cat #AL217C). Briefly, 10  $\mu$ l of 2.5 $\times$  AlphaLISA Anti-IFN $\gamma$  acceptor beads and biotinylated Antibody Anti-IFN $\gamma$  mix were added to the 5  $\mu$ l of NK-92 supernatants and incubated for 1 h at room temperature under shaking. Under subdued light, 2.5  $\mu$ l of 2 $\times$  streptavidin (SA) donor beads were pipetted in each well and incubated for 30 min at room temperature under shaking. AlphaLISA signal was then measured on an Enspire plate reader (Perkin Elmer) using 680 and 615 nm as excitation and emission wavelengths respectively. Half maximal inhibitory concentration (IC<sub>50</sub>) was calculated based on a variable slope, four parameter analysis using GraphPad PRISM software. Results from this experiment are shown in the table below.

**IL-18BP-Mediated Inhibition of IFN $\gamma$  Secretion by NK92 Cells Stimulated with Unconjugated IL-18 Variants and Corresponding IL-18 Immunocytokines**

Composition	Antibody	IL-18 polypeptide	IC <sub>50</sub> (nM)
—	—	SEQ ID NO: 1	0.781
—	—	SEQ ID NO: 30	487.8
—	—	SEQ ID NO: 31	0.282
A	LZM-009	SEQ ID NO: 30	6.87
B	Atezolizumab/Tecentriq	SEQ ID NO: 30	>1000
C	Trastuzumab/Herceptin	SEQ ID NO: 30	21.44
D	LZM-009	SEQ ID NO: 31	0.098

FIG. 25 shows plots measuring the ability of the human IL-18 Binding Protein to inhibit the secretion of IFN $\gamma$  by NK92 cells stimulated with 2 nM of unconjugated IL-18 variants and corresponding IL-18 immunocytokines. The figure shows mean IFN $\gamma$  alphaLISA signal on the y-axis and dosage of the human IL-18 Binding Protein on the x-axis. The unconjugated IL-18 variants are native IL-18 wild-type (SEQ TD NO: 1), SEQ TD NO: 30, and SEQ TD NO: 31. Corresponding IL-18 immunocytokines tested are Compositions A, B, C, and D.

136

**Example 14—In Vivo Antitumor Activity in MC38 Colon Carcinoma Model (FIG. 26-28)**

An in vivo efficacy study was performed in mice. Naïve, 5-8 weeks old, C57BL/6-hPD1 female mice (GemPharmatech Co, Ltd, Nanjing, China) were inoculated subcutaneously at the right upper flank with MC38 tumor cells ( $3 \times 10^5$ ) in 0.1 mL of PBS for tumor development. The animals were randomized (using an Excel-based randomization software 10) performing stratified randomization based upon tumor volumes), and treatment started when the average tumor volume reached approximately 120 mm<sup>3</sup>. Animals treated with unmodified antibodies received two weekly 10 mL/kg bolus intraperitoneal (i.p.) injections. Animals treated with modified IL-18 polypeptide conjugated antibodies received two weekly 10 mL/kg bolus intravenous (i.v.) injections. After inoculation, the animals were checked daily for morbidity and mortality. At the time, animals were checked for effects 15 on tumor growth and normal behavior such as mobility, food and water consumption, body weight gain/loss (body weights were measured twice weekly), eye/hair matting and any other abnormal effect. The major endpoints were delayed tumor growth or complete tumor regression. Tumor sizes were measured three times a week in two dimensions 20 using a caliper, and the volume was expressed in mm<sup>3</sup> using the formula:  $V=0.5 ab^2$  where a and b are the long and short diameters of the tumor, respectively. Death and observed 25 clinical signs were recorded on the basis of the numbers of animals within each subset.

FIG. 26A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The 30 immunocytokine tested in this figure is Composition A tested as a single agent at 0.3 and 1 mg/kg as two weekly i.v. injections. (n=9; mean $\pm$ SEM).

FIG. 26B shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated 40 PD-L1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition B tested as a single agent at 1 and 3 mg/kg as two weekly i.v. injections. (n=9; mean $\pm$ SEM).

FIG. 27A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the body weight of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition A tested as a single agent at 0.3 and 1 mg/kg as two weekly i.v. injections. (n=9; mean $\pm$ SEM).

FIG. 27B shows a plot describing the effect of unmodified 55 PD-L1 antibodies and of IL-18 polypeptide conjugated PD-L1 antibody on the body weight of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure is Composition B tested as a single agent at 1 and 3 mg/kg as two weekly i.v. injections. (n=9; mean $\pm$ SEM).

**Example 15—In Vivo Antitumor Activity in MC38 Colon Carcinoma Model (FIG. 28-31)**

An in vivo efficacy study was performed in mice. Naïve, 6-8 weeks old, C57BL/6-hPD1 female mice (GemPharmatech

137

ech Co, Ltd, Nanjing, China) were inoculated subcutaneously at the right upper flank with MC38 tumor cells ( $3 \times 10^5$ ) in 0.1 mL of PBS for tumor development. The animals were randomized (using an Excel-based randomization software performing stratified randomization based upon tumor volumes), and treatment started when the average tumor volume reached approximately  $110 \text{ mm}^3$ . Animals treated with unmodified antibodies received two weekly 10 mL/kg bolus intraperitoneal (i.p.) injections. Animals treated with modified IL-18 polypeptide conjugated antibodies received two weekly 10 mL/kg bolus intravenous (i.v.) injections. After inoculation, the animals were checked daily for morbidity and mortality. At the time, animals were checked for effects on tumor growth and normal behavior such as mobility, food and water consumption, body weight gain/loss (body weights were measured twice weekly), eye/hair matting and any other abnormal effect. The major endpoints were delayed tumor growth or complete tumor regression. Tumor sizes were measured three times a week in two dimensions using a caliper, and the volume was expressed in  $\text{mm}^3$  using the formula:  $V=0.5 ab^2$  where a and b are the long and short diameters of the tumor, respectively. Death and observed clinical signs were recorded on the basis of the numbers of animals within each subset.

FIG. 28A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with Lzm-009 anti-PD-1 antibody at 1 mg/kg (n=9; mean $\pm$ SEM).

FIG. 28B shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows the mean tumor volume on day 17 post treatment initiation on the y-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with Lzm-009 anti-PD-1 antibody at 10 mg/kg (n=9; One-way Anova test \*\*\*P-value<0.001, \*\*P-value<0.01, \*P-value<0.1, ns not significant, TGI: Tumor Growth Inhibition).

FIG. 28C shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of MC38 syngeneic colon carcinoma tumors in hPD1 C57BL/6 mice. The figure shows the tumor volume of each individual animal on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with Lzm-009 anti-PD-1 antibody at 1 mg/kg (n=9; CR: Complete Response).

FIG. 29 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the on the body weight of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.1, 0.25, and

138

0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with Lzm-009 anti-PD-1 antibody at 1 mg/kg (n=9; mean $\pm$ SEM).

FIG. 30 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the survival of MC38 syngeneic colon carcinoma tumor-bearing hPD1 C57BL/6 mice. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.1, 0.25, and 0.5 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 0.5 mg/kg as a single agent and in combination with Lzm-009 anti-PD-1 antibody at 1 mg/kg (n=9; mean $\pm$ SEM; CR: Complete response).

#### Example 16—In Vivo Antitumor Activity in B16F10 Melanoma Model (FIG. 31-34)

An in vivo efficacy study was performed in mice. Naïve, 6-8 weeks old, C57BL/6-hPD1 female mice (GemPharmatech Co, Ltd, Nanjing, China) were inoculated subcutaneously at the right upper flank with B16F10 tumor cells ( $5 \times 10^4$ ; 1:1 with Matrigel®) in 0.1 mL of PBS for tumor development. The animals were randomized (using an Excel-based randomization software performing stratified randomization based upon tumor volumes), and treatment started when the average tumor volume reached approximately  $70 \text{ mm}^3$ . Animals treated with unmodified antibodies received two weekly 10 mL/kg bolus intraperitoneal (i.p.) injections. Animals treated with modified IL-18 polypeptide conjugated antibodies received two weekly 10 mL/kg bolus intravenous (i.v.) injections. After inoculation, the animals were checked daily for morbidity and mortality. At the time, animals were checked for effects on tumor growth and normal behavior such as mobility, food and water consumption, body weight gain/loss (body weights were measured twice weekly), eye/hair matting and any other abnormal effect. The major endpoints were delayed tumor growth or complete tumor regression. Tumor sizes were measured three times a week in two dimensions using a caliper, and the volume was expressed in  $\text{mm}^3$  using the formula:  $V=0.5 ab^2$  where a and b are the long and short diameters of the tumor, respectively. Death and observed clinical signs were recorded on the basis of the numbers of animals within each subset.

FIG. 31A shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of B16F10 syngeneic melanoma tumors in hPD1 C57BL/6 mice. The figure shows mean tumor volume on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with Lzm-009 anti-PD-1 antibody at 10 mg/kg (n=9; mean $\pm$ SEM).

FIG. 31B shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of B16F10 syngeneic melanoma tumors in hPD1 C57BL/6 mice. The figure shows the mean tumor volume on day 10 post treatment initiation on the y-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as

a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; TGI: Tumor Growth Inhibition).

FIG. 31C shows a plot describing the effect of unmodified PD-1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the growth of B16F10 syngeneic melanoma tumors in hPD1 C57BL/6 mice. The figure shows the tumor volume of each individual animal on the y-axis and time on the x-axis. The immunocytokine tested in this figure are composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; CR: Complete Response).

FIG. 32 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the body weight of B16F10 syngeneic melanoma tumor-bearing hPD1 C57BL/6 mice. The figure shows mean body weight change on the y-axis and time on the x-axis. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; mean±SEM).

FIG. 33 shows a plot describing the effect of unmodified PD-L1 antibodies and of IL-18 polypeptide conjugated PD-1 antibody on the survival of B16F10 syngeneic melanoma tumor-bearing hPD1 C57BL/6 mice. The immunocytokine tested in this figure are Composition A tested as a single agent at 0.3, 1, and 3 mg/kg as two weekly i.v. injections. As a control, Her2-targeted immunocytokine composition C was applied at 3 mg/kg as a single agent and in combination with LZM-009 anti-PD-1 antibody at 10 mg/kg (n=9; mean±SEM; CR: Complete response).

#### Example 17: Characteristics of Additional IL-18 Polypeptides

17A—HEK-Blue Reporter Assay—An IL-18R positive HEK-Blue reporter cell line is used to determine binding of IL-18 variants to IL-18R and subsequent downstream signaling. The general protocol is outlined below.

5×104 cells HEK-Blue IL18R reporter cells (InvivoGen, #hkb-hmi118) are seeded into each well of a 96 well plate and stimulated with 0-100 nM of IL-18 polypeptide variants at 37° C. and 5% CO2. After 20 h incubation, 20 µL of cell culture supernatant is then taken from each well and mixed with 180 µL QUANTI-Blue media in a 96 well plate, incubated for 1 hour at 37° C. and 5% CO2. The absorbance signal at 620 nm is then measured on an Enspire plate reader with 680 and 615 nm as excitation and emission wavelengths, respectively. Half Maximal Effective dose (EC50) is calculated based on a variable slope, four parameter analysis using GraphPad PRISM software.

The HEK-Blue IL-18R reporter assay described above was performed on additional IL-18 polypeptides which can be incorporated into immunocytokine compositions provided herein. It is expected that the IL-18 polypeptides provided below would behave similarly to C086 (SEQ ID NO: 30) when incorporated into an immunocytokine composition as those otherwise provided herein.

SEQ ID NO: or Compo- sition ID	Sequence modifications	EC <sub>50</sub> (pM)
1	Native sequence	3.33
34	E6K, K53A, S55A	272.5
39	E6K, K53A	0.72
42	E6K, K53A, S55A, T63A	0.79
50	E6K, K53A, T63A	1.77
54	E6K, C38S, K53A, C68S, K70C, C76S, C127S	9.12
56	E6K, K53A, C38S, C76S, C127S	3.73
57	E6K, C38S, K53A	0.86
30	E6K, V11I, C38A, K53A, T63A, C76A, C127A	0.034
62	E6K, C38A, K53A, C127A	0.17
60	E6K, C38Q, K53A	0.203
59	E6K, C38A, K53A	0.268
57	E6K, C38S, K53A	0.53
C143	V11I, C38A, K53A, C76A, C127A	0.98
C144	V11I, C38A, K53A, T63A, C76A, C127A	0.17
C145	V11I, C38A, K53A, S55A, C76A, C127A	3.63
C146	V11I, C38A, M51G, K53A, C76A, C127A	0.8
C147	V11I, C38A, K53A, D54A, C76A, C127A	1
C148	F2A, V11I, C38A, K53A, C76A, C127A	7.28
C149	V11I, E31A, C38A, K53A, C76A, C127A	6.6
C150	V11I, T34A, C38A, K53A, C76A, C127A	0.7
C151	V11I, D35A, C38A, K53A, C76A, C127A	13.12
C152	V11I, S36A, C38A, K53A, C76A, C127A	0.25
C153	V11I, D37A, C38A, K53A, C76A, C127A	14.12
C154	V11I, E31A, D37A, C38A, K53A, C76A, C127A	11.95
C155	V11I, C38A, D40A, K53A, C76A, C127A	0.52
C156	V11I, C38A, N41A, K53A, C76A, C127A	11.7
C157	V11I, C38A, K53A, C76A, C127A, D132A	1.95
C158	V11I, C38A, K53A, C76A, G108A, C127A	15.56
C159	V11I, C38A, K53A, C76A, H109A, C127A	19.5
C160	V11I, C38A, K53A, C76A, D110A, C127A	2.02
C161	K8R, V11I, C38A, C76A, Q103E, C127A	2.01
C162	K8E, V11I, C38A, C76A, Q103R, C127A	2.3
C163	V11I, C38A, C76A, Q103K, C127A	1.5
C164	V11I, C38A, S55H, C76A, C127A	3.14
C165	V11I, C38A, S55R, C76A, C127A	1.91
C166	V11I, C38A, S55T, C76A, C127A	4.73
C167	V11I, C38A, C76A, S105I, C127A	5.37
C168	V11I, C38A, C76A, S105K, C127A	7.73
C174	K8L, E6K, V11I, C38A, K53A, T63A, C76A, C127A	0.29
C176	E6K, V11I, C38A, I49M, K53A, T63A, C76A, C127A	0.07
C177	E6K, V11I, C38A, I49R, K53A, T63A, C76A, C127A	0.04
C178	E6K, V11I, C38A, K53A, T63A, C76A, Q103R, C127A	0.26
C179	E6K, K8E, V11I, C38A, K53A, T63A, C76A, Q103R, C127A	0.4
C181	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153E	0.1
C182	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153Y	0.08
C183	E6K, V11I, C38A, M51G, K53A, T63A, C76A, C127A	0.1
C184	E6R, V11I, C38A, K53A, T63A, C76A, C127A	0.04
C140	E6K, V11I, C38A, K53A, T63A, C76A, C127A	2.5
C141	E6K, V11I, C38A, K53A, T63A, C76A, C127A	1.68
C142	Y1M, E6K, V11I, C38A, K53A, T63A, C76A, C127A	0.02
C192	E6K, V11I, C38A, M51G, K53A, T63A, C76A, C127A	13.99

60  
17B—IL-18 BP AlphaLISA assay—An IL-18 binding protein AlphaLISA experiment substantially as described in Example 4 was performed on IL-18 polypeptide which can be incorporated into immunocytokine compositions as provided herein to assess ability to bind to IL-18BP. Results are shown in the Table below.

SEQ ID NO: or Composition ID	Sequence modifications	KD (nM)
1	Native Sequence	0.67
34	E06K, K53A, S55A	>1500
35	Y01G, F02A, E06K, M51G, K53A, D54A, S55A, T63A	969.0
36	K53A	513.8
37	S55A	10.7
38	E06K	0.13
39	E06K, K53A	130.3
40	E06K, S55A	12.3
41	K53A, S55A	500.0
42	E06K, K53A, S55A, T63A	822.0
43	E06K, K53A, S55A, Y01G	
44	E06K, K53A, S55A, F02A	>1000
45	E06K, K53A, S55A, D54A	>1000
46	E06K, K53A, S55A, M51G	>1000
47	C38S, C68S, C76S, C127S	0.03
48	C38S, C68S, C76S, C127S, K70C	0.21
49	E06K, K53A, S55A, C38S, C68S, C76S, C127S, K70C	>1000
50	E06K, K53A, T63A	339.8
51	T63A	2.59
52	E06K, T63A	0.83
53	K53A, T63A	198
54	E06K, K53A, C38S, C68S, C76S, C127S, K70C	446.0
55	K53A, T63A, C38S, C68S, C76S, C127S, K70C	913
56	E6K, K53A, C38S, C76S, C127S	435.5
57	E6K, K53A, C38S	50.2
C143	V11I, C38A, K53A, C76A, C127A	8.86
C144	V11I, C38A, K53A, T63A, C76A, C127A	0.66
C145	V11I, C38A, K53A, S55A, C76A, C127A	9.74
C146	V11I, C38A, M51G, K53A, C76A, C127A	373.30
C147	V11I, C38A, K53A, D54A, C76A, C127A	25.77
C148	F2A, V11I, C38A, K53A, C76A, C127A	57.21
C149	V11I, E31A, C38A, K53A, C76A, C127A	0.64
C150	V11I, T34A, C38A, K53A, C76A, C127A	1.24
C151	V11I, D35A, C38A, K53A, C76A, C127A	2.88
C152	V11I, S36A, C38A, K53A, C76A, C127A	1.12
C153	V11I, D37A, C38A, K53A, C76A, C127A	4.55
C154	V11I, E31A, D37A, C38A, K53A, C76A, C127A	2.12
C155	V11I, C38A, D40A, K53A, C76A, C127A	0.74
C156	V11I, C38A, N41A, K53A, C76A, C127A	18.47
C157	V11I, C38A, K53A, C76A, C127A, D132A	13.70
C158	V11I, C38A, K53A, C76A, G108A, C127A	1.24
C159	V11I, C38A, K53A, C76A, H109A, C127A	0.55
C160	V11I, C38A, K53A, C76A, D110A, C127A	0.71
C161	K8R, V11I, C38A, C76A, Q103E, C127A	0.06

-continued

SEQ ID NO: or Composition ID	Sequence modifications	KD (nM)
5	C162	K8E, V11I, C38A, C76A, Q103R, C127A
	C163	V11I, C38A, C76A, Q103K, C127A
	C164	V11I, C38A, S55H, C76A, C127A
10	C165	V11I, C38A, S55R, C76A, C127A
	C166	V11I, C38A, S55T, C76A, C127A
	C167	V11I, C38A, C76A, S105I, C127A
	C168	V11I, C38A, C76A, S105K, C127A
	C174	K8L, E6K, V11I, C38A, K53A, T63A, C76A, C127A
15	C176	E6K, V11I, C38A, I49M, K53A, T63A, C76A, C127A
	C177	E6K, V11I, C38A, I49R, K53A, T63A, C76A, C127A
	C178	E6K, V11I, C38A, K53A, T63A, C76A, Q103R, C127A
20	C179	E6K, K8E, V11I, C38A, K53A, T63A, C76A, Q103R, C127A
	C180	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153R
	C181	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153E
	C182	E6K, V11I, C38A, K53A, T63A, C76A, C127A, V153Y
	C183	E6K, V11I, C38A, M51G, K53A, T63A, C76A, C127A
25	C184	E6R, V11I, C38A, K53A, T63A, C76A, C127A
	C140	E6K, V11I, C38A, K53A, T63A, C76A, C127A
	C141	E6K, V11I, C38A, K53A, T63A, C76A, C127A
30	C142	Y1M, E6K, V11I, C38A, K53A, T63A, C76A, C127A
	C192	E6K, V11I, C38A, M51G, K53A, T63A, C76A, C127A
	62	E6K, C38A, K53A, C127A
	60	E6K, C38Q, K53A
35	59	E6K, C38A, K53A

17C—IFN $\gamma$  Stimulation and IL-18BP Inhibition Assay—  
The experiments described in Example 4 for determination of IFN $\gamma$  stimulation in NK92 cells (and inhibition by IL-18 BP) were performed substantially as described on modified IL-18 polypeptides in order to assess their activities and their suitability for incorporation into immunocytokine compositions. Results are shown in the table below.

SEQ ID NO: or Composition ID	Sequence modifications	IC <sub>50</sub> (nM)	EC <sub>50</sub> (nM)
1	Native sequence	1.47	0.276
34	E06K, K53A, S55A	229	0.824
35	Y01G, F02A, E06K, M51G, K53A, D54A, S55A, T63A	>55.0	>55.0
36	K53A	27.3	0.444
37	S55A	4.46	0.108
38	E06K	7.79	0.0567
39	E06K, K53A	>703	0.0192
40	E06K, S55A	15	0.067
41	K53A, S55A	37.3	1.58
42	E06K, K53A, S55A, T63A	1060	0.144
43	E06K, K53A, S55A, Y01G	27.8	6.12
44	E06K, K53A, S55A, F02A	NT	>1000
45	E06K, K53A, S55A, D54A	NT	30
46	E06K, K53A, S55A, M51G	0.189	7.4
47	C38S, C68S, C76S, C127S	0.444	0.115
48	C38S, C68S, C76S, C127S, K70C	0.114	0.488
49	E06K, K53A, S55A, C38S, C68S, C76S, C127S, K70C	NT	58.5
50	E06K, K53A, T63A	>1000	0.0268
51	T63A	0.239	0.449
52	E06K, T63A	47.1	0.011
53	K53A, T63A	18.2	0.155
54	E06K, K53A, C38S, C68S, C76S, C127S, K70C	23.5	0.962

-continued

SEQ ID NO: or Composition ID	Sequence modifications	IC <sub>50</sub> (nM)	EC <sub>50</sub> (nM)
55	K53A, T63A, C38S, C68S, C76S, C127S, K70C	>1000	17.2
6	E6K, V11I, C38A, K53A, T63A, C68A, C76A, C127A, D98C	5.847	1.366
5	E6K, V11I, C38A, K53A, T63A, C68A, C76A, C127A, M86C	62.37	0.075
9	E6K, C38A, K53A, C68A, D98C	960.8	0.069
4	E6K, C38A, K53A, C68A, M86C	396.3	0.022
62	E6K, C38A, K53A, C127A	283.6	0.026
60	E6K, C38Q, K53A	780.5	0.006
59	E6K, C38A, K53A	653.5	0.015
57	E6K, C38S, K53A	146.2	0.045
C143	V11I, C38A, K53A, C76A, C127A	1.625	0.138
C144	V11I, C38A, K53A, T63A, C76A, C127A	7.522	0.012
C145	V11I, C38A, K53A, S55A, C76A, C127A	10.24	0.087
C146	V11I, C38A, M51G, K53A, C76A, C127A	732.9	0.037
C147	V11I, C38A, K53A, D54A, C76A, C127A	47.63	0.079
C148	F2A, V11I, C38A, K53A, C76A, C127A	5.055	0.256
C149	V11I, E31A, C38A, K53A, C76A, C127A	1.167	0.187
C150	V11I, T34A, C38A, K53A, C76A, C127A	21.27	0.015
C151	V11I, D35A, C38A, K53A, C76A, C127A	3.622	0.061
C152	V11I, S36A, C38A, K53A, C76A, C127A	7.85	0.033
C153	V11I, D37A, C38A, K53A, C76A, C127A	2.222	0.175
C154	V11I, E31A, D37A, C38A, K53A, C76A, C127A	3.709	0.062
C155	V11I, C38A, D40A, K53A, C76A, C127A	3.233	0.067
C156	V11I, C38A, N41A, K53A, C76A, C127A	0.681	0.558
C157	V11I, C38A, K53A, C76A, C127A, D132A	6.082	0.056
C158	V11I, C38A, K53A, C76A, G108A, C127A	3.981	0.073
C159	V11I, C38A, K53A, C76A, H109A, C127A	1.807	0.123
C160	V11I, C38A, K53A, C76A, D110A, C127A	3.181	0.028
C161	K8R, V11I, C38A, C76A, Q103E, C127A	1.073	0.057
C162	K8E, V11I, C38A, C76A, Q103R, C127A	7.292	0.061
C163	V11I, C38A, C76A, Q103K, C127A	0.823	0.093
C164	V11I, C38A, S55H, C76A, C127A	0.456	0.414
C165	V11I, C38A, S55R, C76A, C127A	0.885	0.176
C166	V11I, C38A, S55T, C76A, C127A	0.44	0.098
C167	V11I, C38A, C76A, S105L, C127A	0.809	0.103
C168	V11I, C38A, C76A, S105K, C127A	0.176	0.098

Although the present disclosure and its advantages have been described in detail, it should be understood that various changes, substitutions and

alterations can be made herein without departing from the spirit and scope of the disclosure as defined in the appended claims.

## SEQUENCE LISTING

```

Sequence total quantity: 408
SEQ ID NO: 1      moltype = AA  length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = Homo sapiens

SEQUENCE: 1
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM 60
AVTISVKCEK ISTLSCENKI ISPKEMNPDD NIKDTKSII FFQRSPVGHD NKMQFESSYY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED                         157

SEQ ID NO: 2      moltype = AA  length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 2
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60
AVTISVKAEK ISTLSCENKI ISPKCMNPDD NIKDTKSII FFQRSPVGHD NKMQFESSYY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED                         157

SEQ ID NO: 3      moltype = AA  length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 3
YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60

```

-continued

---

AVAISVKAEK ISTLSAENKI ISFKCMNPPD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 4 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 4  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKAEK ISTLSCENKI ISFKECNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 5 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 5  
YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAISVKAEK ISTLSAENKI ISFKEMNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 6 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 6  
YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAISVKAEK ISTLSAENKI ISFKEMNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 7 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 7  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKAEK ISTLSAENKI ISFKEMNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 8 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 8  
YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAISVKAEK ISTLSAENKI ISFKEMNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 9 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 9  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKAEK ISTLSCENKI ISFKEMNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 10 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 10  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKAEK ISTLSAENKI ISFKEMNPYD NIKDTKSDII FFQRSVPYHD NKMQFESSYY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 11 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 11

-continued

---

YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 12 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 12  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVANISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 13 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 13  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVANISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 14 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 14  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVANISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 15 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 15  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVANISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 16 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 16  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVANIAVCKEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 17 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 17  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 18 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 18  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKAEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 19 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

-continued

SEQUENCE: 19  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENAI ISFKEMNPPD NIKDTKSDII FFQRSPVGHD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 20            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 20  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENAI ISFKEMNPPD NIKDTKSDII FFQRSPVGHD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 21            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 21  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSPVPAHD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 22            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 22  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSPVGAD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 23            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 23  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSPVGHD NAMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 24            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 24  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
 AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSDII FFQRSPVGHD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 25            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 25  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADSQPRGM 60  
 AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSDII FFQRSPVGHD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 26            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 26  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
 AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSDII FFQRSPVGHD NKMQFESSY 120  
 EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 27            moltype = AA length = 157  
 FEATURE                Location/Qualifiers  
 source                1..157  
 mol\_type = protein

-continued

---

organism = synthetic construct

SEQUENCE: 27  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 28            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 28  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 29            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 29  
YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 30            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 30  
YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 31            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 31  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYKDSQPRGM 60  
AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 32            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 32  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYKDSQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 33            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 33  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 34            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 34  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 35            moltype = AA length = 157  
FEATURE                  Location/Qualifiers  
source                  1..157

-continued

---

```

mol_type = protein
organism = synthetic construct

SEQUENCE: 35
YAGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS GYAAAQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 36      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 36
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 37      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 37
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDAQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 38      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 38
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 39      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 39
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 40      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 40
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDAQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 41      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 41
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 42      moltype = AA length = 157
FEATURE           Location/Qualifiers
source            1..157
mol_type = protein
organism = synthetic construct

SEQUENCE: 42
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM 60
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPGHD NKMQFESSY 120
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 43      moltype = AA length = 157
FEATURE           Location/Qualifiers

```

---

-continued

---

source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 43  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 44 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 44  
YAGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADAQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 45 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 45  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYAAAQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 46 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 46  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS GYADAQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 47 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 47  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYKDSQPRGM 60  
AVTISVKSEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 48 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 48  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYKDSQPRGM 60  
AVTISVKSEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 49 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 49  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADAQPRGM 60  
AVTISVKSEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 50 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 50  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSPVPGHD NMKFESSY 120  
EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 51 moltype = AA length = 157

-continued

---

FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 51  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM 60  
AVAIISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 52 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 52  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYKDSQPRGM 60  
AVAIISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 53 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 53  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 54 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 54  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 55 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 55  
YFGKLESKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM 60  
AVAIISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 56 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 56  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKCEK ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 57 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 57  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 58 moltype = AA length = 157  
FEATURE Location/Qualifiers  
source 1..157  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 58  
YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDSRD NAPRTIFIIS MYADSQPRGM 60  
AVTISVKSEC ISTLSENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
EGYFLASEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

-continued

SEQ ID NO: 59 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 59  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 60 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 60  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 61 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 61  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 62 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 62  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 63 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 63  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 64 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 64  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS MYADAQPRGM 60  
 AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 65 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 65  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDQRD NAPRTIFIIS MYADAQPRGM 60  
 AVAISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 66 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 66  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFAEMNPPD NIKDTKSII FFQRSVPGHD NMKFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

-continued

SEQ ID NO: 67 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 67  
 YFGKLKSKLS VIRNLNDQVL FIDQGNRPLF EDMTDSDCRD NAPRTIFIIS MYADSQPRGM 60  
 AVTISVKCEK ISTLSCENKI ISFKEMNPPD NIKDTKSAII FFQRSVPGHD NKMQFESSY 120  
 EGYFLACEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 68 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 68  
 YFGKLESKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS GYADSQPRGM 60  
 AVTISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NKMQFESSY 120  
 EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 69 moltype = AA length = 157  
 FEATURE Location/Qualifiers  
 source 1..157  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 69  
 YFGKLKSKLS IIRNLNDQVL FIDQGNRPLF EDMTDSDARD NAPRTIFIIS GYADSQPRGM 60  
 AVAISVKCEK ISTLSAENKI ISFKEMNPPD NIKDTKSII FFQRSVPGHD NKMQFESSY 120  
 EGYFLAAEKE RDLFKLILKK EDELGDRSIM FTVQNED 157

SEQ ID NO: 70 moltype = AA length = 158  
 FEATURE Location/Qualifiers  
 source 1..158  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 70  
 GFYFGKLKSKL SIIRNLNDQV LFIDQGNRPL FEDMTDSAR DNAPRTIFII SGYADSQPRG 60  
 MAVAISVKCE KISTLSAENK IISFKEMNPP DNIKDTKSDII IFFQRSVPGH DNKMQFESSS 120  
 YEGYFLAAEKG ERDLFKLILKK KEDELGDRSI MFTVQNED 158

SEQ ID NO: 71 moltype = AA length = 158  
 FEATURE Location/Qualifiers  
 source 1..158  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 71  
 GFYFGKLKSKL SIIRNLNDQV LFIDQGNRPL FEDMTDSAR DNAPRTIFII SMYADSQPRG 60  
 MAVAISVKCE KISTLSAENK IISFKEMNPP DNIKDTKSDII IFFQRSVPGH DNKMQFESSS 120  
 YEGYFLAAEKG ERDLFKLILKK KEDELGDRSI MFTVQNED 158

SEQ ID NO: 72 moltype = AA length = 161  
 FEATURE Location/Qualifiers  
 source 1..161  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 72  
 GGGGYFGKLK SKLSIIRNLN DQVLFIDQGN RPLFEDMTDS DARDNAPRTI FIISMYADSQ 60  
 PRGMAVAISVKCE KISTLSAENK ENKIISFKEM NPPDNIKDTK SDIIFFQRSV PGHDNMQFE 120  
 SSSYEGYFLA AAKERDLFKL ILKKEDELGD RSIMFTVQNE D 161

SEQ ID NO: 73 moltype = length =  
 SEQUENCE: 73  
 000

SEQ ID NO: 74 moltype = length =  
 SEQUENCE: 74  
 000

SEQ ID NO: 75 moltype = length =  
 SEQUENCE: 75  
 000

SEQ ID NO: 76 moltype = length =  
 SEQUENCE: 76  
 000

SEQ ID NO: 77 moltype = length =  
 SEQUENCE: 77

-continued

000

```

SEQ ID NO: 78      moltype = length =
SEQUENCE: 78
000

SEQ ID NO: 79      moltype = length =
SEQUENCE: 79
000

SEQ ID NO: 80      moltype = length =
SEQUENCE: 80
000

SEQ ID NO: 81      moltype = length =
SEQUENCE: 81
000

SEQ ID NO: 82      moltype = length =
SEQUENCE: 82
000

SEQ ID NO: 83      moltype = length =
SEQUENCE: 83
000

SEQ ID NO: 84      moltype = length =
SEQUENCE: 84
000

SEQ ID NO: 85      moltype = length =
SEQUENCE: 85
000

SEQ ID NO: 86      moltype = length =
SEQUENCE: 86
000

SEQ ID NO: 87      moltype = length =
SEQUENCE: 87
000

SEQ ID NO: 88      moltype = length =
SEQUENCE: 88
000

SEQ ID NO: 89      moltype = length =
SEQUENCE: 89
000

SEQ ID NO: 90      moltype = length =
SEQUENCE: 90
000

SEQ ID NO: 91      moltype = length =
SEQUENCE: 91
000

SEQ ID NO: 92      moltype = length =
SEQUENCE: 92
000

SEQ ID NO: 93      moltype = length =
SEQUENCE: 93
000

SEQ ID NO: 94      moltype = length =
SEQUENCE: 94
000

SEQ ID NO: 95      moltype = length =
SEQUENCE: 95
000

SEQ ID NO: 96      moltype = length =
SEQUENCE: 96
000

SEQ ID NO: 97      moltype = length =

```

-continued

```

SEQUENCE: 97          moltype =    length =
000

SEQ ID NO: 98          moltype =    length =
SEQUENCE: 98
000

SEQ ID NO: 99          moltype =    length =
SEQUENCE: 99
000

SEQ ID NO: 100         moltype =    length =
SEQUENCE: 100
000

SEQ ID NO: 101         moltype =    length =
SEQUENCE: 101
000

SEQ ID NO: 102         moltype =    length =
SEQUENCE: 102
000

SEQ ID NO: 103         moltype =    length =
SEQUENCE: 103
000

SEQ ID NO: 104         moltype =    length =
SEQUENCE: 104
000

SEQ ID NO: 105         moltype =    length =
SEQUENCE: 105
000

SEQ ID NO: 106         moltype =    length =
SEQUENCE: 106
000

SEQ ID NO: 107         moltype =    length =
SEQUENCE: 107
000

SEQ ID NO: 108         moltype =    length =
SEQUENCE: 108
000

SEQ ID NO: 109         moltype =    length =
SEQUENCE: 109
000

SEQ ID NO: 110         moltype =    length =
SEQUENCE: 110
000

SEQ ID NO: 111         moltype =    length =
SEQUENCE: 111
000

SEQ ID NO: 112         moltype =    length =
SEQUENCE: 112
000

SEQ ID NO: 113         moltype =    length =
SEQUENCE: 113
000

SEQ ID NO: 114         moltype =    length =
SEQUENCE: 114
000

SEQ ID NO: 115         moltype =    length =
SEQUENCE: 115
000

SEQ ID NO: 116         moltype =    length =
SEQUENCE: 116
000

```

-continued

---

```

SEQ ID NO: 117      moltype =   length =
SEQUENCE: 117
000

SEQ ID NO: 118      moltype =   length =
SEQUENCE: 118
000

SEQ ID NO: 119      moltype =   length =
SEQUENCE: 119
000

SEQ ID NO: 120      moltype =   length =
SEQUENCE: 120
000

SEQ ID NO: 121      moltype =   length =
SEQUENCE: 121
000

SEQ ID NO: 122      moltype =   length =
SEQUENCE: 122
000

SEQ ID NO: 123      moltype =   length =
SEQUENCE: 123
000

SEQ ID NO: 124      moltype =   length =
SEQUENCE: 124
000

SEQ ID NO: 125      moltype =   length =
SEQUENCE: 125
000

SEQ ID NO: 126      moltype =   length =
SEQUENCE: 126
000

SEQ ID NO: 127      moltype =   length =
SEQUENCE: 127
000

SEQ ID NO: 128      moltype =   length =
SEQUENCE: 128
000

SEQ ID NO: 129      moltype =   length =
SEQUENCE: 129
000

SEQ ID NO: 130      moltype =   length =
SEQUENCE: 130
000

SEQ ID NO: 131      moltype =   length =
SEQUENCE: 131
000

SEQ ID NO: 132      moltype =   length =
SEQUENCE: 132
000

SEQ ID NO: 133      moltype =   length =
SEQUENCE: 133
000

SEQ ID NO: 134      moltype =   length =
SEQUENCE: 134
000

SEQ ID NO: 135      moltype =   length =
SEQUENCE: 135
000

SEQ ID NO: 136      moltype =   length =
SEQUENCE: 136
000

```

-continued

```

SEQ ID NO: 137      moltype =   length =
SEQUENCE: 137
000

SEQ ID NO: 138      moltype =   length =
SEQUENCE: 138
000

SEQ ID NO: 139      moltype =   length =
SEQUENCE: 139
000

SEQ ID NO: 140      moltype =   length =
SEQUENCE: 140
000

SEQ ID NO: 141      moltype =   length =
SEQUENCE: 141
000

SEQ ID NO: 142      moltype =   length =
SEQUENCE: 142
000

SEQ ID NO: 143      moltype =   length =
SEQUENCE: 143
000

SEQ ID NO: 144      moltype =   length =
SEQUENCE: 144
000

SEQ ID NO: 145      moltype =   length =
SEQUENCE: 145
000

SEQ ID NO: 146      moltype =   length =
SEQUENCE: 146
000

SEQ ID NO: 147      moltype =   length =
SEQUENCE: 147
000

SEQ ID NO: 148      moltype =   length =
SEQUENCE: 148
000

SEQ ID NO: 149      moltype =   length =
SEQUENCE: 149
000

SEQ ID NO: 150      moltype =   length =
SEQUENCE: 150
000

SEQ ID NO: 151      moltype =   length =
SEQUENCE: 151
000

SEQ ID NO: 152      moltype =   length =
SEQUENCE: 152
000

SEQ ID NO: 153      moltype =   length =
SEQUENCE: 153
000

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

SEQ ID NO: 155      moltype =   length =
SEQUENCE: 155
000

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

```

-continued

```

000

SEQ_ID NO: 157      moltype = length =
SEQUENCE: 157
000

SEQ_ID NO: 158      moltype = length =
SEQUENCE: 158
000

SEQ_ID NO: 159      moltype = length =
SEQUENCE: 159
000

SEQ_ID NO: 160      moltype = length =
SEQUENCE: 160
000

SEQ_ID NO: 161      moltype = length =
SEQUENCE: 161
000

SEQ_ID NO: 162      moltype = length =
SEQUENCE: 162
000

SEQ_ID NO: 163      moltype = length =
SEQUENCE: 163
000

SEQ_ID NO: 164      moltype = length =
SEQUENCE: 164
000

SEQ_ID NO: 165      moltype = length =
SEQUENCE: 165
000

SEQ_ID NO: 166      moltype = length =
SEQUENCE: 166
000

SEQ_ID NO: 167      moltype = length =
SEQUENCE: 167
000

SEQ_ID NO: 168      moltype = length =
SEQUENCE: 168
000

SEQ_ID NO: 169      moltype = length =
SEQUENCE: 169
000

SEQ_ID NO: 170      moltype = length =
SEQUENCE: 170
000

SEQ_ID NO: 171      moltype = length =
SEQUENCE: 171
000

SEQ_ID NO: 172      moltype = length =
SEQUENCE: 172
000

SEQ_ID NO: 173      moltype = length =
SEQUENCE: 173
000

SEQ_ID NO: 174      moltype = length =
SEQUENCE: 174
000

SEQ_ID NO: 175      moltype = length =
SEQUENCE: 175
000

SEQ_ID NO: 176      moltype = length =

```

-continued

```

SEQUENCE: 176          moltype =    length =
000

SEQ ID NO: 177          moltype =    length =
SEQUENCE: 177
000

SEQ ID NO: 178          moltype =    length =
SEQUENCE: 178
000

SEQ ID NO: 179          moltype =    length =
SEQUENCE: 179
000

SEQ ID NO: 180          moltype =    length =
SEQUENCE: 180
000

SEQ ID NO: 181          moltype =    length =
SEQUENCE: 181
000

SEQ ID NO: 182          moltype =    length =
SEQUENCE: 182
000

SEQ ID NO: 183          moltype =    length =
SEQUENCE: 183
000

SEQ ID NO: 184          moltype =    length =
SEQUENCE: 184
000

SEQ ID NO: 185          moltype =    length =
SEQUENCE: 185
000

SEQ ID NO: 186          moltype =    length =
SEQUENCE: 186
000

SEQ ID NO: 187          moltype =    length =
SEQUENCE: 187
000

SEQ ID NO: 188          moltype =    length =
SEQUENCE: 188
000

SEQ ID NO: 189          moltype =    length =
SEQUENCE: 189
000

SEQ ID NO: 190          moltype =    length =
SEQUENCE: 190
000

SEQ ID NO: 191          moltype =    length =
SEQUENCE: 191
000

SEQ ID NO: 192          moltype =    length =
SEQUENCE: 192
000

SEQ ID NO: 193          moltype =    length =
SEQUENCE: 193
000

SEQ ID NO: 194          moltype =    length =
SEQUENCE: 194
000

SEQ ID NO: 195          moltype =    length =
SEQUENCE: 195
000

```

-continued

```

SEQ ID NO: 196      moltype =   length =
SEQUENCE: 196
000

SEQ ID NO: 197      moltype =   length =
SEQUENCE: 197
000

SEQ ID NO: 198      moltype =   length =
SEQUENCE: 198
000

SEQ ID NO: 199      moltype =   length =
SEQUENCE: 199
000

SEQ ID NO: 200      moltype =   length =
SEQUENCE: 200
000

SEQ ID NO: 201      moltype =   length =
SEQUENCE: 201
000

SEQ ID NO: 202      moltype =   length =
SEQUENCE: 202
000

SEQ ID NO: 203      moltype =   length =
SEQUENCE: 203
000

SEQ ID NO: 204      moltype =   length =
SEQUENCE: 204
000

SEQ ID NO: 205      moltype =   length =
SEQUENCE: 205
000

SEQ ID NO: 206      moltype =   length =
SEQUENCE: 206
000

SEQ ID NO: 207      moltype =   length =
SEQUENCE: 207
000

SEQ ID NO: 208      moltype =   length =
SEQUENCE: 208
000

SEQ ID NO: 209      moltype =   length =
SEQUENCE: 209
000

SEQ ID NO: 210      moltype = AA  length = 152
FEATURE
source
1..152
mol_type = protein
organism = Homo sapiens
SEQUENCE: 210
DCDIEGKDGK QYESVLMVSI DQLLDSMKEI GSNCLNNNEFN FFKRHICDAN KEGMFLFRAA 60
RKLRQFLKMN STGDFDLHLL KVSEGTTILL NCTGQVKGRK PAALGEAQPT KSLEENKSLK 120
EQKKLNLDLCF LKRLLQEIKT CWNKILMGTK EH 152

SEQ ID NO: 211      moltype = AA  length = 133
FEATURE
source
1..133
mol_type = protein
organism = Homo sapiens
SEQUENCE: 211
APTSSSSTKKT QLQLEHLLL D LQMILNGINN YKNPKLTRML TFKFYMPKKA TELKHLQCLE 60
EELKPLETEEVL NLAQSKNFHL RPRDLISNIN VIVLELGSE TTFMCEYADE TATIVEFLNR 120
WITFCQSIIS TLT 133

SEQ ID NO: 212      moltype =   length =
SEQUENCE: 212
000

```

-continued

```

SEQ ID NO: 213      moltype = length =
SEQUENCE: 213
000

SEQ ID NO: 214      moltype = length =
SEQUENCE: 214
000

SEQ ID NO: 215      moltype = length =
SEQUENCE: 215
000

SEQ ID NO: 216      moltype = length =
SEQUENCE: 216
000

SEQ ID NO: 217      moltype = length =
SEQUENCE: 217
000

SEQ ID NO: 218      moltype = length =
SEQUENCE: 218
000

SEQ ID NO: 219      moltype = length =
SEQUENCE: 219
000

SEQ ID NO: 220      moltype = length =
SEQUENCE: 220
000

SEQ ID NO: 221      moltype = length =
SEQUENCE: 221
000

SEQ ID NO: 222      moltype = length =
SEQUENCE: 222
000

SEQ ID NO: 223      moltype = length =
SEQUENCE: 223
000

SEQ ID NO: 224      moltype = AA length = 20
FEATURE
source
1..20
mol_type = protein
organism = synthetic construct
VARIANT
1..20
note = This sequence may encompass 1-10 GS repeating units
SEQUENCE: 224
GSGSGSGSGS GSGSGSGSGS                                         20

SEQ ID NO: 225      moltype = AA length = 30
FEATURE
source
1..30
mol_type = protein
organism = synthetic construct
VARIANT
1..30
note = This sequence may encompass 1-10 GGS repeating units
SEQUENCE: 225
GGSGGGSGGSG GGGGGGGGGG SGGGGGGGGS                                         30

SEQ ID NO: 226      moltype = AA length = 40
FEATURE
source
1..40
mol_type = protein
organism = synthetic construct
VARIANT
1..40
note = This sequence may encompass 1-10 GGGS repeating units
SEQUENCE: 226
GGGSGGGGGG GGGGGGGGGG GGGGGGGGGG GGGGGGGGGG                                         40

SEQ ID NO: 227      moltype = AA length = 40
FEATURE
source
1..40
mol_type = protein

```

-continued

organism = synthetic construct  
 1..40  
 note = This sequence may encompass 1-10 GGSG repeating units  
**SEQUENCE: 227**  
 GGGGGGGGGG SGGGGGGGGG GGSGGGGGGG SGGS  
40  
**SEQ ID NO: 228** moltype = AA length = 50  
**FEATURE** Location/Qualifiers  
**source** 1..50  
**mol\_type** = protein  
**organism** = synthetic construct  
**VARIANT** 1..50  
**note** = This sequence may encompass 1-10 GGSGG repeating units  
**SEQUENCE: 228**  
 GGGGGGGGGG GGSGGGGGGG GGSGGGGGGG GGSGGGGGGG GGSGGGGGGG  
50  
**SEQ ID NO: 229** moltype = AA length = 50  
**FEATURE** Location/Qualifiers  
**source** 1..50  
**mol\_type** = protein  
**organism** = synthetic construct  
**VARIANT** 1..50  
**note** = This sequence may encompass 1-10 GGGGS repeating units  
**SEQUENCE: 229**  
 GGGGSGGGGS GGGGSGGGGS GGGGSGGGGS GGGGSGGGGS GGGGSGGGGS  
50  
**SEQ ID NO: 230** moltype = AA length = 15  
**FEATURE** Location/Qualifiers  
**source** 1..15  
**mol\_type** = protein  
**organism** = synthetic construct  
**SEQUENCE: 230**  
 GGGGSGGGGS GGGGS  
15  
**SEQ ID NO: 231** moltype = AA length = 20  
**FEATURE** Location/Qualifiers  
**source** 1..20  
**mol\_type** = protein  
**organism** = synthetic construct  
**SEQUENCE: 231**  
 GGGGSGGGGS GGGGSGGGGS  
20  
**SEQ ID NO: 232** moltype = AA length = 450  
**FEATURE** Location/Qualifiers  
**source** 1..450  
**mol\_type** = protein  
**organism** = synthetic construct  
**SEQUENCE: 232**  
 EVQLESGGG LVQPGGSSLRL SCAASGFTFS SYIMMWVRQA PGKGLEWVSS IYPSGGITFY 60  
 ADTVKGRTFI SRDNNSKNTLY LQMNSLRAED TAVYYCARIK LGTVTTVDYW GQGTLTVSS 120  
 ASTKGPSVFP LAPSSKSTSG GTAAALGCLVK DYFPEPVTVS WNSGALTSGV HTFPFAVLQSS 180  
 GLYSLSVSVT VPSSSLGTQT YICNVNWKPS NTKVDKKVEP KSCDKTHTCP PCPAPELLGG 240  
 LFPPFLFPKPK DKTLMISRTP HEDEPVKPNW YVDGVEVHN A KTKPREEQYN 300  
 STYRVVSVLT VLHQDWLNGK EYKCKVSNKA LPAPIEKATIS KAKGQPREPQ VYTLPSPRDE 360  
 LTKNQVSLTC LVKGFYPSDI AVEWESNGP ENNYKTTTPV LDSDGSFFLY SKLTVDKSRW 420  
 QQGNVFSCSV MHEALHNHYT QKSLSLSPGK  
450  
**SEQ ID NO: 233** moltype = AA length = 216  
**FEATURE** Location/Qualifiers  
**source** 1..216  
**mol\_type** = protein  
**organism** = synthetic construct  
**SEQUENCE: 233**  
 QSALTQPSA SGSPGQSITI SCTGTSSDVG GYNVYWSYQQ HPGKAPKLMY YDVSNRPSGV 60  
 SNRFSGSKSG NTASLTISGL QAEADEADYYC SSYTSSSTRV FGTGKVTVL GQPKANPVT 120  
 LFPPSSEELQ ANKATLVCLI SDFYYPGAVTV AWKADGSPVK AGVETTKPSK QSNNKYAASS 180  
 YLSLTPEQWK SHRSYSCQVT HEGSTVEKTV APTECS  
216  
**SEQ ID NO: 234** moltype = AA length = 451  
**FEATURE** Location/Qualifiers  
**source** 1..451  
**mol\_type** = protein  
**organism** = synthetic construct  
**SEQUENCE: 234**  
 EVQLVESGGG LVQPGGSSLRL SCAASGFTFS RYWMWSWRQA PGKGLEWVAN IKQDGSEKYY 60  
 VDSVKGRTFI SRDNNAKNSLY LQMNSLRAED TAVYYCAREG GWFGELAFDY WGQGTLTVVS 120  
 SASTKGPSVFP LAPSSKSTSG GTAAALGCLVK KYDFPEPVTV SWNSGALTSGV VHTFPAVLQSS 180

## US 12,383,631 B2

181

182

-continued

---

SGLYSLSSVV	TVPSSSLGTQ	TYICVNHNKP	SNTKVDKRVE	PKSCDKTHTC	PPCPAPEFEG	240
GPSVFLPPPK	PKDLMISRT	PEVTCVVVDV	SHEDPEVKFN	WYVDGVEVHN	AKTKPREEQY	300
NSTYRVVSVL	TVLHQDWLNG	KEYCKVSNK	ALPASIEKTI	SKAKGQREP	QVYTLPPSR	360
EMTKNQVSLT	CLVKGFYP PSD	IAVEWESNGQ	PENNYKTPP	VLDSDGSFFL	YSKLTVDKSR	420
WQQGNVFSCS	VMHEALHNHY	TQKSLSLSPG	K			451

---

SEQ ID NO: 235      moltype = AA length = 215  
 FEATURE              Location/Qualifiers  
 source              1..215  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 235  
 EIVLTQSPGT LSLSPGERAT LSCRASQRVS SSYLAWSYQOK PGQAPRLLIY DASSRATGIP 60  
 DRPSGSGSGT DFTLTISRLE PEDFAVYYCQ QYGSLPWTFG QGTVKEIKRT VAAPSVFIFP 120  
 PSDEQLKSGT ASVVCLNNF YPREAKVQWK VDNALQSGNS QESVTEQDSK DSTYLSSTL 180  
 TLSKADYEKHH KVYACEVTHQ GLSSPVTKSF NRGE 215

SEQ ID NO: 236      moltype = AA length = 448  
 FEATURE              Location/Qualifiers  
 source              1..448  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 236  
 EVQLVESGGG LVQPGGSLRL SCAASGFTFS DSWIHWVRQA PGKGLEWVAW ISPYGGSTYY 60  
 ADSVVKGRFTI SADTSKNTAY LQMNSLRAED TAVYYCARH WPGGFYWGQ GTLTVSSAS 120  
 TKGPSVPLA PSSKSTSGGT AALGCLVKDY FPEPVTVSWN SGALTSGVHT FPAVLQSSGL 180  
 YSLSSVVTVP SSSLGTQTYI CNVNHKPSNT KVDKKVPEPKS CDKTHTCPG PAPELLGGPS 240  
 VFLFPPKPKD TLMISRTPEV TCVVVDDVSHE DPEVKFNWYV DGVEVHNAAKT KPREEQYAST 300  
 YRVSVSLTQL HODWLNGKEY KCKVSNKALP APIEKTISKA KGQPREPQVY TLPPSREEMT 360  
 KNQVSLTCLV KGFYPSDIAV EWESNGQOPEN NYKTTTPVLD SDGSFFLYSK LTVDKSRWQQ 420  
 GNVFSCSVMH EALHNHYTQK SLSLSPGK 448

SEQ ID NO: 237      moltype = AA length = 214  
 FEATURE              Location/Qualifiers  
 source              1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 237  
 DIQMTQSPSS LSASVGDRVT ITCRASQDV S TAVAQYQQPK GKAPKLLIYS ASFLYSGVPS 60  
 RFSGSGSGTD FTLTIISSLQP EDFATYYCQQ YLYHPATFGQ GTKVEIKRTV AAPSVFIFP 120  
 SDBQLKSGTA SVVCLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYLSSTLT 180  
 LSKADYEKHH KVYACEVTHQ LSSPVTKSFN RGEC 214

SEQ ID NO: 238      moltype = AA length = 448  
 FEATURE              Location/Qualifiers  
 source              1..448  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 238  
 EVQLVESGGG LVQPGGSLRL SCAASGFTFS SYAMSWVRQA PGKGLEWVSG ISGSGGFTYY 60  
 ADSVVKGRFTI SRDNSKNTLY LQMNSLRAED TAVYYCAKPP RGYNYGPFDY WGQGTLTVS 120  
 SASTKGPSVF PLAPCRSTS ESTAAALGCLV KDYFPEPVTV SWNSGALTSG VHTFPAVLQS 180  
 SGGLYSLSSVV TVPSSSLGTT TYTCNVDHKP SNTKVDKRVE SKYGPCCPC PAPEFLGGPS 240  
 VFLFPPKPKD TLMISRTPEV TCVVVDDVSQE DPEVQFNWYV DGVEVHNAAKT KPREEQFNST 300  
 YRVSVSLTQL HODWLNGKEY KCKVSNKGLP SIEKTISKA KGQPREPQVY TLPPSQEEMT 360  
 KNQVSLTCLV KGFYPSDIAV EWESNGQOPEN NYKTTTPVLD SDGSFFLYSR LTVDKSRWQE 420  
 GNVFSCSVMH EALHNHYTQK SLSLSLGK 448

SEQ ID NO: 239      moltype = AA length = 214  
 FEATURE              Location/Qualifiers  
 source              1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 239  
 SYVLTQPPSV SVAPGQTARI TCGGNNIGSK SVHWFYQQKPG QAPVLVYDD SDRPSGIPER 60  
 FSGSNSGNTA TLTISRVEAG DEADYYCQW DSSSDDHVVFQ GGTKLTVLGQ PKAAPSVTLF 120  
 PPSSEELQAN KATLVCLISD FYPGAVTVAW KADSSPVKAG VETTTPSKQS NNKYAASSYL 180  
 SLTPEQWKSH RSYSCQVTHE GSTVEKTVAP TECS 214

SEQ ID NO: 240      moltype = length =  
 SEQUENCE: 240  
 000

SEQ ID NO: 241      moltype = length =  
 SEQUENCE: 241  
 000

SEQ ID NO: 242      moltype = AA length = 118  
 FEATURE              Location/Qualifiers

-continued

---

source 1..118  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 242  
QGQLQESGPS LVKPSQTL SL TCTVSGDSIT RGYWNWIRKH PGKGLEYIYG ISYTGSTYSN 60  
LSLKSRTVIS RDTSKNQYYL KLSSVTAADT AVYYCATSTG WLDPVDYWQ GTLTVSS 118

SEQ ID NO: 243 moltype = AA length = 107  
FEATURE Location/Qualifiers  
source 1..107  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 243  
DIVMTQSPDS LAVSLGERAT INCKASQNV D TSVAWFQQKP GQPPKALIYS ASFRYSGVPD 60  
RFSGSGSGTD FTLTISLQA EDVAVYFCQQ YYGYPFTFGQ GTKLEIK 107

SEQ ID NO: 244 moltype = AA length = 123  
FEATURE Location/Qualifiers  
source 1..123  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 244  
EVOLVQSGGG LVKPGGSLRL SCAASGFTFS SYTMNWRQ A PGKGLEWVSS ISSGSDYLYY 60  
ADSVKGRFTI SRDNAKNSLY LQMNSLRAED TAVYYCARNE LRWYPQAGAF DRWGQQGMVT 120  
VSS 123

SEQ ID NO: 245 moltype = AA length = 109  
FEATURE Location/Qualifiers  
source 1..109  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 245  
QSVTQPPSM SAAPGQRVTI SCSSGSSYIE SSYVGWYQQL PGTAPRLLIY DDDMRPSSGIP 60  
DRFSGSKSGT SATLAITGLQ TGDEADYYCE IWRSGLGGVF GGGTKLTIVL 109

SEQ ID NO: 246 moltype = length =  
SEQUENCE: 246  
000

SEQ ID NO: 247 moltype = length =  
SEQUENCE: 247  
000

SEQ ID NO: 248 moltype = AA length = 453  
FEATURE Location/Qualifiers  
source 1..453  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 248  
QVQLVQSGAE VKKPGSSVKV SCKASGGTFS SYAISWRQ A PGQGLEWMGG IIPIFGTANY 60  
AQKFQGRVTI TADKSTSTAY MELSSLRSED TAVYYCARSP DYSPYYYYGM DVWGQGTTVT 120  
VSSASTKGPS VFPLAPSSKS TSGGTAALGC LVKDYFPPEPV TVSWNSGALT SGVHTFPAVL 180  
QSSGLYSLSS VVTVPSSSLG TQTYICNVNH KPSNTKVDKR VEKPSCDKTH TCPPCPAPEA 240  
EGAPSFLFP PKPKDITLMS RTPEVTCVVV DVSHEDPEVK FNWYVDGVEV HNAKTKPREE 300  
QYNSTYRVVS LTVLHQDWL NGKAKCCKVS NKLAPSSIEK TISKAKGQPR EPQVYTLPPS 360  
REEMTKNQVS LTCLVKGFYP SDIAVEWESN QOPENNNYKITT PPVLDSDGSF FLYSKLTIVDK 420  
SRWQQGNVFS CSVMEHALHN HYTQKSLSL S PKG 453

SEQ ID NO: 249 moltype = AA length = 216  
FEATURE Location/Qualifiers  
source 1..216  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 249  
QSVLTQPPSA SGTPGQRVTI SCSSGSSNIG SNTVNWYQQL PGTAPKLLIY GNSNRPSGVP 60  
DRFSGSKSGT SASLAISGLQ SEDEADYYCQ SYDSSLGSV FGGGIKLTVL GQPKAAPS VT 120  
LFFPSSEELQ ANKATLVCLI SDFYFGAVTV AWKADSSPVK AGVETTTPSK QSNNKYAASS 180  
YLSLTPEQWK SHRSYSCQVT HEGSTVEKTV APAECS 216

SEQ ID NO: 250 moltype = AA length = 5  
FEATURE Location/Qualifiers  
source 1..5  
mol\_type = protein  
organism = synthetic construct

SEQUENCE: 250  
SYTMN

SEQ ID NO: 251 moltype = AA length = 17  
FEATURE Location/Qualifiers

-continued

---

source	1..17 mol_type = protein organism = synthetic construct	
SEQUENCE: 251 SISSGSDLY YADSVKG		17
SEQ ID NO: 252 FEATURE source	moltype = AA length = 14 Location/Qualifiers 1..14 mol_type = protein organism = synthetic construct	
SEQUENCE: 252 NELRWYPQAG AFDR		14
SEQ ID NO: 253 FEATURE source	moltype = AA length = 13 Location/Qualifiers 1..13 mol_type = protein organism = synthetic construct	
SEQUENCE: 253 SGSSSYIESS YVG		13
SEQ ID NO: 254 FEATURE source	moltype = AA length = 7 Location/Qualifiers 1..7 mol_type = protein organism = synthetic construct	
SEQUENCE: 254 DDDMRPS		7
SEQ ID NO: 255 FEATURE source	moltype = AA length = 10 Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 255 EIWRSGLGGV		10
SEQ ID NO: 256 FEATURE source	moltype = AA length = 360 Location/Qualifiers 1..360 mol_type = protein organism = synthetic construct	
SEQUENCE: 256 QVQLVESGGG LVOPGGSLRL SCAASGKMS RRCMAWFRQA PGKERERVAK LLTTSGSTYL ADSVKGRFTI SRDNSKNTVY LQMNSLRAED TAVYYCAADS FEDPTCTLVT SSGAQYWGQ GTLVTVSSEP KSSDKTHTCP PCPAPELLGG PSVFLFPKP KDTLMISRTP EVTCVVVA HEDPEVKFENW YVDGVEVHN A KTKPREEQYN STYRVVSVLT VLHQDWLNKG EYKCKVSNKA LPAGIEKTIS KAKGQPREPQ VYTLPPSRDE LTKNQVSLTC LVKGFYPSDI AVEWESNGQP ENNYKTTTPV LDSDGSFFLY SKLTVDKSRW QQGNVFSCSV MHEALHNHYT QKSLSLSPKG	60 120 180 240 300 360 420 480 540 600 660 720 759	
SEQ ID NO: 257 FEATURE source	moltype = AA length = 759 Location/Qualifiers 1..759 mol_type = protein organism = synthetic construct	
SEQUENCE: 257 DIQMTQSPAS LSASVGDRVT ITCQASQSIG TYLAWYQQKP GKPPKLLIYR AFILASGVPS RFSGSGSGTD FTTLTISSLQP EDFATYYCQS NFYSDSTTIG PNAFGTGTKV TVLGGGGSE VQLVESGGGL VQPGGSLRLS CAASGFSFSA NYPPCWVRQA PGKGLEWIGC IYGGSSDIT DANWTKGRTF ISRDNSKNTV YLQMNSLRAE DTAVYYCARS AWYSGWGGDL WGQGTLVTVS SGGGGSGGGG SGGGGSGGGG SIQMTQSPSS LSASVGDRVT ITCQASQSIIS NRLAWYQQKP GKAPKLLIYS ASTLASGVPS RFSGSGSGTD FTTLTISSLQP EDFATYYCQS TYYGNDGNAF GTGTKVTVLQ GGGGSEBVQLV ESGGGLVQPG GSRLRSLCAAS GFSFNDSYWI YWVRQAPGKG LEWIASIYGG SSGNTQYASW AQGRFTISRD NSKNTVYQLM NSLRAEDTAV YFCARGYVDY GGATDLWQGQ TLTVTSSGGG GSgggggSIQM TQSPSSLSAS VGDRVTITCQ SSESVYNNQ LSWYQQKPGQ PPKLLIYDFT DLAQGPVPSR SGSGSGTDF LTISISSLQPED FATYYCAGGF SSSSDRAFTSGG GTKLTVLGGG GGSGGGGGSEV QLVESEGGGLV QPGGSLRLSC AASGFSLSSN AMGWRQAPG KGLEYIGIIS VGGFTYYASW AKGRFTISRD NSKNTVYQLM NSLRAEDTAT YFCARDRHGG DSSGAFYLWG QGTLTVSS	60 120 180 240 300 360 420 480 540 600 660 720 759	
SEQ ID NO: 258 FEATURE source	moltype = AA length = 942 Location/Qualifiers 1..942 mol_type = protein organism = synthetic construct	
SEQUENCE: 258 QMQLVQSGAE VKKPQSSVKV SCKASGGTFS SYAISWVRQA PGQGLEWMGR IIPILGIAN AQKFQGRVTI TADKSTSTAY MELSSLRSED TAVYYCAKPR DGYNLVAFDI WGQGTMVTVS	60 120	

-continued

---

SASTKGPSVF PLAPSSKSTS GGTAALGCLV KDYFPEPVTV SWNSGALTSG VHTPPAVLQS 180  
 SGLYSLSSV TVPSSSLGTQ TYICNVNHPK SNKVDKKVE PKSCDKTHTC PPCPAPELLG 240  
 GPGGGGSEVQ LQQSGPELVK PGPSMKISCK ASGYSFTGYT MNWVKQSHGK NLEWMLINP 300  
 YKGVSTYNQK FKDKATLTVD KSSSTAYMEL LSLTSEDSAV YYCARSGYYG DSDWYFDVWG 360  
 QGTTLTFSQ MQLVQSGAEV KKPGGSVVK CSAGSGTFSS YAISWVRQAP GQGLEWMGRI 420  
 IPILGIANYA QKFQGRVTIT ADKSTSTAYM ELSSLRSEDT AVYYCAKPRD GYNLVAFDIW 480  
 GQGTMVTVSS ASTKGPSVFP LAPSSKSTS GTAAALGCLVK DYFPEPVTVS WNSGALTSGV 540  
 HTFPAPLQSS GLYSLSSVVT VPSSSLGTQT YICNVNHPKS NTKVDKKVEP KSCDKTHTCP 600  
 PCPAPELLGG PGGGGSDIQN TQTTSSLAS LGDRVTISCR ASQDIRNYLN WYQQKPDGTV 660  
 KLLIYYTSRL HSGVPSFKFG SGSGTDYSLT ISNLEQEDIA TYFCQQGNTL PWTFAGGTKL 720  
 EIKRQLVLTQ PPSVSGAPGQ RVTISCTGSS SNIGAGYDVH WYQQLPQAAP KLLIYGDINR 780  
 PSGVPDRFSG SKSGISASLA ITGLQAEDEA DYYCQSYDS LSGGVFGGGT KLTVLRTVAA 840  
 PSVFIFPPSD EQLKSGTASV VCLLNNFYPR EAKVQWKDN ALQSGNSQES VTEQDSKDST 900  
 YSLSSTLTLs KADYEKHKVY ACEVTHQGLS SPVTKSFNRG EC 942

SEQ ID NO: 259 moltype = length =  
 SEQUENCE: 259  
 000

SEQ ID NO: 260 moltype = AA length = 230  
 FEATURE Location/Qualifiers  
 source 1..230  
 mol\_type = protein  
 organism = synthetic construct  
 VARIANT 18  
 note = Any naturally occurring amino acid  
 VARIANT 19  
 note = Any naturally occurring amino acid  
 VARIANT 21  
 note = Any naturally occurring amino acid  
 VARIANT 140  
 note = Any naturally occurring amino acid  
 VARIANT 142  
 note = Any naturally occurring amino acid  
 VARIANT 177  
 note = Any naturally occurring amino acid  
 SEQUENCE: 260  
 PKSCDKTHTC PPCPAPEXXG XPSVFLFPPK PKDTLMISR PEVTCVVVDV SHEDPEVKFN 60  
 WXXVGVEVHN AKTKPREEQY DSTYRVVSVL TVLHQDWLNG KEYKCKVSNK ALPAPIEKTI 120  
 SKAKGQPREP QVYTLPPSRX EXTKNQVSLT CLVKGFYPSD IAVEWESNGQ PENNYKXTPP 180  
 VLDSDGSSFL YSKLTVDKSR WQQGNVFSCS VMHEALHNHY TQKSLSLSPG 230

SEQ ID NO: 261 moltype = AA length = 48  
 FEATURE Location/Qualifiers  
 source 1..48  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 261  
 QETNPTENLY FQQKNMQCQR RFYEALHDPN LNEEQRNARI RSIRDDDC 48

SEQ ID NO: 262 moltype = AA length = 61  
 FEATURE Location/Qualifiers  
 source 1..61  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 262  
 QTADNQNMQ CQRRFYEALH DPNLNEEQRN ARIRSIRDDC SQSANLLAEA QQLNDAQAPQ 60  
 A 61

SEQ ID NO: 263 moltype = AA length = 38  
 FEATURE Location/Qualifiers  
 source 1..38  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 263  
 QETKNMQCQR RFYEALHDPN LNEEQRNARI RSIRDDDC 38

SEQ ID NO: 264 moltype = AA length = 38  
 FEATURE Location/Qualifiers  
 source 1..38  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 264  
 QETFNKQCQR RFYEALHDPN LNEEQRNARI RSIRDDDC 38

SEQ ID NO: 265 moltype = AA length = 38  
 FEATURE Location/Qualifiers  
 source 1..38  
 mol\_type = protein

-continued

---

SEQUENCE: 265	organism = synthetic construct
QETFNMQCQR RFYEALHDPN LNKEQRNARI RSIRDDDC	38
SEQ ID NO: 266	moltype = AA length = 37
FEATURE	Location/Qualifiers
source	1..37
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 266	
QETFNMQCQR RFYEALHDPN LNEEQRNARI RSIKDDC	37
SEQ ID NO: 267	moltype = AA length = 35
FEATURE	Location/Qualifiers
source	1..35
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 267	
QETMQCQR RF YEALHDPNLN EQRNARIRS IKDDC	35
SEQ ID NO: 268	moltype = AA length = 34
FEATURE	Location/Qualifiers
source	1..34
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 268	
QETQCQR RFY EALHDPNLNE EQRNARIRSI KDDC	34
SEQ ID NO: 269	moltype = AA length = 33
FEATURE	Location/Qualifiers
source	1..33
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 269	
QETCQR RFYE ALHDPNLNEE QRNARIRSIK DDC	33
SEQ ID NO: 270	moltype = AA length = 20
FEATURE	Location/Qualifiers
source	1..20
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 270	
QETRGNCA YH KGQLVWCTYH	20
SEQ ID NO: 271	moltype = AA length = 20
FEATURE	Location/Qualifiers
source	1..20
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 271	
QETRGNCA YH KGQIIWCTYH	20
SEQ ID NO: 272	moltype = AA length = 6
FEATURE	Location/Qualifiers
source	1..6
	mol_type = protein
	organism = synthetic construct
VARIANT	6
	note = A or P
SEQUENCE: 272	
LLQGPX	6
SEQ ID NO: 273	moltype = AA length = 8
FEATURE	Location/Qualifiers
source	1..8
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 273	
GGLLQGPP	8
SEQ ID NO: 274	moltype = length =
SEQUENCE: 274	
000	
SEQ ID NO: 275	moltype = length =
SEQUENCE: 275	
000	
SEQ ID NO: 276	moltype = length =

---

-continued

```

SEQUENCE: 276      moltype = length =
000

SEQ ID NO: 277      moltype = length =
SEQUENCE: 277
000

SEQ ID NO: 278      moltype = length =
SEQUENCE: 278
000

SEQ ID NO: 279      moltype = length =
SEQUENCE: 279
000

SEQ ID NO: 280      moltype = length =
SEQUENCE: 280
000

SEQ ID NO: 281      moltype = length =
SEQUENCE: 281
000

SEQ ID NO: 282      moltype = length =
SEQUENCE: 282
000

SEQ ID NO: 283      moltype = length =
SEQUENCE: 283
000

SEQ ID NO: 284      moltype = length =
SEQUENCE: 284
000

SEQ ID NO: 285      moltype = length =
SEQUENCE: 285
000

SEQ ID NO: 286      moltype = length =
SEQUENCE: 286
000

SEQ ID NO: 287      moltype = length =
SEQUENCE: 287
000

SEQ ID NO: 288      moltype = length =
SEQUENCE: 288
000

SEQ ID NO: 289      moltype = length =
SEQUENCE: 289
000

SEQ ID NO: 290      moltype = length =
SEQUENCE: 290
000

SEQ ID NO: 291      moltype = length =
SEQUENCE: 291
000

SEQ ID NO: 292      moltype = length =
SEQUENCE: 292
000

SEQ ID NO: 293      moltype = length =
SEQUENCE: 293
000

SEQ ID NO: 294      moltype = length =
SEQUENCE: 294
000

SEQ ID NO: 295      moltype = length =
SEQUENCE: 295
000

```

-continued

---

```

SEQ ID NO: 296      moltype =   length =
SEQUENCE: 296
000

SEQ ID NO: 297      moltype =   length =
SEQUENCE: 297
000

SEQ ID NO: 298      moltype =   length =
SEQUENCE: 298
000

SEQ ID NO: 299      moltype =   length =
SEQUENCE: 299
000

SEQ ID NO: 300      moltype =   length =
SEQUENCE: 300
000

SEQ ID NO: 301      moltype =   length =
SEQUENCE: 301
000

SEQ ID NO: 302      moltype =   length =
SEQUENCE: 302
000

SEQ ID NO: 303      moltype =   length =
SEQUENCE: 303
000

SEQ ID NO: 304      moltype =   length =
SEQUENCE: 304
000

SEQ ID NO: 305      moltype =   length =
SEQUENCE: 305
000

SEQ ID NO: 306      moltype =   length =
SEQUENCE: 306
000

SEQ ID NO: 307      moltype =   length =
SEQUENCE: 307
000

SEQ ID NO: 308      moltype =   length =
SEQUENCE: 308
000

SEQ ID NO: 309      moltype =   length =
SEQUENCE: 309
000

SEQ ID NO: 310      moltype =   length =
SEQUENCE: 310
000

SEQ ID NO: 311      moltype =   length =
SEQUENCE: 311
000

SEQ ID NO: 312      moltype =   length =
SEQUENCE: 312
000

SEQ ID NO: 313      moltype =   length =
SEQUENCE: 313
000

SEQ ID NO: 314      moltype =   length =
SEQUENCE: 314
000

SEQ ID NO: 315      moltype =   length =
SEQUENCE: 315
000

```

-continued

```

SEQ ID NO: 316      moltype =   length =
SEQUENCE: 316
000

SEQ ID NO: 317      moltype =   length =
SEQUENCE: 317
000

SEQ ID NO: 318      moltype =   length =
SEQUENCE: 318
000

SEQ ID NO: 319      moltype =   length =
SEQUENCE: 319
000

SEQ ID NO: 320      moltype =   length =
SEQUENCE: 320
000

SEQ ID NO: 321      moltype =   length =
SEQUENCE: 321
000

SEQ ID NO: 322      moltype =   length =
SEQUENCE: 322
000

SEQ ID NO: 323      moltype =   length =
SEQUENCE: 323
000

SEQ ID NO: 324      moltype =   length =
SEQUENCE: 324
000

SEQ ID NO: 325      moltype =   length =
SEQUENCE: 325
000

SEQ ID NO: 326      moltype =   length =
SEQUENCE: 326
000

SEQ ID NO: 327      moltype =   length =
SEQUENCE: 327
000

SEQ ID NO: 328      moltype =   length =
SEQUENCE: 328
000

SEQ ID NO: 329      moltype =   length =
SEQUENCE: 329
000

SEQ ID NO: 330      moltype = AA  length = 290
FEATURE
source
1..290
mol_type = protein
organism = Homo sapiens
SEQUENCE: 330
MRIFAVFIFM TYWHLLNAFT VTVPKDLYVV EYGSNMTIEC KFPVEKQLDL AALIVYWEME 60
DKNIIQFVHG EEDLKVQHSS YRQRARLLKD QLSLGNAALQ ITDVKLQDAG VYRCMISYGG 120
ADYKRITVKV NAPYNKINQR ILVVDPVITSE HELTCQARGY PKAEVIWITSS DHQVLSGKTT 180
TTNSKREEKL FNVTSRLRIN TTTNEIFYCT FRRLDPEENH TAEELVIPELP LAHPPNERTH 240
LVILGAILLC LGVALTFIFR LRKGRRMMMDVK KCGIQDTNSK KQSDTHLEET 290

SEQ ID NO: 331      moltype = AA  length = 288
FEATURE
source
1..288
mol_type = protein
organism = Homo sapiens
SEQUENCE: 331
MQIPQAPWPV VWAVALQLGWR PGWFLDSPDR PWNPPTFSPA LLVVTEGDN A TFTCSFSNTS 60
ESFVLNWRM SPSNQTDKLA AFPEDRSQPG QDCRFRVTOL PNGRDFHMSV VRARRNDSGT 120
YLCGAISLAP KAQIKESLRA ELRVTERRAE VPTAHPSPSP RPAGQFQTLV VGVVGGLGS 180
LVLLVWVLAV ICSRAARGTI GARRTGQPLK EDPSAVPVFS VDYGELEDFQW REKTPPEPPVP 240

```

-continued

CVPEQTEYAT IVFPGSMGTS SPARRGSADG PRSAQPLRPE DGHCSWPL 288

SEQ ID NO: 332 moltype = AA length = 445  
 FEATURE Location/Qualifiers  
 source 1..445  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 332  
 QVQLQESGPG LVKPSETLSL TCTVSGFSLT SYGVHWRQGP PGKGLEWIGV IYADGSTYN 60  
 PSLKSRVTIS KDTSKNQVSL KLSSVTAADT AVYYCARAYG NYWYIDVWGQ GTTVTVSSAS 120  
 TKGPSVFPFLA PCSRTSEST AALGCLVKD Y FPEPVTWSW N SGALTSQVHT FPAVLQSSGL 180  
 YSSLSSVTVP SSSLGTKTYT CNVDHKPSNT KVDKRVESKY GPPCPCCPAP PVAGGPVFL 240  
 FPPKPKDPLM ISRTPEVTCV VVAVSQEDPE VQFNWYVTDGV EVHNNAKTKPR EEEQFNSTYRV 300  
 VSVLTVVHQD WLNGKEYKCK VSNKGLPSSI EKTISKAKQG PREPVQVTLIP PSQEEMTKNQ 360  
 VSLTCLVKGF YPSDIAVEWE SNGOPENNYK TTTPVLDSDG SFFLYSKLTV DKSRWQEGNV 420  
 FSCSVMHEAL HNHYTQKSL SLSLGK 445

SEQ ID NO: 333 moltype = AA length = 214  
 FEATURE Location/Qualifiers  
 source 1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 333  
 DIVMTQSPDS LAVSLGERAT INCKSSESVS NDVAWYQQKP GQPPKLLINY AFHRFTGVPD 60  
 RFGSGSGYTD FTISSLQA EDVAVYYCHQ AYSSPYTFQG GTKLEIKRTV AAPSVFIFPP 120  
 SDEQLKSGTA SVVCLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYSLSSTLT 180  
 LSKADYEKHK VYACEVTHQG LSSPVTKSFN RGE 214

SEQ ID NO: 334 moltype = AA length = 447  
 FEATURE Location/Qualifiers  
 source 1..447  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 334  
 QVQLVQSGAE VKKPGSSVKV SCKASGGTFS SYAISWVRQA PGQGLEWMGL IIPMFDTAGY 60  
 AQKFQGRVAI TVDESTSTAY MELSSLRSED TAVYYCAREE HSSTGTFDYW GQGTLVTVSS 120  
 ASTKGPSVFP LAPCSRSTSE STAALGCLVK DYPPEPVTVS WNSGALTSGV HTFPAVLQSS 180  
 GLYSLSSVPT VPSSSLGTKT YTCNVDHKPS NTKVDKRVES KYGPPCPCCP APEFLGGPSV 240  
 FLFPPKPKD LMISRTPEVTT C VVVDVQWQED PEVQFNWYVGD GVEVHNNAKTK PREEQFNSTY 300  
 RVVSVLTVLH QDWLNGKEYK CKVSNKGLPS SIEKTISKAK GQPREPQVYT LPPSQEEMTK 360  
 NQVSLTCLVK GFYPSDIAVE WESNGOPENN YKTTPPVLDG DGSFFLYSRL TVDKSRWQEG 420  
 NVFSCSVMHE ALHNHYTQKS LSLSLGK 447

SEQ ID NO: 335 moltype = AA length = 214  
 FEATURE Location/Qualifiers  
 source 1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 335  
 DIQMTQSPSS VSASVGDRVT ITCRASQGIS SWLAWYQQKP GKAPKLLISA ASSLQSGVP 60  
 RFGSGSGYTD FTISSLQP EDFATYYCQQ ANHLPFTFGG GTKVEIKRTV AAPSVFIFPP 120  
 SDEQLKSGTA SVVCLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYSLSSTLT 180  
 LSKADYEKHK VYACEVTHQG LSSPVTKSFN RGE 214

SEQ ID NO: 336 moltype = AA length = 452  
 FEATURE Location/Qualifiers  
 source 1..452  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 336  
 QGQLVQSGAE VKKPGASVKV SCKASGYTFT DYEMHWVRQA PIHGLEWIGV IESETGGTAY 60  
 NQKFQGRVTI TADKSTSTAY MELSSLRSED TAVYYCAREG ITTVATTYYW YFDVWQGQTT 120  
 VTVSSASTKG PSVFLAPCS RSTSESTAAL GCLVKDYPFPE PVTVSWNSGA LTSGVHTFP 180  
 VIQQSSGSLV SSVVTPVSS LGTKTYTCMV DHKPSNTKVD KRVESKYGPP CPPCPAPEF 240  
 GGPSVFLFPP KPKDTLMISR TPEVTCVVVD VSQEDPEVQF NWYVDGVEVH NAKTKPREEQ 300  
 FNSTYRVVSV LTVLHQDWLN GKEYKCKVSN KGLPSSIEKTT ISKAKGQPRE PQVYTLPPSQ 360  
 EEMTKNQVSL TCLVKGFYPS DIAVEWESNG QPENNYKTTP PVLDSDGSFF LYSRLTVDKS 420  
 RWQEGNFS SVMHEALHNH YTQKSLSL SLSLGK 452

SEQ ID NO: 337 moltype = AA length = 219  
 FEATURE Location/Qualifiers  
 source 1..219  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 337  
 DVVMTQSPLS LPVTLGQPAS ISCRSSQSIV HSNGNTYLEW YLQKPGQSPQ LLIYKVSNRF 60  
 SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCFQGSHVP LTFGQGKLE IKRTVAAPSV 120  
 FIFPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALOQ SGNSQESVTE QDSKDSTYSL 180  
 SSSLTLSKAD YEKHKVYACE VTHQGLSSPV TKSFNRGEC 219

-continued

-continued

---

GLYSLSSVVT VPSSSLGTTK YTCNVDHKPS NTKVDKRVES KYGPPCPCCP APEFLGGPSV 240  
 LFPPPKPKDT LMISRTPEVT CVVVDSQED PEVQFNWYVD GVEVHNNAKTK PREEQFNSTY 300  
 RVVSVLTVLH QDWLNGKEYK CKVSNKGLPS SIEKTISKAK GQPREPQVYT LPPSQEEMTK 360  
 NQVSLTCLVK GFYPSDIAVE WESNGQPENN YTTPPVLDs DGSFFLYSRL TVDKSRWQEG 420  
 NVFSCSVMHE ALHNHYTQKS LSLSLGK 447

SEQ ID NO: 347 moltype = AA length = 218  
 FEATURE Location/Qualifiers  
 source 1..218  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 347  
 EIVLTQSPAT LSLSPGERAT LSCRASKGVs TSGSYLHWY QQKPGQAPRL LIYLASYLES 60  
 GVPARFSGSG SGTDFTLTIS SLEPEDFAVY YCQHSRDLPL TFGGGTKEI KRTVAAPSVF 120  
 IFFPSDQEQLK SGTASVVCll NNFYPREAKV QWKVDNALQs GNSQESVTEQ DSKDSTYSLs 180  
 STLTLSKADY EKHKVYACEV THQGLSSPVT KSFNRGEC 218

SEQ ID NO: 348 moltype = AA length = 166  
 FEATURE Location/Qualifiers  
 source 1..166  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 348  
 QVQLVQSGVE VKKPGASVVK SCKASGYTFT NYMYWVRQA PGQGLEWMGG FPSNGGTNFN 60  
 EKPKNRVTLT DSSTTAYM ELKSLQFDTT AVYYCARRDY RFDMGFDYWG QGTTTVSSD 120  
 ISULFIDEBR IDGEHHHLHH HHHHHHHHHHL LLLLLL 166

SEQ ID NO: 349 moltype = AA length = 111  
 FEATURE Location/Qualifiers  
 source 1..111  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 349  
 EIVLTQSPAT LSLSPGERAT LSCRASKGVs TSGSYLHWY QQKPGQAPRL LIYLASYLES 60  
 GVPARFSGSG SGTDFTLTIS SLEPEDFAVY YCQHSRDLPL TFGGGTKEI K 111

SEQ ID NO: 350 moltype = AA length = 113  
 FEATURE Location/Qualifiers  
 source 1..113  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 350  
 QVQLVESGGG VVQPGRSRL DCKASGITFS NSGMHWVRQA PGKGLEWVAV IWYDGSKRYY 60  
 ADSVVKGRFTI SRDNNSKNTLF LQMNSLRAED TAVYYCATND DYWGQGLVLT VSS 113

SEQ ID NO: 351 moltype = AA length = 154  
 FEATURE Location/Qualifiers  
 source 1..154  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 351  
 EIVLTQSPAT LSLSPGERAT LSCRASQSvS SYLAWYQQKP GQAPRLLIYD ASN RATGIPA 60  
 RFSGSGSGTD FTLTISSELP EDFAVYYCQQ SSNWPRTFGQ GTKVEIKDIS ULFIDEBRID 120  
 GEHHHLHHHH HHHHHHHHHHL HHHHHHHLLL LLLL 154

SEQ ID NO: 352 moltype = AA length = 459  
 FEATURE Location/Qualifiers  
 source 1..459  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 352  
 QVQLVQSGGG LVQPGSSLRL SCAASGFTFS SYWMMYWRQV PGKGLEWVSA IDTGGGRTY 60  
 ADSVVKGRFAI SRVNAKNTMY LQMNSLRAED TAVYYCARDE GGTTGWGVLK DWPYGLDAWG 120  
 QGTLVTVSSA STKGPSVFPL APSSKSTSGG TAALGCLVKD YFPEPVTVSW NSGALTSGVH 180  
 TFPAVLQSSG LYSLSSVVTv PSSSLGTQTY ICNVNHKPSN TKVDRKVERPK SCDKHTCPP 240  
 CPAPEAAGGP SVFLFPKPK DTLMSRTPE VTCVVDVSH EDPEVKENWY VDGVEVHNAAK 300  
 TKPREEQYNS TYRVSVLTV LHQDWLNGKE YKCKVSNKAL PAPIETISK AKGQPREPQV 360  
 YTLPPSREEM TKNQVSLTCL VKGFYPSDIA VEWESENQPE NNYKTTPPVL DSDGSFFLYS 420  
 KLTVDKSRWQ QGNVFSCSVm HEALHNHYTQ KSLSLSPGK 459

SEQ ID NO: 353 moltype = AA length = 214  
 FEATURE Location/Qualifiers  
 source 1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 353  
 QPVLTQPLSV SVALGQTARI TCGGNNIGSK NVHWYQQKPG QAPVLIYRD SNRPSGIPER 60  
 FSGSNNSGNTA TLTISRAQAG DEADYYCQW DSSTAVFGTG TKLTVLQRTV AAPSVFIFPP 120  
 SDBQLKSGTA SVVCLLNNFY PREAKVQWVKA DNALQSGNSQ ESVTEQDSKD STYSLSTLT 180

-continued

---

LSKADYEKHK VYACEVTHQG LSSPVTKSFN RGE<sup>C</sup> 214

SEQ ID NO: 354 moltype = AA length = 439  
 FEATURE Location/Qualifiers  
 source 1..439  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 354  
 QVQLVESGGG VVQPGSRL SCAASGFTFS SYGMHWRQQA PGKGLEWVAV IWYDGSNKYY 60  
 ADSVKGRFTI SRDNSKNTLY LQMNSLRAED TAVYYCASNG DHWGQGLTVL VSSASTKGPS 120  
 VFPLAPCRS TSESTAALGG LVKDYFPEPV TVSWNSGALT SGVHTFPVAL QSSGLYSLSS 180  
 VVTVPSSSLG TKTYTCNVDH KPSNTKVDKR VESKYGPPC PCPAPEFLGG PSVFLFPPKP 240  
 KDTLMISRTV EVTCVVVDVQ QEDPEVQFW YVDGVEVHNA KTKPREEQFW STYRVSVLT 300  
 VLHQDWLNGK LPSSIEKTIS KAKGQPREPQ VYTLPSSQEE MTKNQVSLLTC 360  
 LVKGFYPSDI AVEWESNGQP ENNYKTTPPV LDSDGSFFLY SRLTVDKSRW QEGNFSCSV 420  
 MHEALHNHYT QKSLSLSG 439

SEQ ID NO: 355 moltype = AA length = 214  
 FEATURE Location/Qualifiers  
 source 1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 355  
 EIVMTQSPAT LSVSPGERAT LSCRASQSVS SNLAWYQQKP GQAPRLLIYG ASTRATGIPA 60  
 RFGSGSGSTE FTLTISSLQS EDFAVYYCQQ YNNWPRTEFGQ GTKVEIKRTV AAPSVFIFPP 120  
 SDEQLKSGTA SVVCLLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYSLSSTLT 180  
 LSKADYEKHK VYACEVTHQG LSSPVTKSFN RGE<sup>C</sup> 214

SEQ ID NO: 356 moltype = AA length = 443  
 FEATURE Location/Qualifiers  
 source 1..443  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 356  
 EVQLLESGGG LVQPGGSLRL SCAASGFTFS SYDMMSWVRQA PGKGLEWVST ISGGGSYTYY 60  
 QDSVKGRFTI SRDNSKNTLY LQMNSLRAED TAVYYCASPY YAMDYWQGQT TVTSSASTK 120  
 GPSVFP LAPC SRSTSESTAA LGCLVVKDYFP EPVTVWSNNG ALTSGVHTFP AVLQSSGLYS 180  
 LSSVTVTPSS SLGTKTYTCN VDHKPSNTKV DKRVESKYGP PCPPCPAPEF LGGPSVFLPP 240  
 PKPKDTLMIS RTPEVTCVV DVSQEDPEVQ FNWYVDGVEV HNAKTKPREE QFNSTYRVVS 300  
 VLTVLHQDWL NGKEYKCKVS NKGLPSSIEK TISKAKGQPR EPQVYTLPPS QEEMTKNQVS 360  
 LTCLVKGFYPSDI AVEWESN GQPENNYKTT PPVLSDGSF FLYSRLTVDK SRWQEGNVFS 420  
 CSVMHEALHN HYTQKSLSLS LGK 443

SEQ ID NO: 357 moltype = AA length = 214  
 FEATURE Location/Qualifiers  
 source 1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 357  
 DIQLTQSPSF LSAYVGDRVT ITCKASQDVG TAVAWYQQKP GKAPKLLIYW ASTLHTGVPS 60  
 RFGSGSGSTE FTLTISSLQP EDFATYYCQH YSSYPWTFGQ GTKLEIKRTV AAPSVFIFPP 120  
 SDEQLKSGTA SVVCLLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYSLSSTLT 180  
 LSKADYEKHK VYACEVTHQG LSSPVTKSFN RGE<sup>C</sup> 214

SEQ ID NO: 358 moltype = AA length = 443  
 FEATURE Location/Qualifiers  
 source 1..443  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 358  
 QVQLVESGGG LVKPGGSLRL SCAASGFTFS NYGMSWIROA PGKGLEWSTI SGGSNIYYA 60  
 DSVKGRFTI RDNAKNSLYL QMNSLRAEDT AVYVCVSYY GIDFWQGTS VTVSSASKYG 120  
 PSVFP LAPCS RSTSESTAA GLC1VVKDYFP PTVWSNNGA LTSGVHTFPV VLQSSGLYSL 180  
 SSVTVTPSS LGTAKTYTCN DHKPSNTKV KRVESKYGP PCPPCPAPEF LGGPSVFLPP 240  
 KPKDTLMISRTV PEVTCVVVD VSQEDPEVQF NWYVDGVEV NAKTKPREEQ FNSTYRVSV 300  
 VLTVLHQDWL NGKEYKCKVS NKGLPSSIEK TISKAKGQPR EPQVYTLPPS QEEMTKNQVS 360  
 LTCLVKGFYPSDI AVEWESN GQPENNYKTT PPVLSDGSF FLYSRLTVDK SRWQEGNVFS 420  
 CSVMHEALHN HYTQKSLSLS LGK 443

SEQ ID NO: 359 moltype = AA length = 214  
 FEATURE Location/Qualifiers  
 source 1..214  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 359  
 DIQMTQSPSS LSASVGDRVT ITCKASQDV TAVAWYQQKP GKAPKLLIYW ASTRHTGVPS 60  
 RFGSGSGSTD FTLTISSLQP EDFATYYCQH HYTIPWTFGG GTKLEIKRTV AAPSVFIFPP 120  
 SDEQLKSGTA SVVCLLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYSLSSTLT 180  
 LSKADYEKHK VYACEVTHQG LSSPVTKSFN RGE<sup>C</sup> 214

-continued

SEQ ID NO: 360      moltype = AA length = 445  
 FEATURE                Location/Qualifiers  
 source                1..445  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 360  
 QVQLVQSGAE VKKPGASVKV SCKASGYSFT SYWMNWRQQA PGQGLEWIGV IHPSDSETWL 60  
 DQKFKNRTVI TVDKSTSTAY MELSSLRSED TAVYYCAREH YGTSPFAYWG QGTLTVSSA 120  
 STKGPSVPLS ACPSRSTSES TAALGCLVKD YFPEPVTVSW NSGALTSGVH TFPAVLQSSG 180  
 LYSLSSVVTV FSSSLGKTKV TCNDHKPSN TKVDKRVESK YGPPCPCCPA PEFLGGPSVF 240  
 LFPPPKDTL MISRTPEVTC VVVDVSQEDP EVQFNWYVG E VEHNAKTKP REEQFNSTYR 300  
 VVSVLTVLHQ DWLNGKEYKC KVSNKGLPSS IEKTISAKG QPREPVYTT PPSQEEMTKN 360  
 QVSLTCLVKG FYPSDIAVEW ESNGOPENNY KTPPPVLDSD GSFFLYSRLT VDKSRWQEGN 420  
 VFSCSVMHEA LHNTQKSL SLSLG    445

SEQ ID NO: 361      moltype = AA length = 218  
 FEATURE                Location/Qualifiers  
 source                1..218  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 361  
 EIVLTQSPAT LSLSPGERAT LSCRASESVD NYGMSFMNW QQKPGQPPKL LIHAASNQGS 60  
 GVPVSRFGSG SGTDFTLTIS SLEPEDFAVY FCQQSKEVPY TFGGGTKVEI KRTVAAPS 120  
 IFPPSDEQLK SGTASVVCLL NNFYPREAKV QWKVDNALQGS GNSQESVTEQ DSKDSTYSL 180  
 STLTLSKADY EKHKVYACEV THQGLSSPVT KSFNRGEC    218

SEQ ID NO: 362      moltype = AA length = 444  
 FEATURE                Location/Qualifiers  
 source                1..444  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 362  
 QVQLVQSGAE VKKPGASVKV SCKASGYTFT SYWINWRQQA PGQGLEWMGN IYPGSSLTN 60  
 NEKKFKNRVTM TRDTSTSTVY MELSSLRSED TAVYYCARLS TGTFAYWGQ TLTVSSAST 120  
 KGPSVFPLAP CSRSTSESTA ALGCLVKDYF PEPVTVWSNS GALTSGVHTF PAVLQSSGLY 180  
 SLSSVVTVPS SSLGKTKYTC NVDHKPSNTK VDKRVESKYG PPCPPCPAPE FLGGPSVFLF 240  
 PPKPKDTLMI SRTPEVTCVV VDVSQEDPEV QFNWYVG E VHNAAKTKP RE EQFNSTYR 300  
 SVLTVLHQDW LNGKEYKCKV SNKGLPSSIE KTISAKGQ P REPQVYTLPP SQEEMTKNQV 360  
 SLTCLVKGFY PSDIAVEWES NGQPENNYKT TPPVLDSDGS FFLYSLRTVD KSRWQEGNVF 420  
 SCSTMHEALH NHYTQKSL SLSLG    444

SEQ ID NO: 363      moltype = AA length = 220  
 FEATURE                Location/Qualifiers  
 source                1..220  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 363  
 DIVMTQSPDS LAVSLGERAT INCKSSQSLW DSGNQKNFLT WYQQKPGQPP KLLIYWTSYR 60  
 ESGVPRDFSG SGSGTDFTLT ISSLQAEDVA VYYCQNDYFY PHTFGGGTKV EIKRTVAAPS 120  
 VFIFPPSDEQ LKSGTASVVC LLNNFYPREA KVQWKVDNAL QSGNSQESTV EQDSKDSTYS 180  
 LSSTTLSKA DYEKHKVYAC EVTHQGLSSP VTKSFRNRGEC    220

SEQ ID NO: 364      moltype = AA length = 443  
 FEATURE                Location/Qualifiers  
 source                1..443  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 364  
 EVQLVQSGAE VKKPGESLRI SCKGSGYTFT TYWMHWVRQA TGQGLEWMGN IYPGTGGSNF 60  
 DEKFKNRTVI TADKSTSTAY MELSSLRSED TAVYYCTRWT TGTGAYWGQ TT TVVSSAST 120  
 KGPSVFPLAP CSRSTSESTA ALGCLVKDYF PEPVTVWSNS GALTSGVHTF PAVLQSSGLY 180  
 SLSSVVTVPS SSLGKTKYTC NVDHKPSNTK VDKRVESKYG PPCPPCPAPE FLGGPSVFLF 240  
 PPKPKDTLMI SRTPEVTCVV DVDSQEDPEV QFNWYVG E VHNAAKTKP RE EQFNSTYR 300  
 SVLTVLHQDW LNGKEYKCKV SNKGLPSSIE KTISAKGQ P REPQVYTLPP SQEEMTKNQV 360  
 SLTCLVKGFY PSDIAVEWES NGQPENNYKT TPPVLDSDGS FFLYSLRTVD KSRWQEGNVF 420  
 SCSTMHEALH NHYTQKSL SLSLG    443

SEQ ID NO: 365      moltype = AA length = 220  
 FEATURE                Location/Qualifiers  
 source                1..220  
 mol\_type = protein  
 organism = synthetic construct

SEQUENCE: 365  
 EIVLTQSPAT LSLSPGERAT LSCKSSQSLW DSGNQKNFLT WYQQKPGQAP RLLIYWA STR 60  
 ESGVPSRFSG SGSGTDFFT ISSLQAEDAA VYYCQNDYSY PYTFGGQGTKV EIKRTVAAPS 120  
 VFIFPPSDEQ LKSGTASVVC LLNNFYPREA KVQWKVDNAL QSGNSQESTV EQDSKDSTYS 180  
 LSSTTLSKA DYEKHKVYAC EVTHQGLSSP VTKSFRNRGEC    220

-continued

---

SEQ ID NO: 366	moltype = AA length = 450
FEATURE	Location/Qualifiers
source	1..450
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 366	
QVQLVQSGAE VKKPGSSVKV SCKASGGTFS SYAISWVRQA PGQGLEWMGG IIPIDTANY	60
AQKFQGRVTI TADESTSTAY MELSSLRSED TAVYYCARPG LAAAYDTGSL DYWGQGTLVT	120
VSSASTKGPS VPPLAPCSRS TSESTAALGC LVKDYFPPEPV TVSWNSGALT SGVHTFPVAL	180
QSSGLYSSLKP VFTVTPSSLLG TKTYTCNVDH KPSNTKVDKR VESKYGPPCP PCPAPEFLGG	240
PSVFLFPKPKD KDTLMISRTP EVTCVVVDWS QDPFVQFWN VYDGVEVHNA KTKPREEQFN	300
STYRVSVLTL VHLDWLNKG EYKCKVSNKG LPSSIEKTIS KAKGQPREPQ VYTLPPSQEE	360
MTKNQVSLLTC LVKGFYPSDI AVEWESNGQP ENNYKTTTPV LDSDGSFFLY SRLTVDKSRW	420
QEGNQVFSCSV MHEALHNHYT QKSLSLSLGK	450
SEQ ID NO: 367	moltype = AA length = 214
FEATURE	Location/Qualifiers
source	1..214
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 367	
EIVLTQSPAT LSLSPGERAT LSCRASQSVR SYLAWSQQKP GQAPRLLIYD ASNRTGIPA	60
RFSGSGSGTD FTLTISSLEP EDFAVYYCQQ RNYWPLTFQG GTKVEIKRTV AAPSVFIFPP	120
SDEQLKSGTA SVVCLLNNFY PREAKVQWKV DNALQSGNSQ ESVTEQDSKD STYSLSSTLT	180
LSKADYEKHKK VYACEVTHQG LSSPVTKSFN RGECA	214
SEQ ID NO: 368	moltype = AA length = 496
FEATURE	Location/Qualifiers
source	1..496
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 368	
DIGMTQSPSS LSASVGDRVT ITCRASQDV SVVAWYQQKP GKAPKLLIYS ASYRYTGVP	60
RFSGSGSGTD FTLSITSSLQF EDFATYVYQQ HYPSTPWTFGG GTKLEIKGGG SGGGQVQLV	120
QSGAEVKGPK AVSKVSKAS GYSFTSYWMN WVRQAPGGL EWIGVHIPS D SETWLQDKF	180
DRVITITVDKS TSTAYMELSS LRSEDTAVYY CAREHYGTSP FAYWGQGTLV TVSSGGCGGG	240
EVAAACEKEVA ALEKEVAALE KEVAALEKES KYGPPCPCCP APEFLGGPSV FLFPKPKDT	300
LYITREPEVTT CVVVDSQED PEVQFNWYVD GVEVHNNAKTK PREEQFNSTY RVVSVLTVLH	360
QDWLNGKEYK CKVSNKGKLP SIEKTISKAK GQPREPVQVYT LPPSQEEMTK NQVSLTCLVK	420
GFYPSDIAVE WESNGQOPENN YKTTPPVLDs DGSSFFLYSRL TVDKSRWQEG NVFSCSVMHE	480
ALHNHYTQKS LSLSLG	496
SEQ ID NO: 369	moltype = AA length = 271
FEATURE	Location/Qualifiers
source	1..271
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 369	
EIVLTQSPAT LSLSPGERAT LSCRASESVD NYGMSFMNW QQKPGQPPKL LIHAASNQGS	60
GVPSRSGSG SGTDFTLTIS SLEPDTFVY FCQQSKEVPTY TFGGGTKVEI KGGGSGGGQ	120
VQVLVQSGAEV KRPKGASVKS CKASGYTFTD YNMDWVRQAP GQGLEWMGDI NPDNGVTIYN	180
QKPEGRVTMT TDTSTSTAYM ELRSLRSDDT AVYYCAREAD YFYFDYWGQG TTLTVSSGGC	240
GGGKVAACKE KVAALKVA ALKEKVAALK E	271
SEQ ID NO: 370	moltype = AA length = 447
FEATURE	Location/Qualifiers
source	1..447
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 370	
QVQLVQSGSE LKKPGASVKA SCKASGYTFT NYGMNWRQA PGQGLQWMGW INTDSGESTY	60
AEEFKGRFVFL SLDTSVNTAY LQITSLTAED TGMYFCVRVG YDALDXYWQG TLTVTSSAST	120
KGPSVFPLAP SSKSTVSGGA ALGCLVYDFY PEPVTWSNNS GALTSVGHTF PAVLQSSGLY	180
SLSVVTVTPS SSLGQTQTYIC VNHNKPSNTK VDKRVEPKSC DKTHTCPCCP APPELLGGPSV	240
FLFPPKPKDT LMISRTPEVT CVVVDSHED PEVQFNWYVD GVEVHNNAKTK PREEQFNSTY	300
RVVSVLTVLH QDWLNGKEYK CKVSNKALPA PIEKTISKAK GQPREPVQVYT LPPSRREEMTK	360
NQVSLTCLVK GFYPSDIAVE WESNGQOPENN YKTTPPVLDs DGSSFFLYSRL TVDKSRWQOG	420
NVFSCSVMHE ALHNHYTQKS LSLSPGK	447
SEQ ID NO: 371	moltype = AA length = 213
FEATURE	Location/Qualifiers
source	1..213
	mol_type = protein
	organism = synthetic construct
SEQUENCE: 371	
EIVLTQSPSS LSASVGDRVT ITCSARSSVS YMHWFFQQKPG KAPKLWIYRT SNLASGVPSR	60
FSFGSGSGTSY CLTINSLQPE DFATYYCQQR SSFPLTFGGG TKLEIKRTVA APSVFIFPPS	120
DEQLKSGTAS SVVCLLNNFY REAKVQWKV DALQSGNSQES VTQEQDSKD STYSLSSTLT	180
SKADYEKHKK VYACEVTHQGL SSPVTKSFNR GEC	213

---

-continued

SEQ ID NO: 372 moltype = AA length = 113  
 FEATURE Location/Qualifiers  
 source 1..113  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 372  
 QVQLVESGGG VVQPGRSLRL TCKASGLTFS SSGMHWVRQAA PGKGLEWVAV IWYDGSKRYY 60  
 ADSVKGRFTI SRDN SKNTLF LQMNSLRAED TAVYYCATNN DYWGQQTLVT VSS 113

SEQ ID NO: 373 moltype = AA length = 107  
 FEATURE Location/Qualifiers  
 source 1..107  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 373  
 EIVLTQSPAT LSLSPGERAT LSCRASQSVS SYLAWYQQKP QQAPRLLIYT ASN RATGIPA 60  
 RFSGSGSGTD FTLTIS SLEP YSNWPRTFGQ GTKVEIK 107

SEQ ID NO: 374 moltype = AA length = 120  
 FEATURE Location/Qualifiers  
 source 1..120  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 374  
 EVQLQQSGPV LVKPGASVKM SCKASGYTFT SYYMYWVKQS HGKSLEWIGG VNPSNGGTNF 60  
 NEKFKSKATL TVDKSSSTAY MELNSLTSED SAVYYCAR RD YRYDMGFDYW GQGTTLVSS 120

SEQ ID NO: 375 moltype = AA length = 111  
 FEATURE Location/Qualifiers  
 source 1..111  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 375  
 QIVLTQSPAI MSASPGEKVT MTCRASKGV S TSGSYLHWY QQKPGSSPR L IYLASYLES 60  
 GVPVRFSGSG SGT SYSLTIS RMEAEDAATY YCQHSRELPL TFGTGTRLEI K 111

SEQ ID NO: 376 moltype = AA length = 120  
 FEATURE Location/Qualifiers  
 source 1..120  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 376  
 QVQLVQSGAE VKKPGASVKV SCKASGYTFT SYYMYWVRQAA PGQGLEWMGG VNPSNGGTNF 60  
 NEKFKSRVTI TADKSTSTAY MELSSLRSED TAVYYCAR RD YRYDMGFDYW GQGTTTVSS 120

SEQ ID NO: 377 moltype = AA length = 111  
 FEATURE Location/Qualifiers  
 source 1..111  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 377  
 EIVLTQSPAT LSLSPGERAT ISCRASKGV S TSGSYLHWY QQKPGQAPRL LIYLASYLES 60  
 GVPARFSGSG SGT DFTLTIS SLEPEDFAT YCQHSRELPL TFGTGTRLEI K 111

SEQ ID NO: 378 moltype = AA length = 448  
 FEATURE Location/Qualifiers  
 source 1..448  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 378  
 EIQLVQSGAE VKKPGSSVKV SCKASGYTFT HYGMNHWVRQAA PGQGLEWVGW VNTYTGEPY 60  
 ADDFKGRLTF TLD TSTSTAY MELSSLRSED TAVYYCTREG EGLGFGDWGQ GTTVTVSSAS 120  
 TKGPSVFP LA PSSKSTSGGT AALGCLVKD Y FPEPVTVSWN SGALTSGVHT FPAVLQSSGL 180  
 YSLSSVVTV P SSLGTQTYI CNVNHKPSNT KVDKKV EP KCS CDKTHTCP C PAPEAAGGPS 240  
 VFLFPPPKD TL MIRSTPEV TCVVV DVSH DPEVKFNWVY DGVEVHNAAKT KPREEQYNST 300  
 YRVVSVLTVL HQDWLNGKEY KCKVSNKALP APIEKTISKA KGQPREPQVY TLPPSREEMT 360  
 KNQVSLTCLV KGFPYPSDI AV EWESNGQPEN NYKTTPPVLD SDGSFFLYSK LTVDKSRWQQ 420  
 GNVFSCSVMH EALHNHYTQK SLSLSPKG 448

SEQ ID NO: 379 moltype = AA length = 219  
 FEATURE Location/Qualifiers  
 source 1..219  
 mol\_type = protein  
 organism = synthetic construct  
 SEQUENCE: 379  
 DVVMTQSPLS LPVTPGEPAS ISCRSSQSIV HSHGDTYLEW YLQKPGQSPQ LLIYKVSNRF 60  
 SGVPDRFSGS GSGTDFTLKI SRVEAEDVGV YYCFQGSHIP VTFGQQTKLE IKRTVAAPSV 120  
 FIFPPSDEQL KSGTASVVCL LNNFYPREAK VQWKVDNALQ SGNSQESVTE QDSKDSTYSL 180

-continued

SSTLTLSKAD YEKHKVYACE VTHQGLSSPV TKSFNRGEC	219
SEQ ID NO: 380                    moltype = AA length = 5	
FEATURE                         Location/Qualifiers	
source                         1..5	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 380	
NYYMY	5
SEQ ID NO: 381                    moltype = AA length = 17	
FEATURE                         Location/Qualifiers	
source                         1..17	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 381	
GINPSNGGTN PNEKFKN	17
SEQ ID NO: 382                    moltype = AA length = 11	
FEATURE                         Location/Qualifiers	
source                         1..11	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 382	
RDYRFDMGFD Y	11
SEQ ID NO: 383                    moltype = AA length = 15	
FEATURE                         Location/Qualifiers	
source                         1..15	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 383	
RASKGVSTSG YSYLH	15
SEQ ID NO: 384                    moltype = AA length = 7	
FEATURE                         Location/Qualifiers	
source                         1..7	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 384	
LASYLES	7
SEQ ID NO: 385                    moltype = AA length = 9	
FEATURE                         Location/Qualifiers	
source                         1..9	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 385	
QHSRDLPLT	9
SEQ ID NO: 386                    moltype = AA length = 5	
FEATURE                         Location/Qualifiers	
source                         1..5	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 386	
NSGMH	5
SEQ ID NO: 387                    moltype = AA length = 17	
FEATURE                         Location/Qualifiers	
source                         1..17	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 387	
VIWYDGSKRY YADSVKG	17
SEQ ID NO: 388                    moltype = AA length = 4	
FEATURE                         Location/Qualifiers	
source                         1..4	
mol_type = protein	
organism = synthetic construct	
SEQUENCE: 388	
NDDY	4
SEQ ID NO: 389                    moltype = AA length = 11	
FEATURE                         Location/Qualifiers	
source                         1..11	
mol_type = protein	
organism = synthetic construct	

-continued

---

SEQUENCE: 389	RASQSVSSYL A	11
SEQ ID NO: 390	moltype = AA length = 7 FEATURE source Location/Qualifiers 1..7 mol_type = protein organism = synthetic construct	
SEQUENCE: 390	DASN RAT	7
SEQ ID NO: 391	moltype = AA length = 9 FEATURE source Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 391	QQSSNWPRT	9
SEQ ID NO: 392	moltype = AA length = 9 FEATURE source Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 392	FTFSNYGMS	9
SEQ ID NO: 393	moltype = AA length = 10 FEATURE source Location/Qualifiers 1..10 mol_type = protein organism = synthetic construct	
SEQUENCE: 393	TISGGGSNIY	10
SEQ ID NO: 394	moltype = AA length = 9 FEATURE source Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 394	VSYYYGIDF	9
SEQ ID NO: 395	moltype = AA length = 11 FEATURE source Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 395	KASQDVTTAV A	11
SEQ ID NO: 396	moltype = AA length = 7 FEATURE source Location/Qualifiers 1..7 mol_type = protein organism = synthetic construct	
SEQUENCE: 396	WASTRHT	7
SEQ ID NO: 397	moltype = AA length = 9 FEATURE source Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 397	QQHYTIPWT	9
SEQ ID NO: 398	moltype = AA length = 8 FEATURE source Location/Qualifiers 1..8 mol_type = protein organism = synthetic construct	
SEQUENCE: 398	GLTFSSSG	8
SEQ ID NO: 399	moltype = AA length = 8 FEATURE source Location/Qualifiers 1..8 mol_type = protein	

-continued

---

	organism = synthetic construct	
SEQUENCE: 399 IWYDGSKR		8
SEQ ID NO: 400 FEATURE source	moltype = AA length = 6 Location/Qualifiers 1..6 mol_type = protein organism = synthetic construct	
SEQUENCE: 400 ATNNDY		6
SEQ ID NO: 401 FEATURE source	moltype = AA length = 11 Location/Qualifiers 1..11 mol_type = protein organism = synthetic construct	
SEQUENCE: 401 RASQSVSSYL A		11
SEQ ID NO: 402 FEATURE source	moltype = AA length = 7 Location/Qualifiers 1..7 mol_type = protein organism = synthetic construct	
SEQUENCE: 402 TASN RAT		7
SEQ ID NO: 403 FEATURE source	moltype = AA length = 9 Location/Qualifiers 1..9 mol_type = protein organism = synthetic construct	
SEQUENCE: 403 QQYSNWPRT		9
SEQ ID NO: 404 FEATURE source	moltype = AA length = 831 Location/Qualifiers 1..831 mol_type = protein organism = synthetic construct	
SEQUENCE: 404 MQIPQAPWPW WAVLQLGWWRP GWFLDSPDRP WNPPTFSPAL LVVTEGDNAT FTCSFSNTSE 60 SFVLNWYRMS PSNQTDKLA FPEDRSQPGQ DCRFRVTQLP NGRDFHMSVV RARRNDSGTY 120 LCGAISLAPK AQIKESLRAE LRVTERRAEW PTAHPSPSPR PAGQFQS KYG PPCPSCPAP 180 FLGGPSVFLF PPKPKDLM SRTPEVTCWV DVSQEDPEVQ FNWYVVDGVEV HNAKTKPREE 240 QFNSTYRVVS VLTVLHQDWL SGKEYKCKVNS SKGLPSSIEK TISNATGQPR EPQVYTLPPS 300 QEEMTKNQVS LTCLVKGFYP SDIAVEWESN GQPENNYKTT PPVLDSDGSF FLYSRLTVDK 360 SSWQEGNVFS CSVHMHEALHN HYTQKSLSL LGKIEGRMDQ VSHRYPRIQS IKVQFTEYKK 420 EKGFIILTSQK EDEIMKVQNN SVIINCDGFY LISLKGYFSQ EVNISLHYQK DEEPLFQLKK 480 VRSVNSLMVA SLTYDKVYL NVTTDNTSLD DFHVNGGELI LIHQNPGEFC VLMQIPQAPW 540 PWWAVLQLGW RRGWFLDSPD RPWNPTFSP ALLVVTEGDN ATFTCSFSNT SESFVLNWYR 600 MSPSNQTDKL AAFPEDRSQP QGDCCRFRVTQ LPNGRDFHMS VVRARRNDSG TYLCGAISLA 660 PKAQIKEYSLR AELRVTERRA EVPTAHPSPS PRPAGQFQCV SHRYPRIQS K VQFTEYKK 720 KGFIILTSQK DEIMKVQNN VIINCDGFY ISLKGYFSQ EVNISLHYQK DEEPLFQLKK 780 RSVNSLMVAS LTYDKVYL VTTDNTSLD FHVNNGGELI IHQNPGEFCV L 831		
SEQ ID NO: 405 FEATURE source	moltype = AA length = 5 Location/Qualifiers 1..5 mol_type = protein organism = synthetic construct	
SEQUENCE: 405 GGGGS		5
SEQ ID NO: 406 FEATURE source	moltype = AA length = 17 Location/Qualifiers 1..17 mol_type = protein organism = synthetic construct	
SITE	1 note = Acetylated residue	
SITE	8 note = Lysine-3-((3-((2,5-dioxopyrrolidin-1-yl)oxy)-3-oxopropyl)disulfaneyl)propanoate	
SITE	17 note = Amidated residue	
DISULFID	4..14	

---

-continued

---

```

SEQUENCE: 406          note = Intrachain disulfide bridge
RGNCAHKQGQ LVWCTYH

SEQ ID NO: 407          moltype = AA length = 17
FEATURE          Location/Qualifiers
source           1..17
                  mol_type = protein
                  organism = synthetic construct
SITE             1
note = Acetylated residue
SITE             8
note =
N-(4-Oxo-4-((3-Oxo-3-(phenylthio)propyl)thio)butanoyl)-L-Lysine
SITE             17
note = Amidated residue
DISULFID        4..14
note = Intrachain disulfide bridge

SEQUENCE: 407          note = Intrachain disulfide bridge
RGNCAHKQGQ LVWCTYH

SEQ ID NO: 408          moltype = AA length = 34
FEATURE          Location/Qualifiers
source           1..34
                  mol_type = protein
                  organism = synthetic construct
SITE             1
note = Acetylated residue
SITE             31
note =
N-(3-(((3-Oxo-3-(phenylthio)propyl)thio)carbonyl)benzoyl)-L-Lysine
SITE             34
note = Amidated residue
DISULFID        5..34
note = Intrachain disulfide bridge

SEQUENCE: 408          note = Intrachain disulfide bridge
FNMQCQRFFY EALHDPNLNE EQRNARIRSI KDDC

```

---

17

17

34

What is claimed is:

1. An immunocytokine composition, comprising:  
an IL-18 polypeptide, wherein the IL-18 polypeptide  
comprises the amino acid sequence set forth in SEQ ID NO: 30; and  
an antibody or an antigen binding fragment thereof specific for an immune cell associated antigen; and  
a linker, wherein the linker comprises:  
a first point of attachment to the IL-18 polypeptide; and  
a second point of attachment to the antibody or antigen binding fragment thereof.

2. The immunocytokine composition of claim 1, wherein the linker comprises a polymer.

3. The immunocytokine composition of claim 1, wherein the first point of attachment is at a residue which is not the N-terminus or the C-terminus of the IL-18 polypeptide.

4. The immunocytokine composition of claim 1, wherein the first point of attachment is at residue 68 of the IL-18 polypeptide.

5. The immunocytokine composition of claim 1, wherein the second point of attachment is at an amino acid residue in an Fc region of the antibody or antigen binding fragment.

6. The immunocytokine composition of claim 5, wherein the linker is covalently attached at Lys 248 of the Fc region of the antibody, wherein amino acid residue position number is based on EU numbering.

7. The immunocytokine composition of claim 1, wherein the IL-18 polypeptide displays reduced binding to IL-18 binding protein (IL-18BP) compared to the wild type IL-18 polypeptide of SEQ ID NO: 1.

8. The immunocytokine composition of claim 1, wherein the antibody or antigen binding fragment thereof is a monoclonal antibody.

9. The immunocytokine composition of claim 1, wherein the antibody or antigen binding fragment thereof comprises an IgG1 or an IgG4.

10. The immunocytokine composition of claim 1, wherein the immune cell associated antigen is programmed cell death protein 1 (PD-1).

11. The immunocytokine composition of claim 10, wherein the antibody or antigen binding fragment thereof comprises nivolumab, pembrolizumab, LZM-009, or cemiplimab.

12. The immunocytokine composition of claim 1, wherein the immune cell associated antigen is programmed death-ligand 1 (PD-L1).

13. The immunocytokine composition of claim 12, wherein the antibody is durvalumab, atezolizumab, or avolimumab.

14. An immunocytokine composition, comprising:  
an IL-18 polypeptide having the amino acid sequence set forth in SEQ ID NO: 30;  
an antibody or an antigen binding fragment thereof comprising an Fc region and having a heavy chain variable region (VH) comprising the amino acid sequence set forth in SEQ ID NO: 376 and a light chain variable region (VL) comprising the amino acid sequence set forth in SEQ ID NO: 377; and  
a linker, wherein the linker comprises:  
a first point of attachment to residue C68 of the IL-18 polypeptide; and

50

55

60

65

**219****220**

a second point of attachment to the antibody or antigen binding fragment thereof at Lys 248 of the Fc region, wherein amino acid residue position number is based on EU numbering.

**15.** The immunocytokine composition of claim **14**,<sup>5</sup> wherein the antibody or antigen binding fragment thereof is LZM-009.

**16.** The immunocytokine composition of claim **14**, wherein the linker comprises polyethylene glycol.

**17.** An immunocytokine composition, comprising:<sup>10</sup>  
an IL-18 polypeptide having the amino acid sequence set forth in SEQ ID NO: 30;

an antibody or an antigen binding fragment thereof specific for programmed cell death protein 1 (PD-1) or programmed death-ligand 1 (PD-L1); and<sup>15</sup>

a linker, wherein the linker comprises:

a first point of attachment to residue C68 the IL-18 polypeptide; and

a second point of attachment to the antibody or antigen binding fragment thereof.<sup>20</sup>

**18.** The immunocytokine composition of claim **17**, wherein the antibody or antigen binding fragment thereof is specific for PD-1.

**19.** The immunocytokine composition of claim **18**,<sup>25</sup> wherein the antibody or antigen binding fragment thereof comprises an Fc region, and wherein the second point of attachment to the antibody or antigen binding fragment thereof is Lys 248 of the Fc region, wherein amino acid residue position number is based on EU numbering.

**20.** The immunocytokine composition of claim **19**,<sup>30</sup> wherein the antibody or antigen binding fragment thereof comprises an IgG1 or an IgG4.

\* \* \* \*