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(54) **METHOD OF TREATING UVEITIS WITH
MULTIVALENT PROTEIN-HYALURONIC
ACID POLYMER CONJUGATE**

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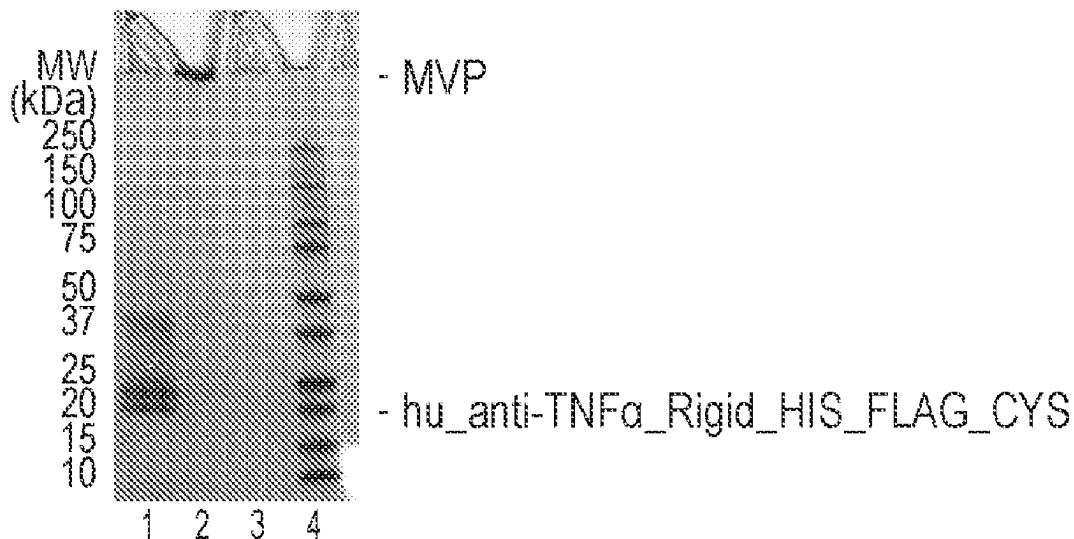
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(57)

ABSTRACT

The present invention relates to multivalent protein-polymer
conjugates, compositions, and methods for treating uveitis,
such as chronic non-infectious uveitis.

Specification includes a Sequence Listing.



1. 5.0 µg hu_anti-TNFα_Rigid_HIS_FLAG_CYS
2. 5.0 µg hu_anti-TNFα_Rigid_HIS_FLAG_CYS MVP0.25 µg hu_anti-TNFα
3. Nothing
4. Ladder

FIG. 1A

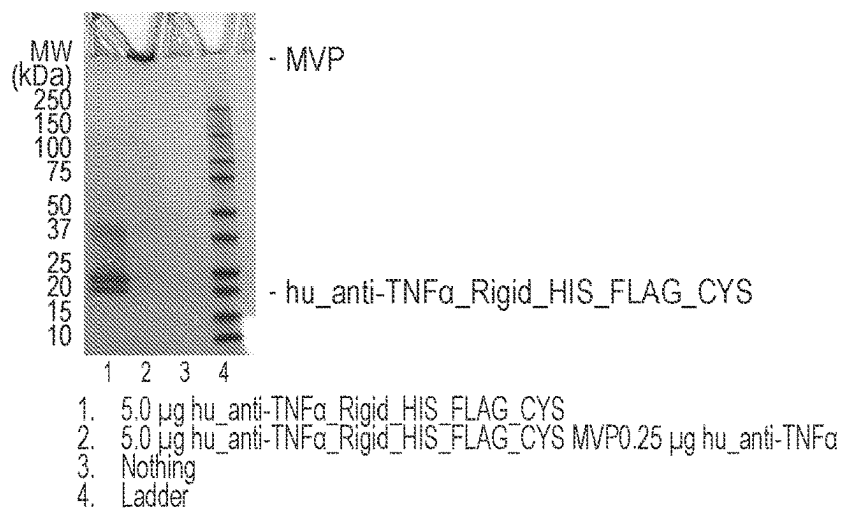


FIG. 1B

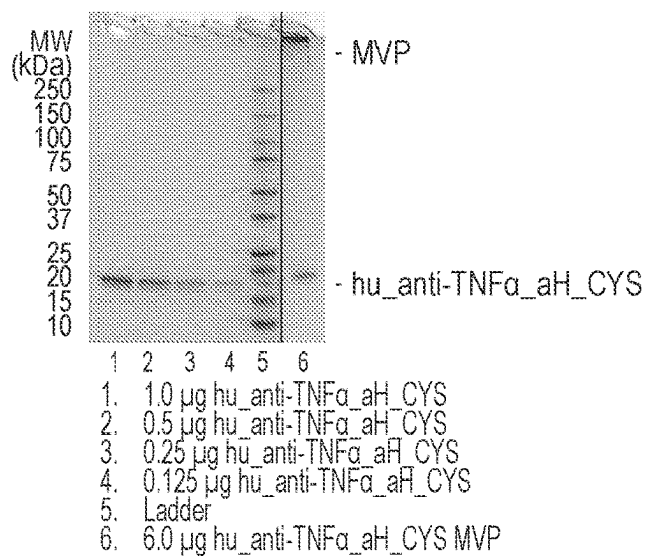


FIG. 1C

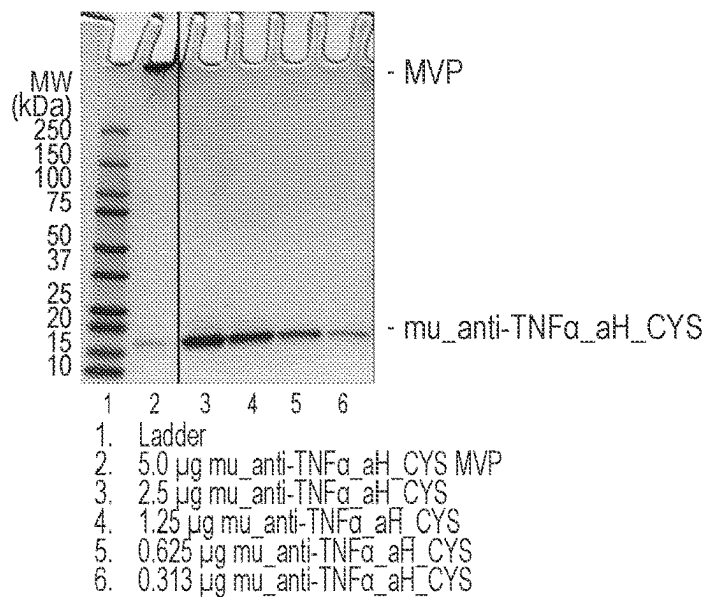


FIG. 1D

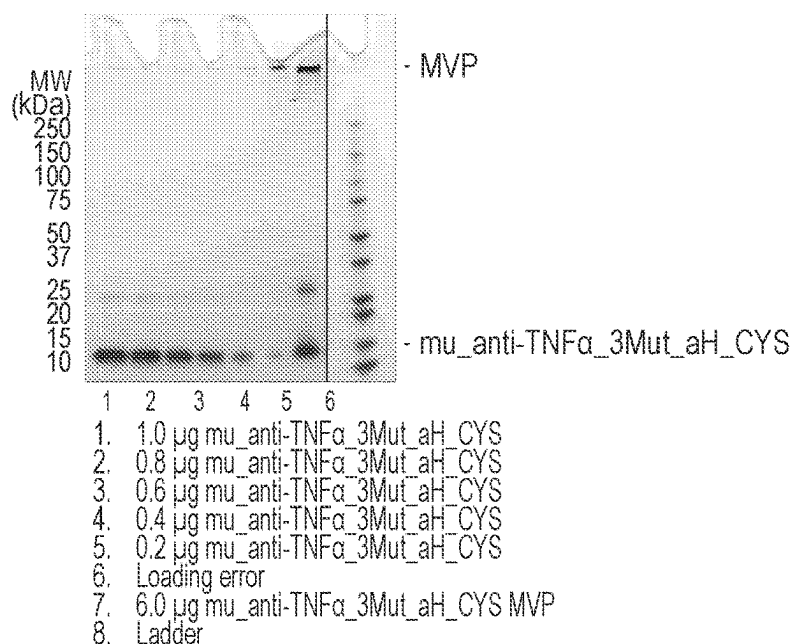


FIG. 1E

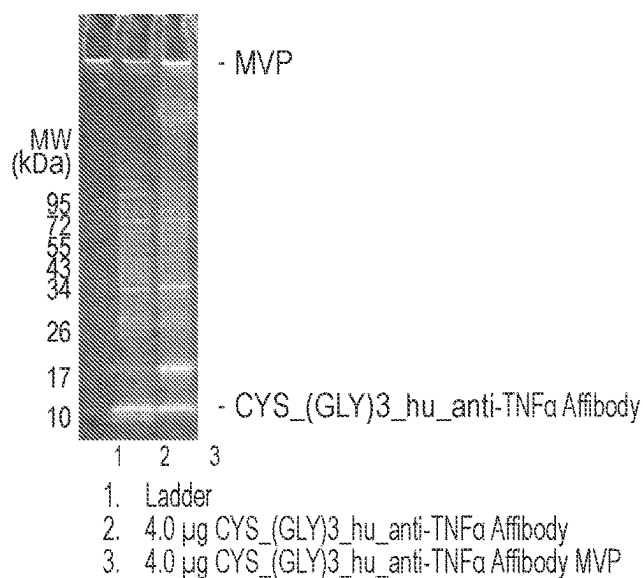


FIG. 1F

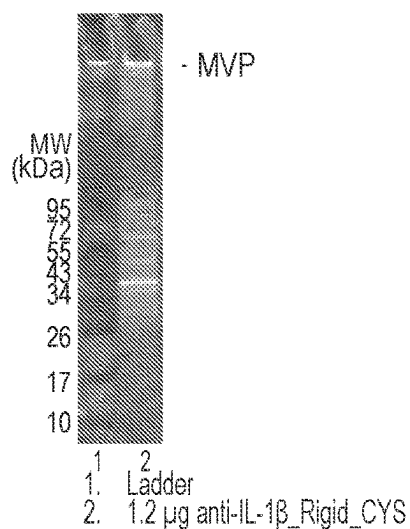


FIG. 2A

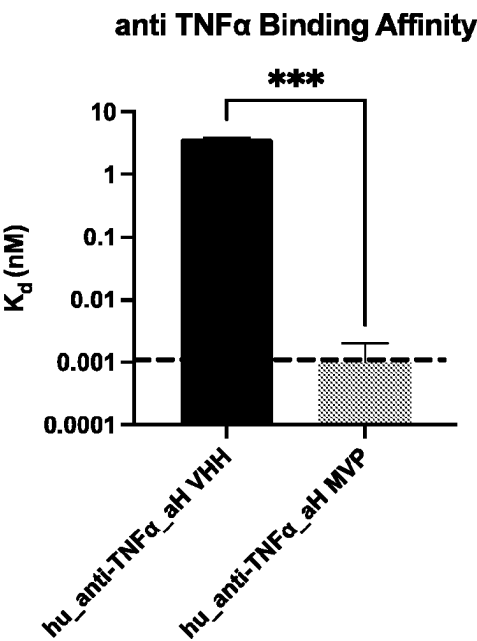


FIG. 2B

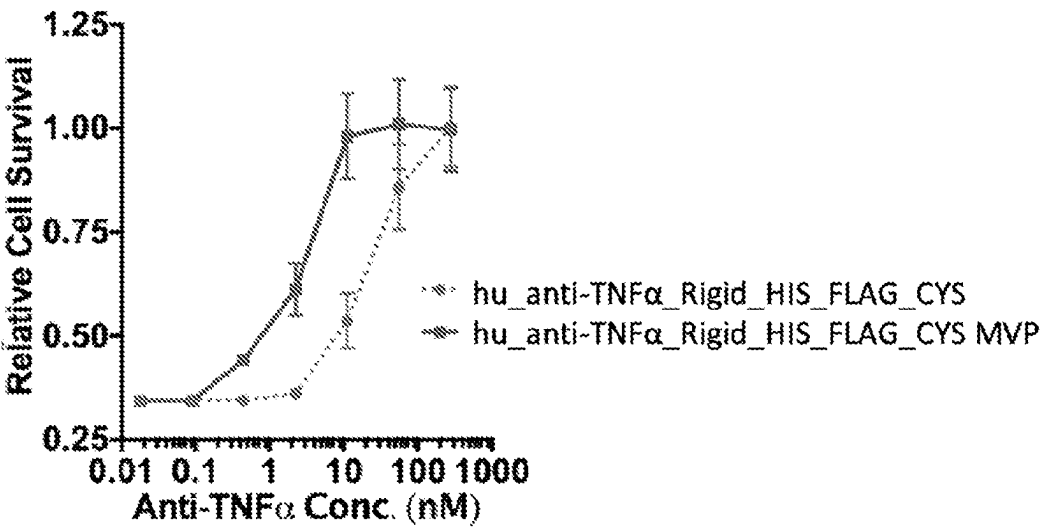


FIG. 3

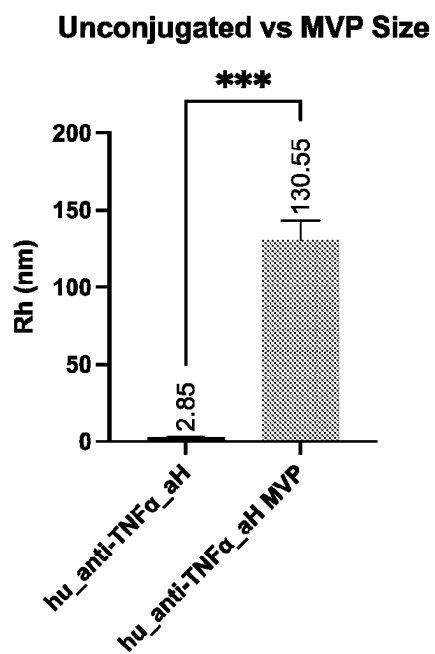


FIG. 4A

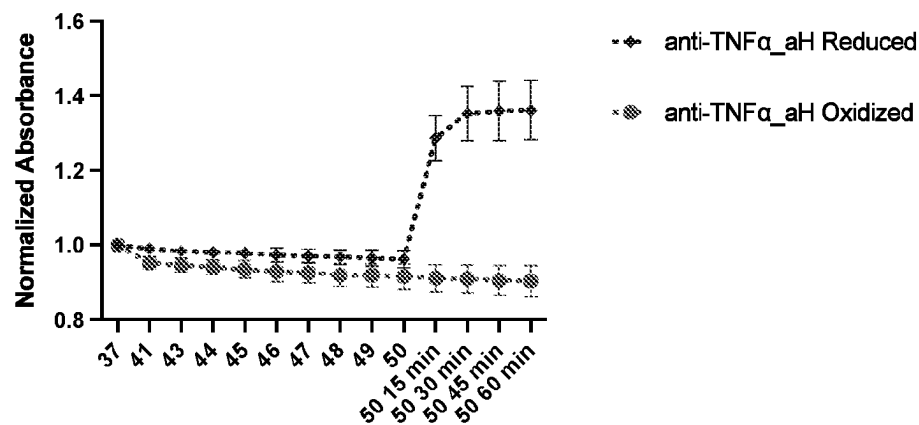


FIG. 4B

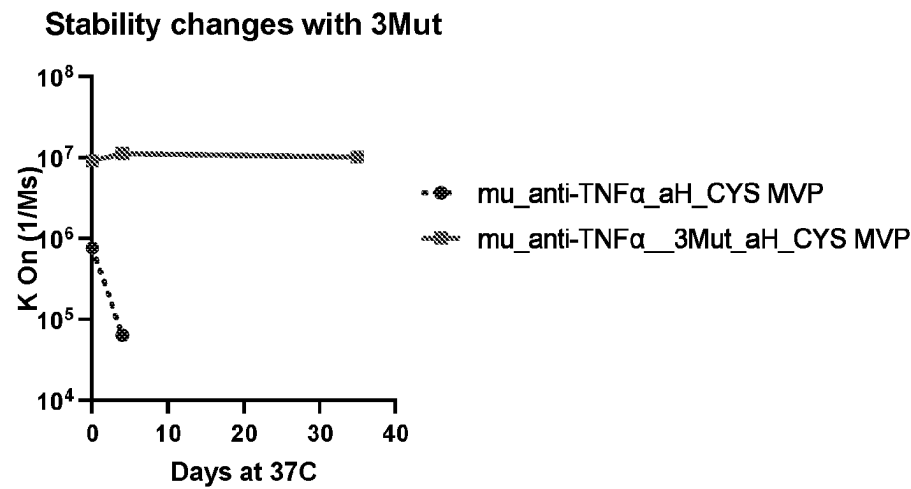


FIG. 4C

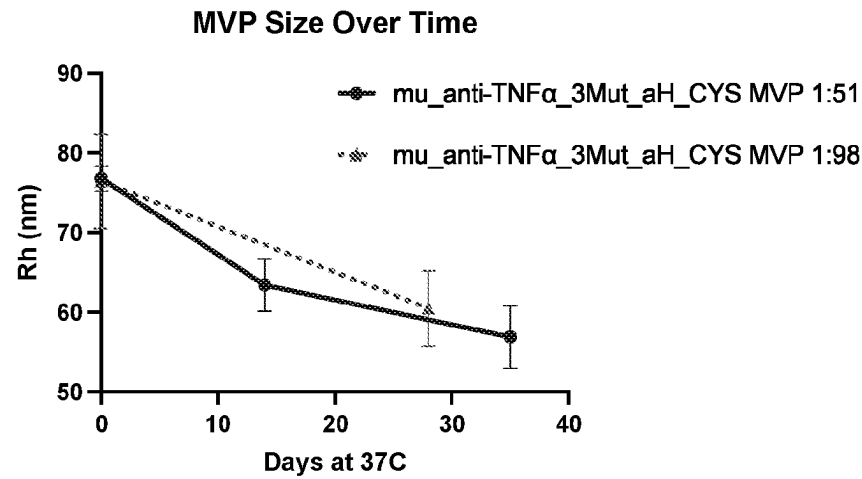


FIG. 5

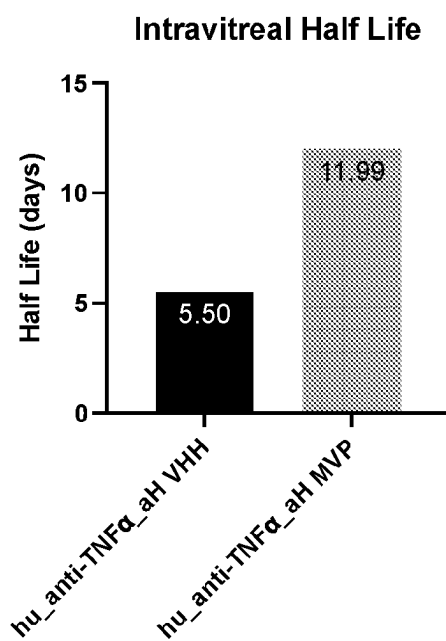


FIG. 6A

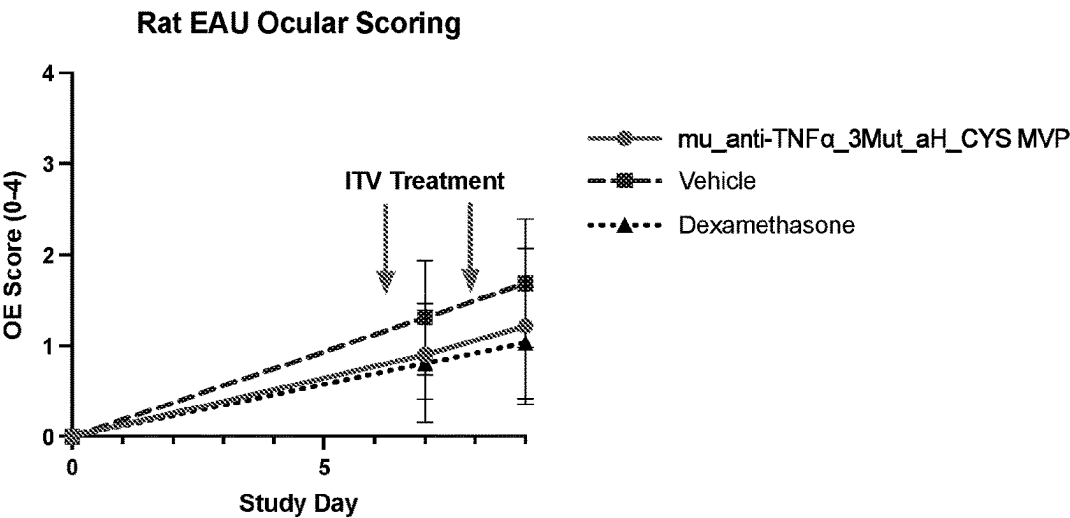


FIG. 6B

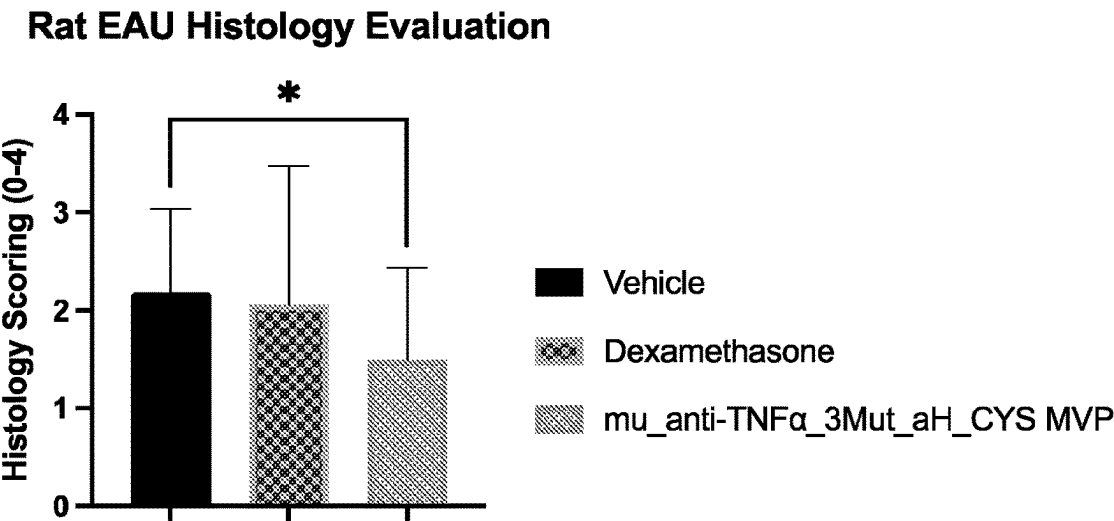


FIG. 7A

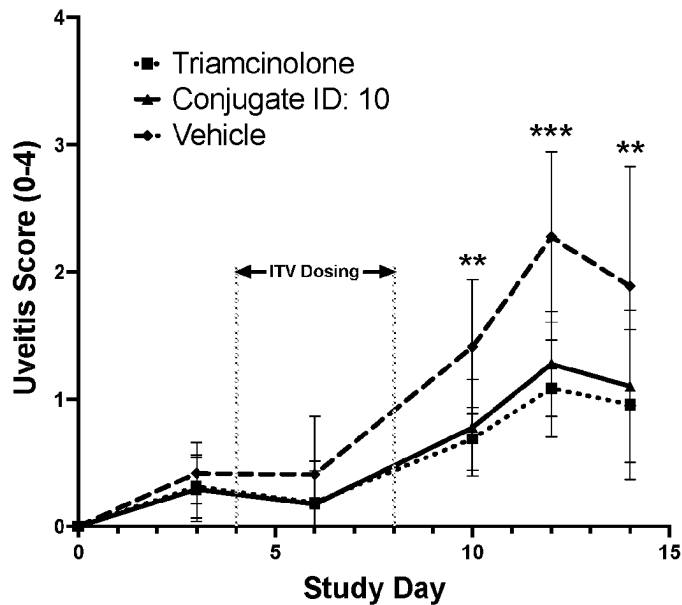


FIG. 7B

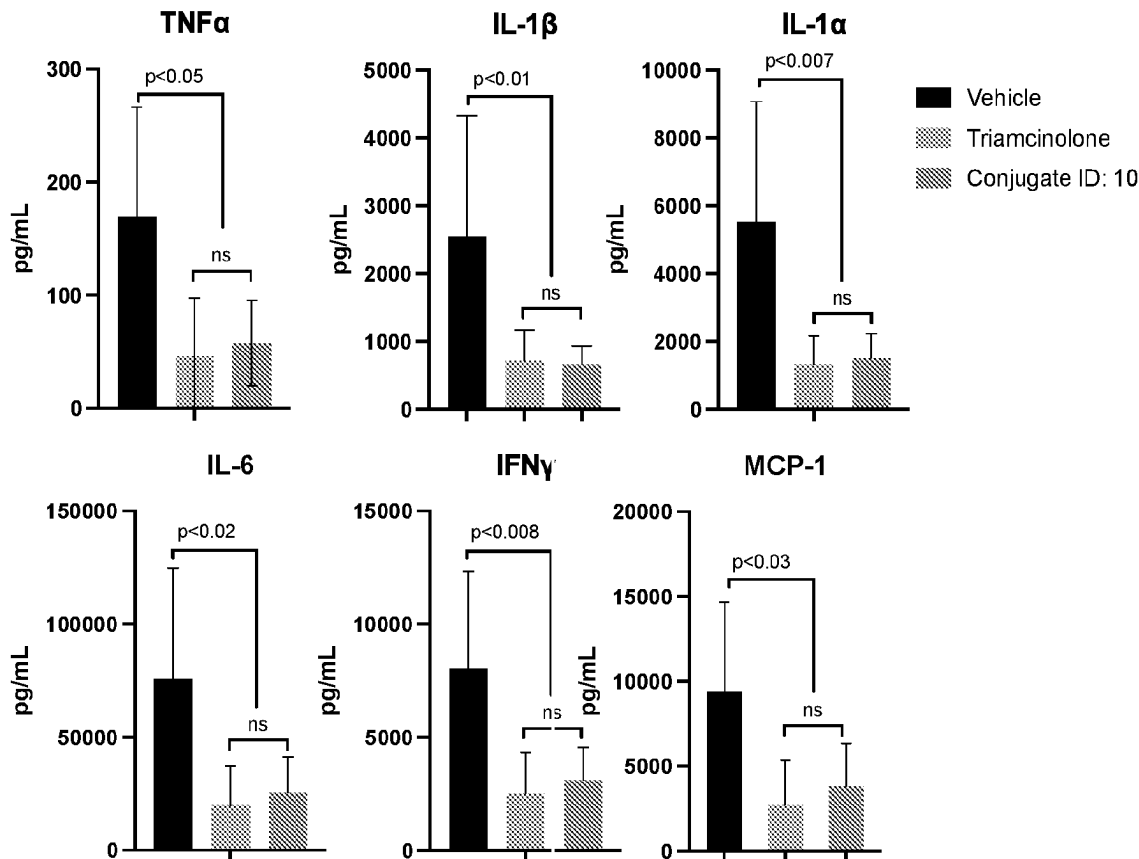


FIG. 8A

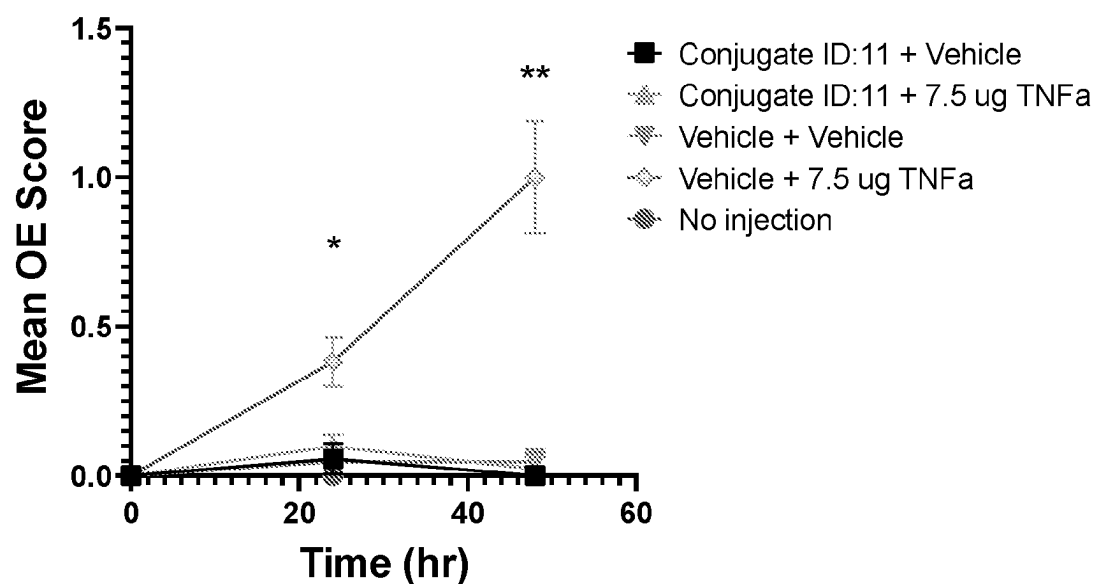
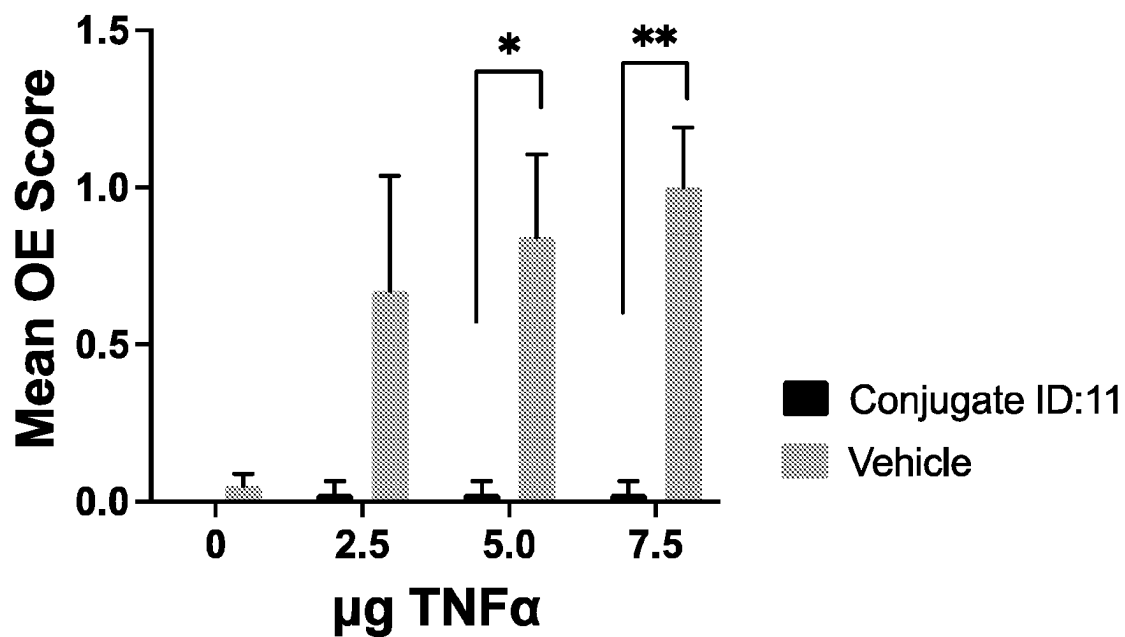


FIG. 8B



METHOD OF TREATING UVEITIS WITH MULTIVALENT PROTEIN-HYALURONIC ACID POLYMER CONJUGATE

CROSS-REFERENCES TO RELATED APPLICATIONS

[0001] This application claims priority to U.S. Provisional Application No. 63/331,554, filed Apr. 15, 2022, which is incorporated herein in its entirety for all purposes.

SEQUENCE LISTING

[0002] The material in the accompanying sequence listing is hereby incorporated by reference in its entirety. The accompanying file, named 2023-04-11 Sequence Listing_ST26 052566-508001WO.xml was created on Apr. 11, 2023, and is 117,702 bytes in size.

BACKGROUND OF THE INVENTION

[0003] The use of biopolymers to modify the properties of biologically active agents is a recurring theme across a wide range of medical and biological applications. A variety of chemical linkers can be used to attach bioactive peptides or proteins to biopolymers to modify the pharmacological properties of the resulting conjugate for use as a drug that can provide optimal treatment of specific diseases. Peptide-polymer conjugates comprising multiple copies of one or more species of peptide conjugated to a single biopolymer chain have been employed to impart specific improvements to the pharmacological properties of the peptides, including: (1) higher binding affinity to the biological target, (2) slower diffusivity through a target tissue, and (3) inhibition of proteases that could deactivate the biological activity of the peptides or proteins.

[0004] These improved pharmacological properties of peptide-polymer conjugates are particularly useful for the delivery of potent drugs that are delivered directly into the diseased tissue. The dose delivered directly into the tissue can be lower than would be required to achieve the same therapeutic effect after systemic administration because the drug has been administered locally to the target tissue. It is also possible to administer to drugs to tissues that otherwise have poor transport properties from the blood. Specific examples of tissues where direct drug administration is common include the posterior eye chamber via intravitreal injection and articular joints via intra-articular injection.

[0005] However, local tissue administration requires a professional to safely provide the required injection, which makes them more burdensome and costly to administer compared to systemic administration. When the peptide

drug is administered as part of a peptide-polymer conjugate, it is possible to substantially reduce the frequency of drug administration, thereby reducing the burden on the patient to receive effective treatment. Furthermore, a reduction in the number of local injections reduces the risk of local tissue injury or adverse effects to the injection. Finally, the need for less frequent administrations can reduce the amount of time that the drug concentration in the target tissue is below the therapeutic concentration, thereby improving the overall efficacy of the drug. Based on these advantages, there is a strong motivation to develop protein-polymer drug products for a variety of diseases.

[0006] Uveitis is a group of sight-threatening intraocular inflammation diseases that is responsible for roughly 5-10% of blindness cases worldwide. Chronic non-infectious uveitis can result in nerve damage and vision loss. Most patients are treated using corticosteroids, which can lead to serious side effects. Intravitreal administration of biologic TNF α inhibitors can substantially reduce the need for steroids. However, these products were not designed or validated for intravitreal use, and off-label intravitreal treatment with existing TNF α inhibitors is not recommended.

[0007] Therefore, there is a need to develop purified peptide-polymer conjugates and methods for treating uveitis, such as chronic non-infectious uveitis. The present invention meets this and other needs.

BRIEF SUMMARY OF THE INVENTION

[0008] In some embodiments, the method of the present invention is a method for treating uveitis in a subject in need thereof, comprising administering to the subject a therapeutically effective amount of a conjugate of Formula V:



[0009] wherein

[0010] each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0011] each Y is an organic linker;

[0012] Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and

[0013] subscript n is an integer of from 1 to 1000.

[0014] In some embodiments, the conjugate of the present invention is a random polymer of Formula VI:



[0015] having a molecular weight of from about 0.1 MDa to about 3 MDa;

[0016] wherein

[0017] each X is independently an anti-TNF- α or anti-IL-1 β peptide comprising:

(SEQ ID NO: 101)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG

KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNPLEPETA

VYYCAARDGI PTERSVESYN YWQGTQVTV SSPSTPPTPS PSTPPGGCDD

DDK,

(SEQ ID NO: 102)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG

KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNPLEPETA

-continued

VYYCAARDGI PSTRSVESYN YWGQGTQVTV SSAEAAAKEA AAKEAAKAG

C,

(SEQ ID NO: 103)

QVQLQDS GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKERE

VGAVSWSGGT TVYADSVLGR FEISRDSARK SVYLMNSLK PEDTAVYYCA

ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAAKEA AAKAGC,

(SEQ ID NO: 104)

QVQLQES GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKERE

VGAVSWSGGT TVYADSVKGR FTISRDSARK SVYLMNSLK PEDTAVYYCA

ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAAKEA AAKAGC,

(SEQ ID NO: 105)

CGGVDNKFN KEVGWAFGEI GALPNLNALQ FRAFIISLWD DPSQSANLLA

EAKKLNDQA PK,

or

(SEQ ID NO: 106)

EIVMTQS PSTLSASVGD RVIITCQASQ SIDNWLSWYQ QKPGKAPKLL

IYRASTLASG VPSRFGSGGS GAFTLTIS LQPDDEFATYY CQNTGGGVSI

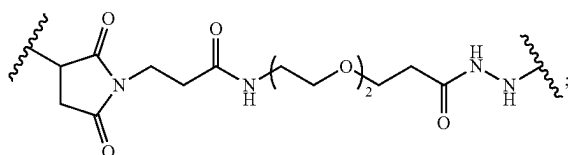
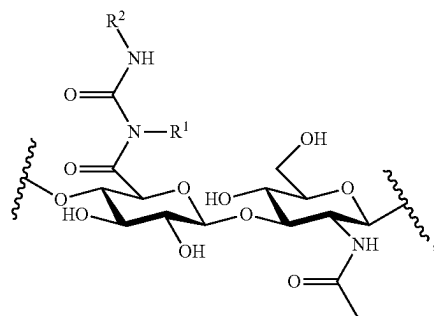
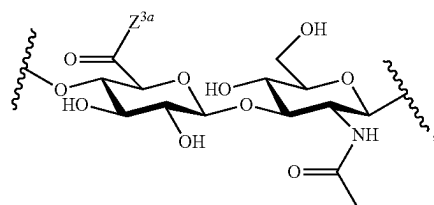
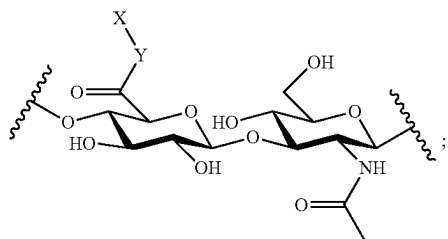
APGQGTKLTV LGGGGGSGGG GSGGGGSGGG GSEVQLVESG GGLVQPGGSL

RLSCTASGFS LSSAAMAWVR QAPKGLEWV GIIYDSASTY YASWAKGRFT

ISRDTSKNTV YLMNSLRAE DTAVYYCARE RAIFSGDFVL WGQGTSLTVS

SSPSTPPTPS PSTPPGGC;

[0018] each Y is an organic linker having the structure:

[0020] each Z^2 has the structure:[0021] each Z^3 independently has the structure:[0019] each X—Y— Z^1 moiety has the structure:[0022] each R^1 and R^2 is independently C_1 - C_6 alkyl, $-(C_1$ - C_6 alkyl)- NR^3R^4 , or C_5 - C_8 cycloalkyl;[0023] each R^3 and R^4 is independently H or C_1 - C_6 alkyl;

- [0024] each $Z^{3\alpha}$ is independently OH or Y';
 [0025] each Y' is an unreacted organic linker;
 [0026] subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;
 [0027] subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and
 [0028] subscript q is an integer of from 100 to 10000.

BRIEF DESCRIPTION OF THE DRAWINGS

[0029] FIG. 1 shows SDS-PAGE images of representative anti-inflammatory peptide polymer conjugates for each sequence ID. (A) SDS-PAGE image of (SEQ ID NO:101)+HyA (850 kDa) conjugate #2 with a valency of 55 compared to the unconjugated VHH. (B) SDS-PAGE image of (SEQ ID NO:102)+HyA (850 kDa) conjugate #3 with a valency of 65 compared to the unconjugated VHH. (C) SDS-PAGE image of (SEQ ID NO:103)+HyA (850 kDa) conjugate #5 with a valency of 121 compared to the unconjugated VHH. (D) SDS-PAGE image of (SEQ ID NO:104)+HyA (850 kDa) conjugate #6 with a valency of 51 compared to the unconjugated VHH. (E) SDS-PAGE image of (SEQ ID NO:105)+HyA (850 kDa) conjugate #8 with a valency of 21 compared to the unconjugated affibody. (F) SDS-PAGE image of (SEQ ID NO:106)+HyA (850 kDa) conjugate #9 with a valency of 15.

[0030] FIG. 2 shows (A) the TNF α binding affinity of a (SEQ ID NO:102)+HyA (850 kDa) conjugate #4 with a valency of 120 is greater than that of an unconjugated TNF α , as determined by biolayer interferometry (** $p < 0.001$ Student's t-tests with $n=3$) Dashed line indicates the limit of detection for the instrument and the binding affinity of the conjugate is below that limit. (B) The bioactivity of a (SEQ ID NO: 101)+HyA (850 kDa) conjugate #1 with a valency of 9 to inhibit TNF α -induced apoptosis in L929 fibroblasts is ~ 10 fold greater than the unconjugated VHH.

[0031] FIG. 3 shows the hydrodynamic radius of conjugate #3 consisting of (SEQ ID NO:102)+HyA (850 kDa) with a valency of 65 is greater than that of an unconjugated VHH. (** $p < 0.001$ Student's t-tests with $n=3$).

[0032] FIG. 4 shows (A) the normalized absorbance at 280 nm ("A280") in unconjugated protein SEQ ID NO:102 as the temperature increased. The oxidized version of the VHH showed minimal change in absorbance when the temperature increased from 37° C.-50° C. whereas the reduced construct showed increased absorbance starting at 50° C., indicating that it has unfolded and was less thermally stable. Error bars represent SD; (B) change in association constant of the TNF α binding affinity to mu_anti-TNF α _aH_CYS conjugates with or without the 3Mut stability enhancement mutation after incubation at 37° C. for the indicated number of days as determined by biolayer interferometry. The samples used were either (SEQ ID NO:103)+HyA (850 kDa) conjugate #5 with a valency of 121 ("mu_anti-TNF α _aH_CYS MVP") or (SEQ ID NO:104)+HyA (850 kDa) conjugate #6 with a valency of 51 ("mu_anti-TNF α _3Mut_aH_CYS MVP"). After 4 days at 37° C., the association constant of conjugate #6 had minimal change after 35 days at 37° C. In contrast, the association constant of the non-mutated conjugate #5 dramatically decreased after 5 days at 37° C., indicative of decreased stability; (C) representative DLS data conjugate size after incubation at 37° C. in vitreous mimetic buffer for up to 35 days. Conjugates were made

with (SEQ ID NO:104)+HyA (850 kDa) with a valency of 51 (conjugate #6, "mu_anti-TNF α _3Mut_aH_CYS MVP 1:51") or 98 (conjugate #7, "mu_anti-TNF α _3Mut_aH_CYS MVP 1:98"). There was no significant difference in MVP size based on valency range from ~ 50 to ~ 100 antibodies per polymer. The conjugates slowly decreased in size to about 75% of the original radius after 35 days at 37° C. Error bars represent SD.

[0033] FIG. 5 shows that conjugation can increase the intravitreal half-life of an anti-inflammatory therapeutic in rabbit intravitreal pharmacokinetics model. Each rabbit received an equal molar 50 μ L intravitreal injection of either unconjugated SEQ ID NO:102 or (SEQ ID NO:102)+HyA (850 kDa) conjugate #4 with a valency of 120. The intravitreal half-life was determined using a non-linear fit of the VHH concentration at each timepoint. Multivalent conjugation increased the half-life at least 2 \times compared to unconjugated VHH.

[0034] FIG. 6 shows that an anti-TNF α conjugate sufficiently suppressed ocular inflammation in a rat experimental autoimmune uveoretinitis model. Conjugate #7 was made with (SEQ ID NO:104)+HyA (850 kDa) and a valency of 98. (A) Average inflammation score (0=none to 4=severe) observed in rat eyes after EAU induction in rats injected intravitreally with either vehicle, dexamethasone (5 μ g) or Conjugate #7 ("mu_anti-TNF α _3Mut_aH_CYS MVP") (12.5 μ g) ($n=8$). Both dexamethasone and conjugate treated eyes showed decreased inflammation compared to the vehicle control one day after ITV treatment. (B) Average histology inflammation scores (0=none to 4=severe) in the same cohort of rats that were sacrificed 14 days after model induction. Left bar: vehicle, middle bar: dexamethasone treated, right bar: Conjugate #7 treated. The conjugate treated eyes (right bar) were less inflamed than vehicle treated eyes (left bar) (* $p < 0.05$ Student's t-tests).

[0035] FIG. 7A-7B show that an anti-TNF α conjugate sufficiently suppressed ocular inflammation in a rat experimental autoimmune uveoretinitis model. Conjugate #10 was made with (SEQ ID NO:104)+HyA (850 kDa) and a valency of 96.5. (A) Average inflammation score (0=none to 4=severe) observed in rat eyes after EAU induction in rats injected intravitreally with either vehicle ($n=24$), triamcinolone (40 μ g) ($n=22$) or conjugate #10 ("Conjugate ID: 10") (19 g) ($n=20$). Both triamcinolone and conjugate treated eyes showed decreased inflammation compared to the vehicle control one day after ITV treatment. No statistical difference was measured between triamcinolone and conjugate #10 at any timepoint. Conjugate #10 and vehicle were significantly different at Day 10, $p=0.002$, at day 12, $p < 0.001$ and at day 14, $p=0.007$. (B) Cytokine analysis of vitreous samples prepared from rat eyes injected intravitreally with either vehicle ($n=24$), triamcinolone (40 μ g) ($n=22$) or conjugate #10 ("Conjugate ID: 10") (19 g) ($n=20$). Bar graphs indicate the concentration of key pro-inflammatory cytokines and inflammatory regulators.

[0036] FIGS. 8A-8B show that an anti-TNF α conjugate sufficiently suppressed ocular inflammation in a rabbit TNF α -induced ocular inflammation model. Conjugate #11 was made with (SEQ ID NO:102)+HyA (850 kDa) and a valency of 132. (A) Average inflammation score over time (0=none to 4=severe) observed in rabbit eyes after inflammation induction using 7.5 μ g of TNF α in rabbits injected intravitreally with either vehicle, or conjugate #11 ("Conjugate ID 11") (0.26 mg) or no injection ($n=26$). In the

rabbits treated with 7.5 μg of $\text{TNF}\alpha$, there was a statistical significance between conjugate #11 and vehicle treated eyes at 24 and 48 hours. (B) After 48 hours, at all concentrations of $\text{TNF}\alpha$ dosed, conjugate #11 treated eyes showed decreased inflammation compared to the vehicle control 48 hours after inflammation induction. A statistically significant difference was observed at 5.0 μg ($p=0.03$) and 7.5 μg ($p<0.001$) of $\text{TNF}\alpha$ dosed.

DETAILED DESCRIPTION OF THE INVENTION

I. Definitions

[0037] Unless specifically indicated otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by those of ordinary skill in the art to which this invention belongs. In addition, any method or material similar or equivalent to a method or material described herein can be used in the practice of the present invention. For purposes of the present invention, the following terms are defined.

[0038] “About” when referring to a value includes the stated value \pm 10% of the stated value. For example, about 50% includes a range of from 45% to 55%, while about 20 molar equivalents includes a range of from 18 to 22 molar equivalents. Accordingly, when referring to a range, “about” refers to each of the stated values \pm 10% of the stated value of each end of the range. For instance, a ratio of from about 1 to about 3 (weight/weight) includes a range of from 0.9 to 3.3.

[0039] “Alkyl” is a linear or branched saturated monovalent or divalent hydrocarbon. For example, an alkyl group can have 1 to 10 carbon atoms (i.e., C_{1-10} alkyl) or 1 to 8 carbon atoms (i.e., C_{1-8} alkyl) or 1 to 6 carbon atoms (i.e., C_{1-6} alkyl) or 1 to 4 carbon atoms (i.e., (C_{1-4}) alkyl). Examples of alkyl groups include, but are not limited to, methyl (Me, $-\text{CH}_3$), ethyl (Et, $-\text{CH}_2\text{CH}_3$), 1-propyl (n-Pr, n-propyl, $-\text{CH}_2\text{CH}_2\text{CH}_3$), 2-propyl (i-Pr, i-propyl, $-\text{CH}(\text{CH}_3)_2$), 1-butyl (n-Bu, n-butyl, $-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 2-methyl-1-propyl (i-Bu, i-butyl, $-\text{CH}_2\text{CH}(\text{CH}_3)_2$), 2-butyl (s-Bu, s-butyl, $-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_3$), 2-methyl-2-propyl (t-Bu, t-butyl, $-\text{C}(\text{CH}_3)_3$), 1-pentyl (n-pentyl, $-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 2-pentyl ($-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_2\text{CH}_3$), 3-pentyl ($-\text{CH}(\text{CH}_2\text{CH}_3)_2$), 2-methyl-2-butyl ($-\text{C}(\text{CH}_3)_2\text{CH}_2\text{CH}_3$), 3-methyl-2-butyl ($-\text{CH}(\text{CH}_3)\text{CH}(\text{CH}_3)_2$), 3-methyl-1-butyl ($-\text{CH}_2\text{CH}_2\text{CH}(\text{CH}_3)_2$), 2-methyl-1-butyl ($-\text{CH}_2\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_3$), 1-hexyl ($-\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 2-hexyl ($-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$), 3-hexyl ($-\text{CH}(\text{CH}_2\text{CH}_3)\text{CH}_2\text{CH}_2\text{CH}_3$), 2-methyl-2-pentyl ($-\text{C}(\text{CH}_3)_2\text{CH}_2\text{CH}_2\text{CH}_3$), 3-methyl-2-pentyl ($-\text{CH}(\text{CH}_3)\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}_3$), 4-methyl-2-pentyl ($-\text{CH}(\text{CH}_3)\text{CH}_2\text{CH}(\text{CH}_3)_2$), 3-methyl-3-pentyl ($-\text{C}(\text{CH}_3)(\text{CH}_2\text{CH}_3)_2$), 2-methyl-3-pentyl ($-\text{CH}(\text{CH}_2\text{CH}_3)\text{CH}(\text{CH}_3)_2$), 2,3-dimethyl-2-butyl ($-\text{C}(\text{CH}_3)_2\text{CH}(\text{CH}_3)_2$), 3,3-dimethyl-2-butyl ($-\text{CH}(\text{CH}_3)\text{C}(\text{CH}_3)_3$), and octyl ($-(\text{CH}_2)_7\text{CH}_3$).

[0040] “Cycloalkyl” refers to a single saturated or partially unsaturated all carbon ring having 3 to 20 annular carbon atoms (i.e., C_{3-20} cycloalkyl), for example from 3 to 12 annular atoms, for example from 3 to 10 annular atoms, or 3 to 8 annular atoms, or 3 to 6 annular atoms, or 3 to 5 annular atoms, or 3 to 4 annular atoms. The term “cycloalkyl” also includes multiple condensed, saturated and partially unsaturated all carbon ring systems (e.g., ring systems

comprising 2, 3 or 4 carbocyclic rings). Accordingly, cycloalkyl includes multicyclic carbocycles such as a bicyclic carbocycles (e.g., bicyclic carbocycles having about 6 to 12 annular carbon atoms such as bicyclo[3.1.0]hexane and bicyclo[2.1.1]hexane), and polycyclic carbocycles (e.g. tricyclic and tetracyclic carbocycles with up to about 20 annular carbon atoms). The rings of a multiple condensed ring system can be connected to each other via fused, spiro and bridged bonds when allowed by valency requirements. Non-limiting examples of monocyclic cycloalkyl include cyclopropyl, cyclobutyl, cyclopentyl, 1-cyclopent-1-enyl, 1-cyclopent-2-enyl, 1-cyclopent-3-enyl, cyclohexyl, 1-cyclohex-1-enyl, 1-cyclohex-2-enyl and 1-cyclohex-3-enyl.

[0041] “Organic linker” as used herein refers to a chemical moiety that directly or indirectly covalently links the peptide to the polymer. Organic linkers useful in the present invention can be about 100 Da to 500 Da. The types of organic linkers of the present invention include, but are not limited to, imides, amides, amines, esters, carbamates, ureas, thioethers, thiocarbamates, thiocarbonate and thioureas. One of skill in the art will appreciate that other types of organic linkers are useful in the present invention.

[0042] “Thiol” refers to the $-\text{SH}$ functional group.

[0043] “Thiol reactive group” refers to a group capable of reacting with a thiol to form a covalent bond to the sulfur atom. Representative thiol reactive groups include, but are not limited to, thiol, TNB-thiol, haloacetyl, aziridine, acryloyl, vinylsulfone, APN (3-arylpropionitrile), maleimide and pyridyl disulfide. Reaction of the thiol reactive group with a thiol can form a disulfide or a thioether.

[0044] “Peptide,” “polypeptide,” and “protein” are used interchangeably herein, and refer to naturally occurring and synthetic amino acids of any length, as well as amino acid analogs and amino acid mimetics that function in a manner similar to the naturally occurring amino acids. The term “peptide” includes fusion proteins, including, but not limited to, fusion proteins with a heterologous amino acid sequence, fusions with heterologous and homologous leader sequences, with or without N-terminal methionine residues; immunologically tagged proteins; and the like. Peptides further include post-translationally modified peptides.

[0045] “VHH” as used herein refers to a single-domain heavy chain antibody.

[0046] “DARPin” refers to a designed ankyrin repeat protein, which is a genetically engineered antibody mimetic protein that can exhibit highly specific and high-affinity target protein binding.

[0047] An “alpha-helix” or “ α -helix” is a common motif in the secondary structure of proteins and is a right hand-helix conformation in which every backbone N—H group hydrogen bonds to the backbone C=O group of the amino acid located four residues earlier along the protein sequence. The alpha-helix is also known as a classic Pauling-Corey-Branson α -helix, or 3.6₁₃-helix, which denotes the average number of residues per helical turn (3.6) with 13 atoms being involved in the ring formed by the hydrogen bond. Peptides that contain an alpha-helix is said to be alpha-helical. Such peptides may be partly or entirely alpha-helical. As understood in the art, an alpha-helix has at least four amino acid residues. In some embodiments, an alpha-helix has from 4 to 40 amino acids.

[0048] Provided are also pharmaceutically acceptable salts of the compounds or peptides described herein. “Pharmaceutically acceptable” or “physiologically acceptable” refer

to compounds, salts, compositions, dosage forms and other materials which are useful in preparing a pharmaceutical composition that is suitable for veterinary or human pharmaceutical use.

[0049] “Pharmaceutical composition” as used herein refers to a product comprising the specified ingredients in the specified amounts, as well as any product, which results, directly or indirectly, from combination of the specified ingredients in the specified amounts. The pharmaceutical composition is generally safe for biological use.

[0050] “Pharmaceutically acceptable excipient” as used herein refers to a substance that aids the administration of an active agent to an absorption by a subject. Pharmaceutically acceptable excipients useful in the present invention include, but are not limited to, binders, fillers, disintegrants, lubricants, coatings, sweeteners, flavors and colors. One of skill in the art will recognize that other pharmaceutically acceptable excipients are useful in the present invention.

[0051] The conjugates described herein may be prepared and/or formulated as pharmaceutically acceptable salts or when appropriate as a free base. Pharmaceutically acceptable salts are non-toxic salts of a free base form of a compound that possess the desired pharmacological activity of the free base. These salts may be derived from inorganic or organic acids or bases. For example, a conjugate that contains a basic nitrogen may be prepared as a pharmaceutically acceptable salt by contacting the compound with an inorganic or organic acid. Non-limiting examples of pharmaceutically acceptable salts include sulfates, pyrosulfates, bisulfates, sulfites, bisulfites, phosphates, monohydrogenphosphates, dihydrogenphosphates, metaphosphates, pyrophosphates, chlorides, bromides, iodides, acetates, propionates, decanoates, caprylates, acrylates, formates, isobutyrate, caproates, heptanoates, propiolates, oxalates, malonates, succinates, suberates, sebacates, fumarates, maleates, butyne-1,4-dioates, hexyne-1,6-dioates, benzoates, chlorobenzoates, methylbenzoates, dinitrobenzoates, hydroxybenzoates, methoxybenzoates, phthalates, sulfonates, methylsulfonates, propylsulfonates, besylates, xylenesulfonates, naphthalene-1-sulfonates, naphthalene-2-sulfonates, phenylacetates, phenylpropionates, phenylbutyrates, citrates, lactates, γ -hydroxybutyrates, glycolates, tartrates, and mandelates. Lists of other suitable pharmaceutically acceptable salts are found in Remington: The Science and Practice of Pharmacy, 21st Edition, Lippincott Williams and Wilkins, Philadelphia, Pa., 2006.

[0052] Examples of “pharmaceutically acceptable salts” of the conjugates disclosed herein also include salts derived from an appropriate base, such as an alkali metal (for example, sodium, potassium), an alkaline earth metal (for example, magnesium), ammonium and NR_4^+ (wherein R is $\text{C}_1\text{--C}_4$ alkyl). Also included are base addition salts, such as sodium or potassium salts.

[0053] “Therapeutically effective amount” as used herein refers to a dose that produces therapeutic effects for which it is administered. The exact dose will depend on the purpose of the treatment, and will be ascertainable by one skilled in the art using known techniques (see, e.g., Lieberman, *Pharmaceutical Dosage Forms* (vols. 1-3, 1992); Lloyd, *The Art, Science and Technology of Pharmaceutical Compounding* (1999); Pickar, *Dosage Calculations* (1999); and Remington: *The Science and Practice of Pharmacy*, 20th Edition, 2003, Gennaro, Ed., Lippincott, Williams & Wilkins). In

sensitized cells, the therapeutically effective dose can be lower than the conventional therapeutically effective dose for non-sensitized cells.

[0054] “Treatment” or “treat” or “treating” as used herein refers to an approach for obtaining beneficial or desired results. For purposes of the present disclosure, beneficial or desired results include, but are not limited to, alleviation of a symptom and/or diminishment of the extent of a symptom and/or preventing a worsening of a symptom associated with a disease or condition. In one embodiment, “treatment” or “treating” includes one or more of the following: a) inhibiting the disease or condition (e.g., decreasing one or more symptoms resulting from the disease or condition, and/or diminishing the extent of the disease or condition); b) slowing or arresting the development of one or more symptoms associated with the disease or condition (e.g., stabilizing the disease or condition, delaying the worsening or progression of the disease or condition); and c) relieving the disease or condition, e.g., causing the regression of clinical symptoms, ameliorating the disease state, delaying the progression of the disease, increasing the quality of life, and/or prolonging survival.

[0055] “Prophylaxis” refers to preventing or retarding the progression of clinical illness in patients suffering from a disease.

[0056] A “subject” of the present invention is a mammal, which can be a human or a non-human mammal, for example a companion animal, such as a dog, cat, rat, or the like, or a farm animal, such as a horse, donkey, mule, goat, sheep, pig, or cow, and the like. In some embodiments, the subject is human.

II. Conjugates

[0057] In some embodiments, the conjugate of the present invention is a conjugate of Formula V:



[0058] wherein

[0059] each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0060] each Y is an organic linker;

[0061] Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and

[0062] subscript n is an integer of from 1 to 1000.

[0063] In some embodiments, each X is independently an anti-TNF- α peptide or an anti-interleukin-1 β peptide.

[0064] In some embodiments, each X is a monoclonal IgG, an IgG fragment, single chain scFv, single-domain heavy-chain VHH, adnectin, affibody, anticalin, DARPIn, or an engineered Kunitz-type inhibitor. In some embodiments, each X is a monoclonal IgG. In some embodiments, each X is an IgG fragment. In some embodiments, each X is a single-domain heavy-chain VHH. In some embodiments, each X is a DARPIn.

[0065] In some embodiments, each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 61-73, 81-85, 91-98, 101-109, 111-118, and 145-151. In some embodiments, each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 101-109 and 148-154.

[0066] In some embodiments, each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 61-73, 81-85, 91-95, 101-106, and 111-118.

[0067] In some embodiments, each X is a peptide having an amino acid sequence comprising:

(SEQ ID NO: 101)
 QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
 KEREFPVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
 VYYCAARDGI PTSRSVESYN YWGQGTQVTV SSPSTPPTPS PSTPPGGCDD
 DDK,

(SEQ ID NO: 102)
 QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
 KEREFPVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
 VYYCAARDGI PTSRSVESYN YWGQGTQVTV SSAEAAAKEA AAKEAAKAG
 C,

(SEQ ID NO: 103)
 QVQLQDS GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKERE
 VGAVSWSGGT TVYADSVLGR FEISRDSARK SVYLQMNLSK PEDTAVYYCA
 ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAAKEA AAKAGC,

(SEQ ID NO: 104)
 QVQLQES GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKERE
 VGAVSWSGGT TVYADSVKGR FTISRDSARK SVYLQMNLSK PEDTAVYYCA
 ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAAKEA AAKAGC,

(SEQ ID NO: 105)
 CGGGVDNKFN KEVGWAFGEI GALPNLNALQ FRAFIISLWD DPSQSANLLA
 EAKKLNDQA PK,
 or

(SEQ ID NO: 106)
 IEVMTQS PSTLSASVGD RVIITCQASQ SIDNWLWYQ QKPGKAPKLL
 IYRASTLASG VPSRFGSGS GAFTLTIS LQPDDEFATYY CQNTGGGVSI
 AFGQGTKLTV LGGGGGSGG GSGGGGSGG GSEVQLVESG GGLVQPGGSL
 RLCTASGFS LSSAAMAVR QAPGKLEWV GIIYDSASTY YASWAKGRFT
 ISRDTSKNTV YLQMNLSRAE DTAVYYCARE RAIFSGDFVL WGQGTSLTVS
 SSPSTPPTPS PSTPPGGC.

[0068] In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 101. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 102. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 103. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 104. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 105. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 106.

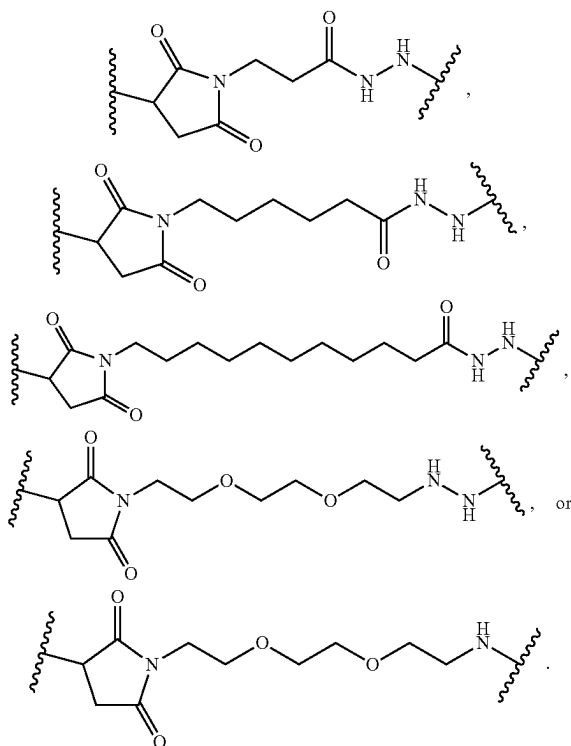
[0069] Each peptide can be linked to the biocompatible polymer by a variety of organic linkers generally known in the art for forming antibody-drug conjugates, such as those provided by BroadPharm of San Diego, CA. Methods for

forming bioconjugate bonds are described in Bioconjugate Techniques, 3rd Edition, Greg T. Hermanson. The organic linkers can be reactive with amines, carbonyls, carboxyl and activated esters, can react via Click-chemistry (with or without copper), or be reactive with thiols.

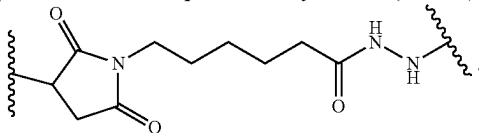
[0070] Representative organic linkers include an amide or disulfide, or are formed from a reactive group such as succinic anhydride, succinimide, N-hydroxy succinimide, N-chlorosuccinimide, N-bromosuccinimide, maleic anhydride, maleimide, hydantoin, phthalimide, and others. The organic linkers useful in the present invention are small and generally have a molecular weight from about 100 Da to about 500 Da containing two functional groups consisting of a maleimide and either an amine or hydrazide. In some embodiments, the peptide is covalently linked to the polymer via a sulfide bond and an organic linker having a

molecular weight of from about 100 Da to about 500 Da. In some embodiments, the organic linker has a molecular weight of from about 100 Da to about 300 Da. In some embodiments, the organic linker comprises a succinimide. In some embodiments, the organic linker is formed using N-beta-maleimidopropionic acid hydrazide (BMPH), N-epsilon-maleimidocaproic acid hydrazide (EMCH), N-aminoethylmaleimide, N-kappa-maleimideundecanoic acid hydrazide (KUMH), hydrazide-PEG2-maleimide, amine-PEG2-maleimide, hydrazide-PEG3-maleimide, or amine-PEG3-maleimide.

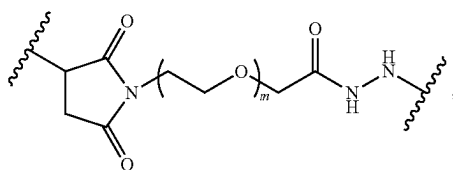
[0071] In some embodiments, the organic linker has the structure:



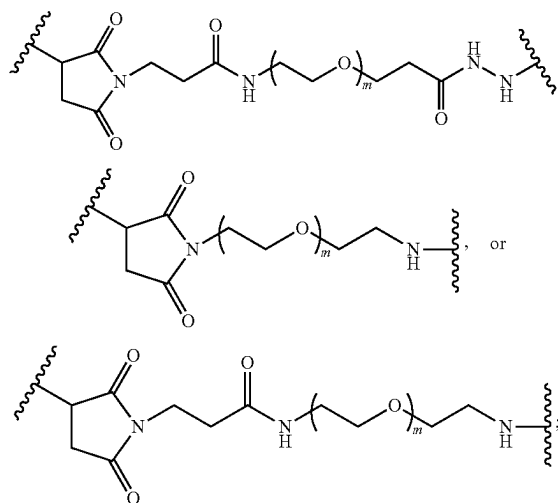
[0072] In some embodiments, the organic linker can be N-epsilon-maleimidocaproic acid hydrazide (EMCH):



[0073] In some embodiments, the organic linker has the structure:

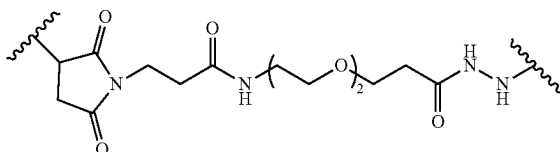


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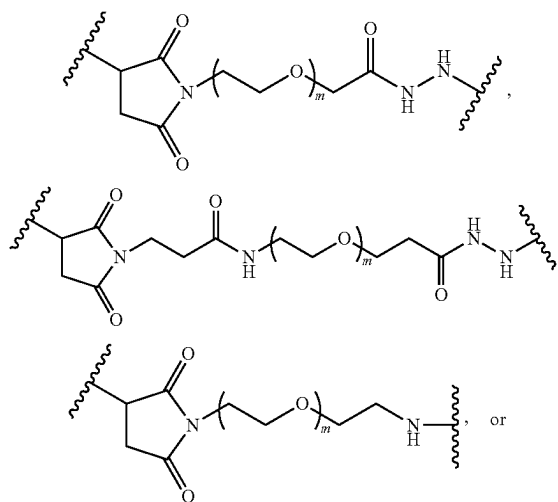
[0074] subscript m is an integer from 1 to 300. In some embodiments, subscript m is an integer from 1 to 100.

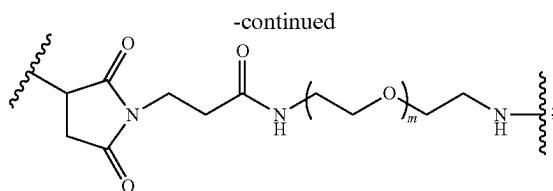
[0075] In some embodiments, the organic linker has the structure:



[0076] The organic linker with the above structure is known as MP2H.

[0077] In some embodiments, each Y is an organic linker having the structure:





and

[0078] subscript m is an integer of from 1 to 300.

[0079] In some embodiments, Z has a molecular weight of from about 0.4 MDa to about 2 MDa. In some embodiments, Z has a molecular weight of from about 0.7 MDa to about 1.5 MDa. In some embodiments, Z has a molecular weight of about 0.8 MDa.

[0080] In some embodiments, the conjugate of Formula V has the structure of Formula Va:

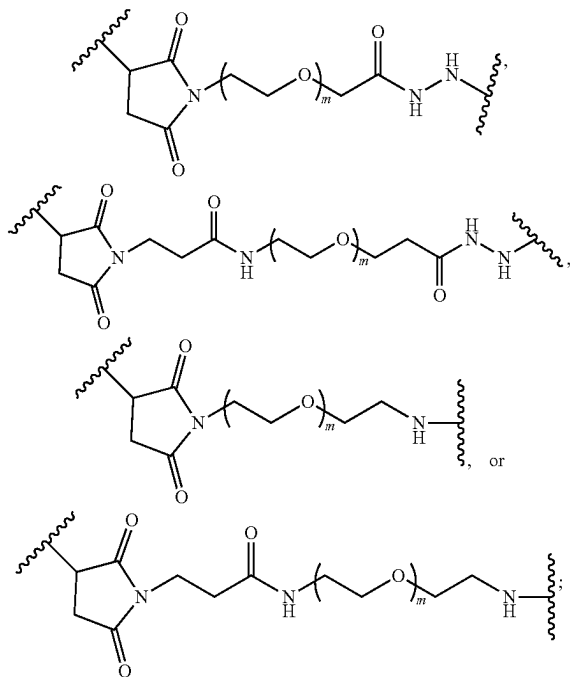


[0081] wherein

[0082] each X^1 is an anti-TNF- α peptide or an anti-interleukin-1 β peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0083] each X^2 is a peptide linker that comprises an alpha-helix;

[0084] each Y is an organic linker having the structure:



[0085] Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and

[0086] subscript m is an integer of from 1 to 300.

[0087] In some embodiments, each X^1 is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 61-73, 81-85, 91-98, 101-109, and 111-118.

[0088] In some embodiments, each X^2 is a peptide linker having an amino acid sequence comprising:

(SEQ ID NO: 21)

AEAAAKEAAAKEAAAKAGC,

(SEQ ID NO: 22)

AEEKKRAEEKKRAEEEGAGC,

(SEQ ID NO: 23)

AEEKKRAEEKKRAEEKKRAEEEGAGC,

(SEQ ID NO: 24)

AEEEEKKKKEEEKKKAGC,

(SEQ ID NO: 25)

AEAAAKEAAAKAGC,

(SEQ ID NO: 26)

PSRLEELRRRLTEGC,

(SEQ ID NO: 27)

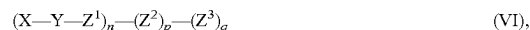
or

AEEEEKKKQEEEAERLRRIQEEMEKERKRREDEERRRKEEEER
RMKLEMAKRKQEEERKKREDEKRRKKAGC.

[0089] In some embodiments, each X^2 is a peptide linker having an amino acid sequence comprising AEAAAKEAAAKEAAAKAGC (SEQ ID NO: 21).

[0090] In some embodiments, each X^1 is a peptide having an amino acid sequence comprising SEQ ID NO: 107, and each X^2 is a peptide linker having an amino acid sequence comprising SEQ ID NO: 21. In some embodiments, each X^1 is a peptide having an amino acid sequence comprising SEQ ID NO: 108, and each X^2 is a peptide linker having an amino acid sequence comprising SEQ ID NO: 21. In some embodiments, each X^1 is a peptide having an amino acid sequence comprising SEQ ID NO: 109, and each X^2 is a peptide linker having an amino acid sequence comprising SEQ ID NO: 21.

[0091] In some embodiments, the conjugate of Formula V is a random polymer of Formula VI:



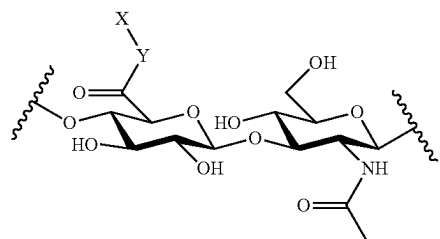
[0092] having a molecular weight of from about 0.1 MDa to about 3 MDa;

[0093] wherein

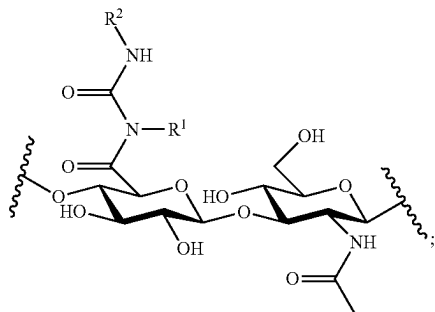
[0094] each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0095] each Y is an organic linker;

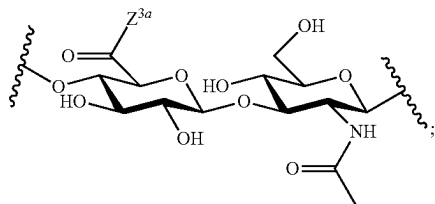
[0096] each $X\text{—}Y\text{—}Z^1$ moiety has the structure:



[0097] each Z^2 has the structure:



[0098] each Z^3 independently has the structure:



[0099] each R^1 and R^2 is independently C_1 - C_6 alkyl, $-(C_1$ - C_6 alkyl)- NR^3R^4 , or C_5 - C_8 cycloalkyl;

[0100] each R^3 and R^4 is independently H or C_1 - C_6 alkyl;

[0101] each Z^{3a} is independently OH or Y' ;

[0102] each Y' is an unreacted organic linker;

[0103] subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;

[0104] subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and

[0105] subscript q is an integer of from 100 to 10000.

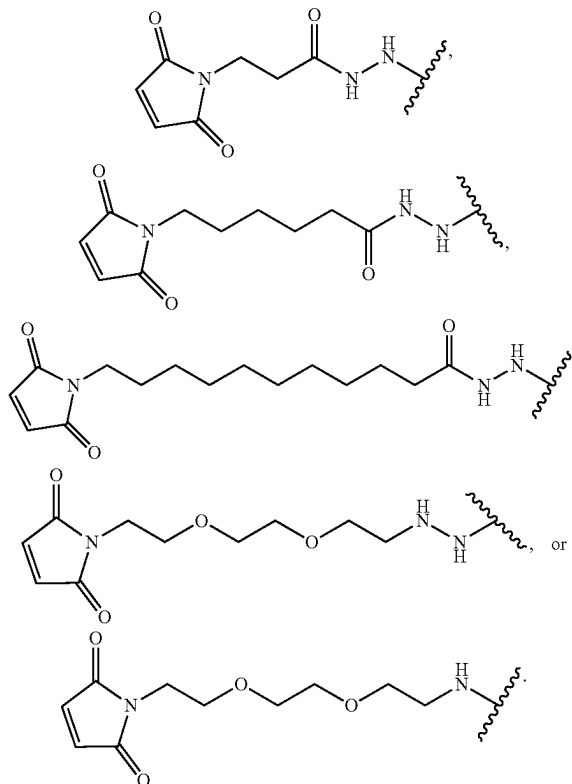
[0106] In some embodiments, each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 61-73, 81-85, 91-98, 101-109, 111-118, and 145-154. In some embodiments, each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 101-109 and 148-154.

[0107] In some embodiments, each R^1 and R^2 is independently C_1 - C_3 alkyl or $-(C_1$ - C_3 alkyl)- NR^3R^4 . In some embodiments, each R^1 and R^2 is ethyl or $-(CH_2)_3$ - NMe_2 . In some embodiments, each R^1 is ethyl; and each R^2 is $-(CH_2)_3$ - NMe_2 . In some embodiments, each R^1 is $(CH_2)_3$ - NMe_2 ; and each R^2 is ethyl.

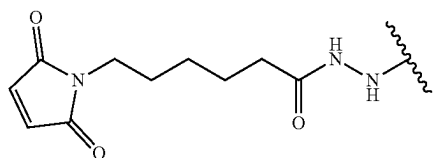
[0108] In some embodiments, each R^3 and R^4 is independently C_1 - C_3 alkyl.

[0109] In some embodiments, preparing the conjugates of the present invention comprises covalently attaching the organic linker to the biocompatible polymer and then covalently attaching the peptide to the organic linker. In some embodiments, after preparing the conjugate of the present invention, unreacted organic linker is present on the biocompatible polymer. The structure of the unreacted organic linker depends on the organic linker and would be understood by a person skilled in the art.

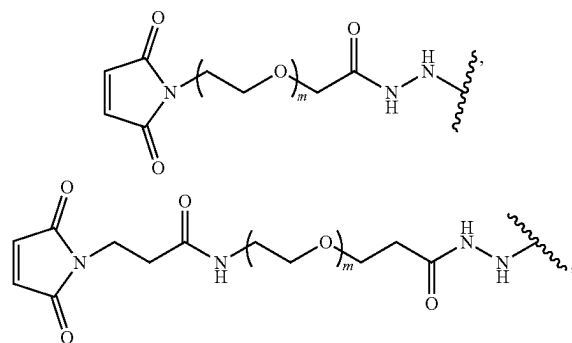
[0110] Representative unreacted organic linkers include, but are not limited to,

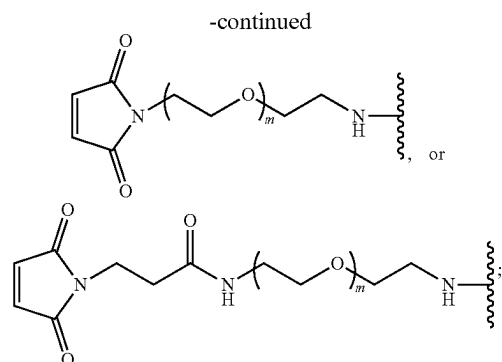


[0111] In some embodiments, the unreacted organic linker has the structure:



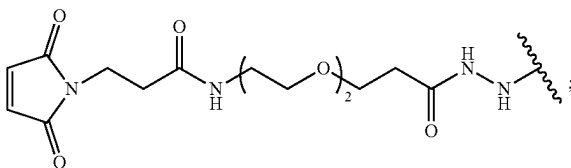
[0112] In some embodiments, the unreacted organic linker has the structure:



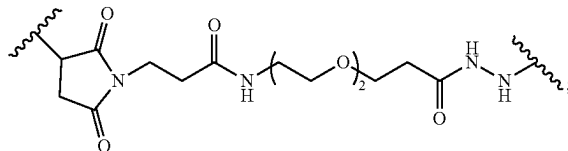


[0113] wherein subscript m is an integer of from 1 to 300. In some embodiments, subscript m is an integer from 1 to 100.

[0114] In some embodiments, the unreacted organic linker has the structure:

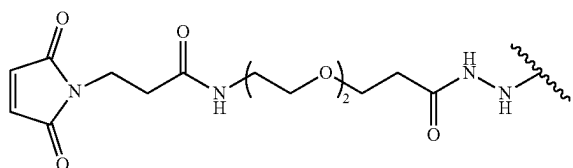


[0115] In some embodiments, the organic linker has the structure:



and

[0116] the unreacted organic linker has the structure:



[0117] In some embodiments, the conjugate is a random polymer of Formula VI:



[0118] having a molecular weight of from about 0.1 MDa to about 3 MDa;

[0119] wherein

[0120] each X is independently an anti-TNF- α or anti-IL-1 β peptide comprising:

(SEQ ID NO: 101)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG

KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA

VYYCAARDGI PTERSVESYN YWGQGTQVTV SSPSTPPTPS PSTPPGGCDD

DDK,

(SEQ ID NO: 102)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG

KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA

VYYCAARDGI PTERSVESYN YWGQGTQVTV SAEAAAKEA AKEAAAKAG

C,

(SEQ ID NO: 103)

QVQLQDS GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKREF

VGAVSWSGGT TVYADSVLGR FEISRDSARK SVYLQMNSLK PEDTAVYYCA

ARPYQKYNWA SASYNVWGQG TQVTSSAEA AKEAAAKEA AAKAGC,

(SEQ ID NO: 104)

QVQLQES GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKREF

VGAVSWSGGT TVYADSVKGR FTISRDSARK SVYLQMNSLK PEDTAVYYCA

ARPYQKYNWA SASYNVWGQG TQVTSSAEA AKEAAAKEA AAKAGC,

(SEQ ID NO: 105)

CGGGVDNKFN KEVGWAFGEI GALPNLNLQ FRAFIISLWD DPSQSANLLA

EAKKLNDQA PK,

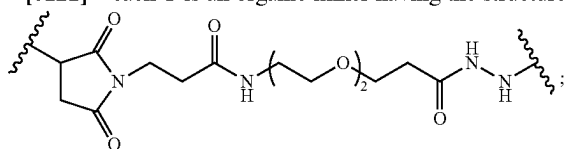
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or

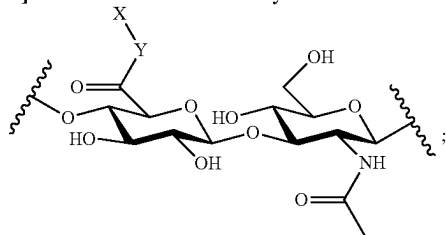
(SEQ ID NO: 106)

EIVMTQS PSTLSASVGD RVIITCQASQ SIDNWLWYQ QKPGKAPKLL
 IYRASTLASG VPSRFGSGSGS GAFTLTISS LQPDDEFATYY CQNTGGGVSI
 AFGQGTCLTV LGGGGGSGGG GSGGGGSGGG GSEVQLVESG GGLVQPGGSL
 RLSCASGFS LSSAAMAVR QAPGKLEWV GIIYDSASTY YASWAKGRFT
 ISRDTSKNTV YLQMNLSRAE DTAVYYCARE RAIFSGDFVL WGQGTLVTVS
 SSPSTPPTPS PSTPPGGC;

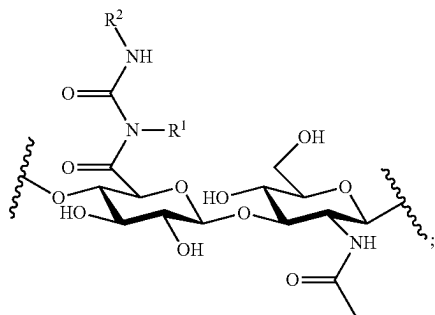
[0121] each Y is an organic linker having the structure:



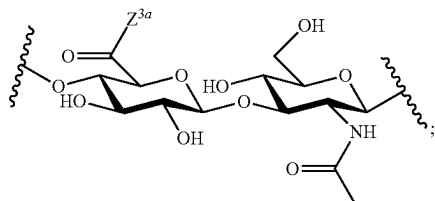
[0122] each X—Y—Z¹ moiety has the structure:



[0123] each Z² has the structure:



[0124] each Z³ independently has the structure:



[0125] each R¹ and R² is independently C₁-C₆ alkyl, —(C₁-C₆ alkyl)-NR³R⁴, or C₅-C₈cycloalkyl;

[0126] each R³ and R⁴ is independently H or C₁-C₆ alkyl;

[0127] each Z^{3a} is independently OH or Y¹;

[0128] each Y¹ is an unreacted organic linker;

[0129] subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;

[0130] subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and

[0131] subscript q is an integer of from 100 to 10000.

[0132] In some embodiments, subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and subscript q is an integer of from 100 to 10000. In some embodiments, subscript n is an integer of from 1 to 1000 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 800 and less than about 8% of the sum of subscripts n, p, and q; and subscript q is an integer of from 100 to 10000. In some embodiments, subscript n is an integer of from 10 to 450 and less than about 15% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 300 and less than about 10% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 240 and less than about 8% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 60 and less than about 2% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 30 and less than about 1% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 15 and less than about 0.5% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000.

[0133] In some embodiments, a conjugate of the present invention is for use in a method of treating uveitis as described herein.

III. Compositions

[0134] In some embodiments, the present invention relates to a pharmaceutical composition as described herein. In some embodiments, the pharmaceutical composition is a pharmaceutical composition comprising a conjugate as described herein, and a pharmaceutically acceptable excipient.

A. Formulation

[0135] For preparing pharmaceutical compositions from the conjugates of the present invention, pharmaceutically acceptable carriers can be either solid or liquid. Solid form preparations include powders, cachets, and dispersible granules. A solid carrier can be one or more substances, which may also act as diluents, binders, preservatives, disintegrating agents, or an encapsulating material. Details on techniques for formulation and administration are well described in the scientific and patent literature, see, e.g., the latest edition of Remington's Pharmaceutical Sciences, Maack Publishing Co, Easton PA ("Remington's").

[0136] In powders, the carrier is a finely divided solid, which is in a mixture with the finely divided active component. In tablets, the active component is mixed with the carrier having the necessary binding properties in suitable proportions and compacted in the shape and size desired. The powders and tablets preferably contain from 5% or 10% to 70% of the conjugates of the present invention.

[0137] Liquid form preparations include solutions, suspensions, and emulsions, for example, water or water/propylene glycol solutions. For parenteral injection, liquid preparations can be formulated in solution in aqueous polyethylene glycol solution.

[0138] Aqueous solutions suitable for oral use can be prepared by dissolving the conjugates of the present invention in water and adding suitable colorants, flavors, stabilizers, and thickening agents as desired. Aqueous suspensions suitable for oral use can be made by dispersing the finely divided active component in water with viscous material, such as natural or synthetic gums, resins, methylcellulose, sodium carboxymethylcellulose, hydroxypropylmethylcellulose, sodium alginate, polyvinylpyrrolidone, gum tragacanth and gum acacia, and dispersing or wetting agents such as a naturally occurring phosphatide (e.g., lecithin), a condensation product of an alkylene oxide with a fatty acid (e.g., polyoxyethylene stearate), a condensation product of ethylene oxide with a long chain aliphatic alcohol (e.g., heptadecaethylene oxycetanol), a condensation product of ethylene oxide with a partial ester derived from a fatty acid and a hexitol (e.g., polyoxyethylene sorbitol mono-oleate), or a condensation product of ethylene oxide with a partial ester derived from fatty acid and a hexitol anhydride (e.g., polyoxyethylene sorbitan mono-oleate). The aqueous suspension can also contain one or more preservatives such as ethyl or n-propyl p-hydroxybenzoate, one or more coloring agents, one or more flavoring agents and one or more sweetening agents, such as sucrose, aspartame or saccharin. Formulations can be adjusted for osmolality.

[0139] Also included are solid form preparations, which are intended to be converted, shortly before use, to liquid form preparations for oral administration. Such liquid forms include solutions, suspensions, and emulsions. These preparations may contain, in addition to the active component,

colorants, flavors, stabilizers, buffers, artificial and natural sweeteners, dispersants, thickeners, solubilizing agents, and the like.

[0140] Oil suspensions can be formulated by suspending the conjugates of the present invention in a vegetable oil, such as arachis oil, olive oil, sesame oil or coconut oil, or in a mineral oil such as liquid paraffin; or a mixture of these. The oil suspensions can contain a thickening agent, such as beeswax, hard paraffin or cetyl alcohol. Sweetening agents can be added to provide a palatable oral preparation, such as glycerol, sorbitol or sucrose. These formulations can be preserved by the addition of an antioxidant such as ascorbic acid. As an example of an injectable oil vehicle, see Minto, J. Pharmacol. Exp. Ther. 281:93-102, 1997. The pharmaceutical formulations of the invention can also be in the form of oil-in-water emulsions. The oily phase can be a vegetable oil or a mineral oil, described above, or a mixture of these. Suitable emulsifying agents include naturally-occurring gums, such as gum acacia and gum tragacanth, naturally occurring phosphatides, such as soybean lecithin, esters or partial esters derived from fatty acids and hexitol anhydrides, such as sorbitan mono-oleate, and condensation products of these partial esters with ethylene oxide, such as polyoxyethylene sorbitan mono-oleate. The emulsion can also contain sweetening agents and flavoring agents, as in the formulation of syrups and elixirs. Such formulations can also contain a demulcent, a preservative, or a coloring agent.

[0141] The compositions of the present invention can also be delivered as microspheres for slow release in the body. For example, microspheres can be formulated for administration via intradermal injection of drug-containing microspheres, which slowly release subcutaneously (see Rao, J. Biomater Sci. Polym. Ed. 7:623-645, 1995; as biodegradable and injectable gel formulations (see, e.g., Gao Pharm. Res. 12:857-863, 1995); or, as microspheres for oral administration (see, e.g., Eyles, J. Pharm. Pharmacol. 49:669-674, 1997). Both transdermal and intradermal routes afford constant delivery for weeks or months.

[0142] In another embodiment, the compositions of the present invention can be formulated for parenteral administration into a body cavity such as intravitreal administration into an eye or the intra-articular space of a joint. The formulations for administration will commonly comprise a solution of the compositions of the present invention dissolved in a pharmaceutically acceptable carrier. Among the acceptable vehicles and solvents that can be employed are water and Ringer's solution, an isotonic sodium chloride. In addition, sterile fixed oils can conventionally be employed as a solvent or suspending medium. For this purpose any bland fixed oil can be employed including synthetic mono- or diglycerides. In addition, fatty acids such as oleic acid can likewise be used in the preparation of injectables. These solutions are sterile and generally free of undesirable matter. These formulations may be sterilized by conventional, well known sterilization techniques. The formulations may contain pharmaceutically acceptable auxiliary substances as required to approximate physiological conditions such as pH adjusting and buffering agents, toxicity adjusting agents, e.g., sodium acetate, sodium chloride, potassium chloride, calcium chloride, sodium lactate and the like. The concentration of the compositions of the present invention in these formulations can vary widely, and will be selected primarily based on fluid volumes, viscosities, body weight, and the like, in accordance with the particular mode of administra-

tion selected and the patient's needs. For IV or intravitreal administration, the formulation can be a sterile injectable preparation, such as a sterile injectable aqueous or oleaginous suspension. This suspension can be formulated according to the known art using those suitable dispersing or wetting agents and suspending agents. The sterile injectable preparation can also be a sterile injectable solution or suspension in a nontoxic parenterally-acceptable diluent or solvent, such as a solution of 1,3-butanediol.

[0143] In another embodiment, the formulations of the compositions of the present invention can be delivered by the use of liposomes which fuse with the cellular membrane or are endocytosed, i.e., by employing ligands attached to the liposome, or attached directly to the oligonucleotide, that bind to surface membrane protein receptors of the cell resulting in endocytosis. By using liposomes, particularly where the liposome surface carries ligands specific for target cells, or are otherwise preferentially directed to a specific organ, one can focus the delivery of the compositions of the present invention into the target cells in vivo. (See, e.g., Al-Muhammed, J. Microencapsul. 13:293-306, 1996; Chonn, Curr. Opin. Biotechnol. 6:698-708, 1995; Ostro, Am. J. Hosp. Pharm. 46: 1576-1587, 1989).

[0144] Lipid-based drug delivery systems include lipid solutions, lipid emulsions, lipid dispersions, self-emulsifying drug delivery systems (SEDDS) and self-microemulsifying drug delivery systems (SMEDDS). In particular, SEDDS and SMEDDS are isotropic mixtures of lipids, surfactants and co-surfactants that can disperse spontaneously in aqueous media and form fine emulsions (SEDDS) or microemulsions (SMEDDS). Lipids useful in the formulations of the present invention include any natural or synthetic lipids including, but not limited to, sesame seed oil, olive oil, castor oil, peanut oil, fatty acid esters, glycerol esters, Labrafil®, Labrasol®, Cremophor®, Solutol®, Tween®, Capryol®, Capmul®, Captex®, and Peceol®.

B. Administration

[0145] The conjugates and compositions of the present invention can be delivered by any suitable means, including oral, parenteral and topical methods. In some embodiments, the delivery method is intravitreal.

[0146] The pharmaceutical preparation is preferably in unit dosage form. In such form the preparation is subdivided into unit doses containing appropriate quantities of the conjugates and compositions of the present invention. The unit dosage form can be a packaged preparation, the package containing discrete quantities of preparation, such as packaged tablets, capsules, and powders in vials or ampoules.

[0147] The conjugates and compositions of the present invention can be co-administered with other agents. Co-administration includes administering the conjugate or composition of the present invention within 0.5, 1, 2, 4, 6, 8, 10, 12, 16, 20, or 24 hours of the other agent. Co-administration also includes administering simultaneously, approximately simultaneously (e.g., within about 1, 5, 10, 15, 20, or 30 minutes of each other), or sequentially in any order. Moreover, the conjugates and compositions of the present invention can each be administered once a day, or two, three, or more times per day so as to provide the preferred dosage level per day.

[0148] In some embodiments, co-administration can be accomplished by co-formulation, i.e., preparing a single pharmaceutical composition including the conjugates and

compositions of the present invention and any other agent. Alternatively, the various components can be formulated separately.

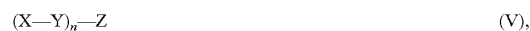
[0149] The conjugates and compositions of the present invention, and any other agents, can be present in any suitable amount, and can depend on various factors including, but not limited to, weight and age of the subject, state of the disease, etc. Suitable dosage ranges include from about 0.1 mg to about 10,000 mg, or about 1 mg to about 1000 mg, or about 10 mg to about 750 mg, or about 25 mg to about 500 mg, or about 50 mg to about 250 mg. Suitable dosages also include about 1 mg, 5, 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 200, 300, 400, 500, 600, 700, 800, 900 or 1000 mg. The composition can also contain other compatible therapeutic agents. The conjugates described herein can be used in combination with one another, with other active agents known to be useful in modulating a glucocorticoid receptor, or with adjunctive agents that may not be effective alone, but may contribute to the efficacy of the active agent.

[0150] In some embodiments, a composition of the present invention is for use in a method of treating uveitis as described herein.

IV. Methods of Treatment

[0151] In some embodiments, the present invention relates to a method and/or use comprising a conjugate or a composition as described herein for the treatment of uveitis in a subject in need thereof. Uveitis is an eye disease that occurs when the middle layer of the eyeball is inflamed, red and/or swollen. This layer, called the uvea, has many blood vessels that nourish the eye. Uveitis can damage vital eye tissue, leading to permanent vision loss. The uveitis can be anterior uveitis, intermediate uveitis, and/or posterior uveitis.

[0152] In some embodiments, the method of the present invention is a method for treating uveitis in a subject in need thereof, comprising administering to the subject a therapeutically effective amount of a conjugate of Formula V:



[0153] wherein

[0154] each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0155] each Y is an organic linker;

[0156] Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and

[0157] subscript n is an integer of from 1 to 1000.

[0158] In some embodiments, each X is independently an anti-TNF- α peptide or an anti-interleukin-1 β peptide.

[0159] In some embodiments, each X is a monoclonal IgG, an IgG fragment, single chain scFv, single-domain heavy-chain VHH, adnectin, affibody, anticalin, DARPin, or an engineered Kunitz-type inhibitor. In some embodiments, each X is a monoclonal IgG. In some embodiments, each X is an IgG fragment. In some embodiments, each X is a single-domain heavy-chain VHH. In some embodiments, each X is a DARPin.

[0160] In some embodiments, each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 61-73, 81-85, 91-98, 101-109, 111-118, and 145-154.

[0161] In some embodiments, each X is a peptide having an amino acid sequence comprising:

(SEQ ID NO: 101)
 QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
 KREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
 VYCAARDGI PTSRSVESYN YWGQGTQVTV SSPSTPPTPS PSTPPGGCDD
 DDK,

(SEQ ID NO: 102)
 QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
 KREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
 VYCAARDGI PTSRSVESYN YWGQGTQVTV SAAEAAAKEA AAKEAAKAG
 C,

(SEQ ID NO: 103)
 QVQLQDS GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKREF
 VGAVSWSGGT TVYADSVLGR FEISRDSARK SVYLMNSLK PEDTAVYYCA
 ARPQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAKEA AAKAGC,

(SEQ ID NO: 104)
 QVQLQES GGGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKREF
 VGAVSWSGGT TVYADSVKGR FTISRDSARK SVYLMNSLK PEDTAVYYCA
 ARPQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAKEA AAKAGC,

(SEQ ID NO: 105)
 CGGGVDNKFN KEVGWAFGEI GALPNLNLALQ FRAFIISLWD DPSQSANLLA
 EAKKLNDQA PK,
 or

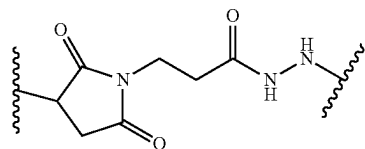
(SEQ ID NO: 106)
 EIVMTQS PSTLSASVGD RVIITCQASQ SIDNWLWYQ QKPGKAPKLL
 IYRASTLASG VPSRFGSGGS GAFTLTISS LQPDDEFATYY CQNTGGGVSI
 AFGQGTCLTV LGGGGGSGGG GSGGGGSGGG GSEVQLVESG GGLVQPGGSL
 RLCTASGFS LSSAAMAVR QAPGKLEWV GIIYDSASTY YASWAKGRFT
 ISRDTSKNTV YLMNSLRAE DTAVYYCARE RAIFSGDFVL WGQGTLVTVS
 SSPSTPPTPS PSTPPGGC.

[0162] Each peptide can be linked to the biocompatible polymer by a variety of organic linkers generally known in the art for forming antibody-drug conjugates, such as those provided by Conju-Probe or BroadPharm of San Diego, CA or Creative Biolabs of Shirley, NY. Methods for forming bioconjugate bonds are described in Bioconjugate Techniques, 3rd Edition, Greg T. Hermanson. The organic linkers can be reactive with amines, carbonyls, carboxyl and activated esters, can react via Click-chemistry (with or without copper), or be reactive with thiols.

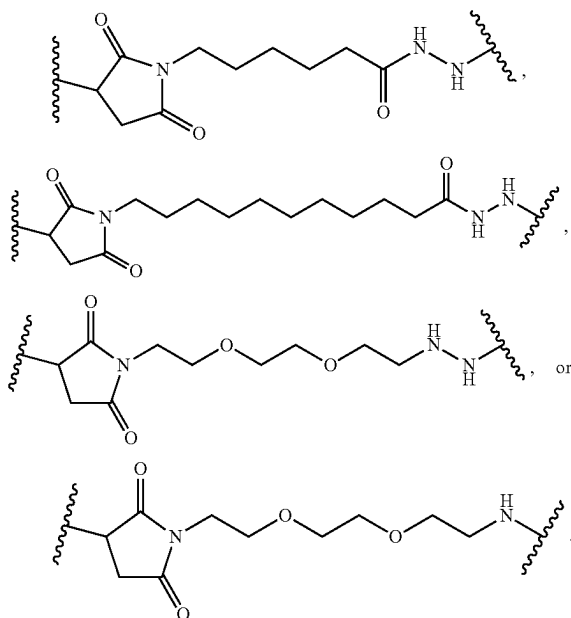
[0163] Representative organic linkers include an amide or disulfide, or are formed from a reactive group such as succinic anhydride, succinimide, N-hydroxy succinimide, N-chlorosuccinimide, N-bromosuccinimide, maleic anhydride, maleimide, hydantoin, phthalimide, and others. The organic linkers useful in the present invention are small and generally have a molecular weight from about 100 Da to about 500 Da containing two functional groups consisting of a maleimide and either an amine or hydrazide. In some embodiments, the peptide is covalently linked to the polymer via a sulfide bond and an organic linker having a

molecular weight of from about 100 Da to about 500 Da. In some embodiments, the organic linker has a molecular weight of from about 100 Da to about 300 Da. In some embodiments, the organic linker comprises a succinimide. In some embodiments, the organic linker is formed using N-beta-maleimidopropionic acid hydrazide (BMPH), N-epsilon-maleimidocaproic acid hydrazide (EMCH), N-epsilon-ethylmaleimide, N-kappa-maleimidoundecanoic acid hydrazide (KUMH), hydrazide-PEG2-maleimide, amine-PEG2-maleimide, hydrazide-PEG3-maleimide, or amine-PEG3-maleimide.

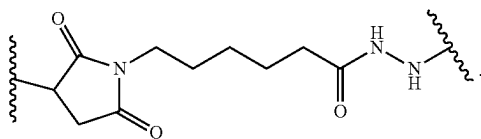
[0164] In some embodiments, the organic linker has the structure:



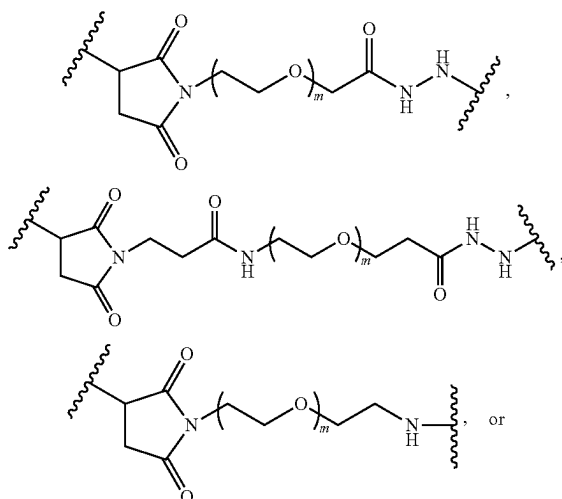
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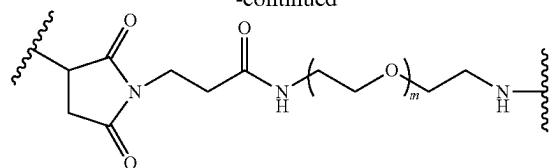
[0165] In some embodiments, the organic linker can be N-epsilon-maleimidocaproic acid hydrazide (EMCH):



[0166] In some embodiments, the organic linker has the structure:



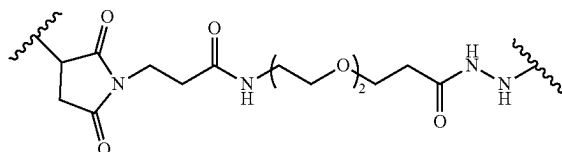
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and

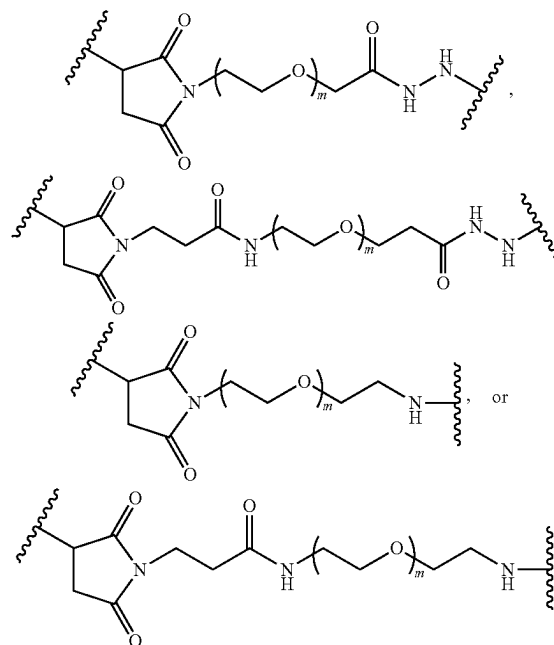
[0167] subscript m is an integer from 1 to 300. In some embodiments, subscript m is an integer from 1 to 100.

[0168] In some embodiments, the organic linker has the structure:



[0169] The organic linker with the above structure is known as MP2H.

[0170] In some embodiments, each Y is an organic linker having the structure:



and

[0171] subscript m is an integer of from 1 to 300.

[0172] In some embodiments, Z has a molecular weight of from about 0.4 MDa to about 2 MDa. In some embodiments, Z has a molecular weight of from about 0.7 MDa to about 1.5 MDa. In some embodiments, Z has a molecular weight of about 0.8 MDa.

[0173] In some embodiments, the conjugate of Formula V has the structure of Formula Va:

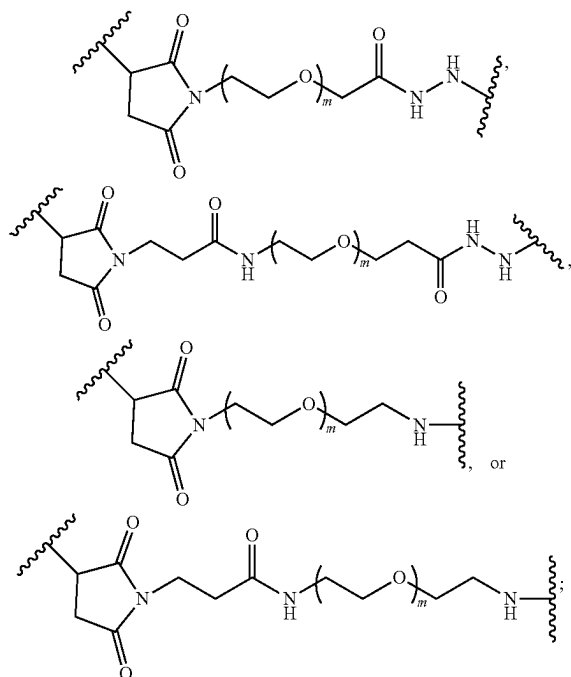


[0174] wherein

[0175] each X^1 is an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0176] each X^2 is a peptide linker that comprises an alpha-helix;

[0177] each Y is an organic linker having the structure:



[0178] Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and

[0179] subscript m is an integer of from 1 to 300.

[0180] In some embodiments, each X^1 is independently an anti-TNF- α peptide or an anti-interleukin-1 β peptide.

[0181] In some embodiments, each X^2 is a peptide linker having an amino acid sequence comprising:

AEAAAKEAAAKEAAAKAGC, (SEQ ID NO: 21)

AEEKKRAEEKKRAEEKEAGC, (SEQ ID NO: 22)

AEEKKRAEEKKRAEEKKRAEEKEAGC, (SEQ ID NO: 23)

AEEKKKKKEEEKKKKKAGC, (SEQ ID NO: 24)

AEAAAKEAAAKAGC, (SEQ ID NO: 25)

PSRLEELRRRLTEGC, (SEQ ID NO: 26)

-continued

or

(SEQ ID NO: 27)

AEEEEKKQEEEEERLRRIQEEMEKERKRREDEERRRKEEEER
RMKLEMEAKRKQEEERKKREDEKRKKKAGC.

[0182] In some embodiments, each X^1 is a peptide having an amino acid sequence comprising SEQ ID NO: 107, and each X^2 is a peptide linker having an amino acid sequence comprising SEQ ID NO: 21. In some embodiments, each X^1 is a peptide having an amino acid sequence comprising SEQ ID NO: 108, and each X^2 is a peptide linker having an amino acid sequence comprising SEQ ID NO: 21. In some embodiments, each X^1 is a peptide having an amino acid sequence comprising SEQ ID NO: 109, and each X^2 is a peptide linker having an amino acid sequence comprising SEQ ID NO: 21.

[0183] In some embodiments, the conjugate of Formula V is a random polymer of Formula VI:



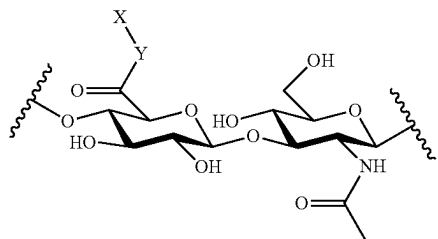
[0184] having a molecular weight of from about 0.1 MDa to about 3 MDa;

[0185] wherein

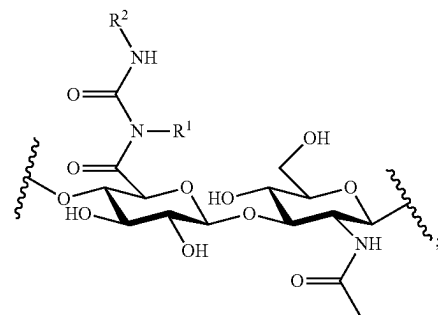
[0186] each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

[0187] each Y is an organic linker;

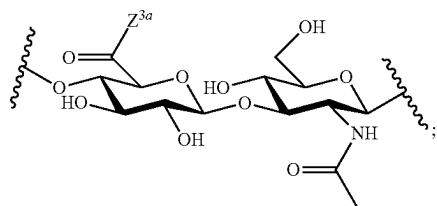
[0188] each $X-Y-Z^1$ moiety has the structure:



[0189] each Z^2 has the structure:



[0190] each Z^3 independently has the structure:



[0191] each R^1 and R^2 is independently C_1 - C_6 alkyl, $-(C_1$ - C_6 alkyl)- NR^3R^4 , or C_5 - C_8 cycloalkyl;

[0192] each R^3 and R^4 is independently H or C_1 - C_6 alkyl;

[0193] each Z^{3a} is independently OH or Y' ;

[0194] each Y' is an unreacted organic linker;

[0195] subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;

[0196] subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and

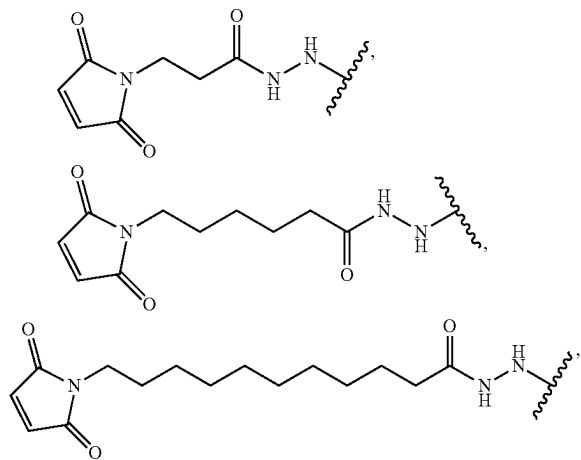
[0197] subscript q is an integer of from 100 to 10000.

[0198] In some embodiments, each R^1 and R^2 is independently C_1 - C_3 alkyl or $-(C_1$ - C_3 alkyl)- NR^3R^4 . In some embodiments, each R^1 and R^2 is ethyl or $-(CH_2)_3-NMe_2$. In some embodiments, each R^1 is ethyl; and each R^2 is $-(CH_2)_3-NMe_2$. In some embodiments, each R^1 is $-(CH_2)_3-NMe_2$; and each R^2 is ethyl.

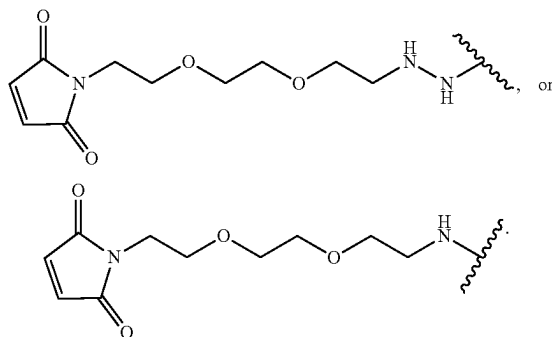
[0199] In some embodiments, each R^3 and R^4 is independently C_1 - C_3 alkyl.

[0200] In some embodiments, preparing the conjugates of the present invention comprises covalently attaching the organic linker to the biocompatible polymer and then covalently attaching the peptide to the organic linker. In some embodiments, after preparing the conjugate of the present invention, unreacted organic linker is present on the biocompatible polymer. The structure of the unreacted organic linker depends on the organic linker and would be understood by a person skilled in the art.

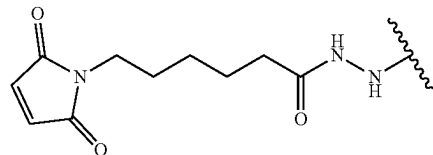
[0201] Representative unreacted organic linkers include, but are not limited to,



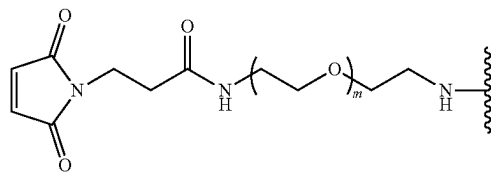
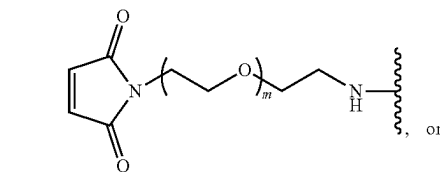
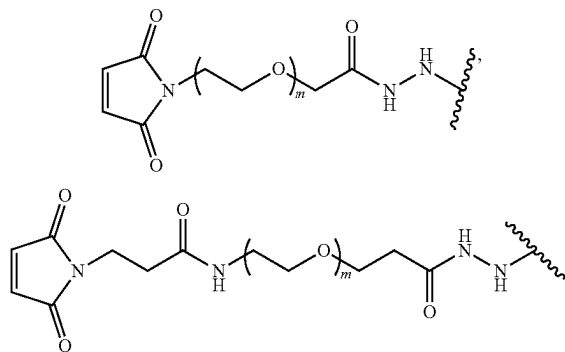
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[0202] In some embodiments, the unreacted organic linker has the structure:

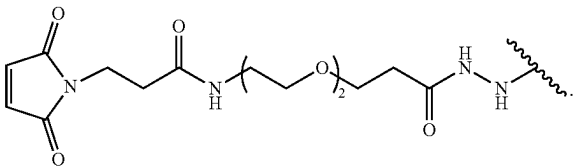


[0203] In some embodiments, the unreacted organic linker has the structure:



[0204] wherein subscript m is an integer of from 1 to 300. In some embodiments, subscript m is an integer from 1 to 100.

[0205] In some embodiments, the unreacted organic linker has the structure:



[0206] In some embodiments, subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and subscript q is an integer of from 100 to 10000. In some embodiments, subscript n is an integer of from 1 to 1000 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 800 and less than about 8% of the sum of subscripts n, p, and q; and subscript q is an integer of from 100 to 10000. In some embodiments, subscript n is an integer of from 10 to 450 and less than about 15% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 300 and less than about 10% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 240 and less than about 8% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 60 and less than about 2% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In

some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 30 and less than about 1% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000. In some embodiments, subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q; subscript p is an integer of from 1 to 15 and less than about 0.5% of the sum of subscripts n, p, and q; and subscript q is an integer of from 1000 to 3000.

[0207] In some embodiments, the uveitis is chronic uveitis. In some embodiments, the uveitis is chronic non-infectious uveitis.

[0208] In some embodiments, the method comprises intravitreal administration. In some embodiments, the method comprises multiple administrations of the conjugate. In some embodiments, the method comprises administering the conjugate every month, every two months, or every three months. In some embodiments, the method comprises administering the conjugate twice or three times yearly. In some embodiments, the method comprises administering the conjugate yearly.

[0209] In some embodiments, the method of the present invention is a method for treating chronic non-infectious uveitis in a subject in need thereof, comprising administering to the subject a therapeutically effective amount of the conjugate that is a random polymer of Formula VI:



[0210] having a molecular weight of from about 0.1 MDa to about 3 MDa;

[0211] wherein

[0212] each X is independently an anti-TNF- α or anti-IL-1 β peptide comprising:

(SEQ ID NO: 101)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
KEREVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
VYYCAARDGI PTERSVE SYN YWGQGTQVTV SSPSTPPTPS PSTPPGGCDD
DDK,

(SEQ ID NO: 102)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
KEREVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
VYYCAARDGI PTERSVE SYN SAEAAKEA AKEAAKAG
C,

(SEQ ID NO: 103)

QVQLQDS GGGLVQAGGS LRLSCAASGG TFSSIIAMWF RQAPGKERE
VGA VSWSGGT TVYADSVLGR FEISRDSARK SVYLQMNSLK PEDTAVYYCA
ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AKEAAKEA AAKAGC,

(SEQ ID NO: 104)

QVQLQES GGGLVQAGGS LRLSCAASGG TFSSIIAMWF RQAPGKERE
VGA VSWSGGT TVYADSVKGR FTISRDSARK SVYLQMNSLK PEDTAVYYCA
ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AKEAAKEA AAKAGC,

-continued

(SEQ ID NO: 105)

CGGGVDNKFN KEVGWAFGEI GALPNLNALQ FRAFIISLWD DPSQSANLLA

EAKKLNDQA PK,
or

(SEQ ID NO: 106)

EIVMTQS PSTLSASVGD RVIITCQASQ SIDNWLWSYQ QKPGKAPKLL

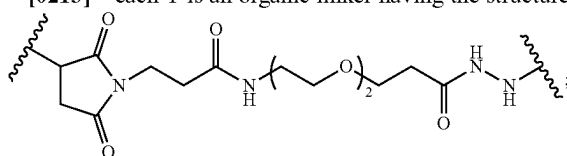
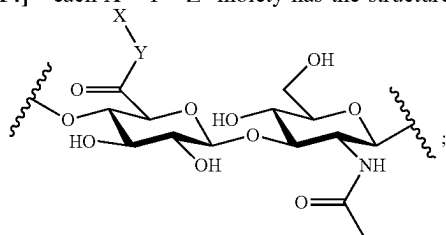
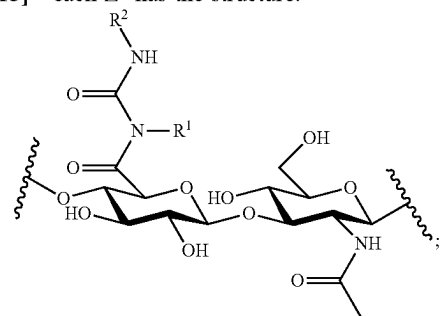
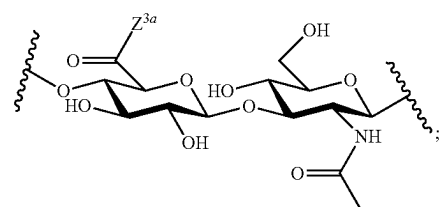
IYRASTLASG VPSRFSGSGS GAFTLTIS LQPDDFATYY CQNTGGGVSI

AFQGQTKLTV LGGGGSGSGG GSGGGSGGG GSEVQLVESG GGLVQPGGSL

RLSCTASGFS LSSAAMAVR QAPKGLEWV GIIYDSASTY YASWAKGRFT

ISRDTSKNTV YLQMSLRAE DTAVYYCARE RAIFSGDFVL WGQGLTVTS

SSPSTPPTPS PSTPPGGC;

[0213] each Y is an organic linker having the structure:**[0214]** each X—Y—Z¹ moiety has the structure:**[0215]** each Z² has the structure:**[0216]** each Z³ independently has the structure:**[0217]** each R¹ and R² is independently C₁-C₆ alkyl, —(C₁-C₆ alkyl)-NR³R⁴, or C₅-C₈cycloalkyl;**[0218]** each R³ and R⁴ is independently H or C₁-C₆ alkyl;**[0219]** each Z^{3a} is independently OH or Y';**[0220]** each Y' is an unreacted organic linker;**[0221]** subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;**[0222]** subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and**[0223]** subscript q is an integer of from 100 to 10000.**[0224]** In some embodiments, the random polymer of Formula VI has a molecular weight of from about 0.4 MDa to about 2 MDa. In some embodiments, the random polymer of Formula III has a molecular weight of from about 0.7 MDa to about 1.5 MDa. In some embodiments, the random polymer of Formula III has a molecular weight of about 0.8 MDa.**[0225]** In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 101. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 102. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 103. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 104. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 105. In some embodiments, each X is a peptide having an amino acid sequence comprising SEQ ID NO: 106.**[0226]** In some embodiments, a use of the present invention comprises the preparation of a medicament for a method of treating uveitis as described herein.**[0227]** In some embodiments, the subject is a human.**[0228]** In some embodiments, a use of the present invention is a use for treating uveitis comprising a conjugate or pharmaceutical composition as described herein.**[0229]** In some embodiments, a pharmaceutical composition of the present invention is a pharmaceutical composition for use in treating uveitis comprising a conjugate as described herein.**[0230]** In some embodiments, a conjugate of the present invention is a conjugate for use in treating uveitis as described herein.

V. EXAMPLES

[0231] Certain abbreviations and acronyms are used in describing the experimental details. Although most of these would be understood by one skilled in the art, the Table below contains a list of many of these abbreviations and acronyms.

TABLE 1

List of abbreviations and acronyms.	
Abbreviation	Meaning
aH	alpha-helix
BLI	biolayer interferometry
CBB	Coomassie brilliant blue
Da	daltons
DLS	dynamic light scattering
DMSO	dimethyl sulfoxide
DMTMM	4-(4,6-dimethoxy-1,3,5-triazin-2-yl)-4-methylmorpholinium chloride
DPBS	Dulbecco's phosphate buffered saline
DTT	dithiothreitol
EAU	experimental autoimmune uveoretinitis
EDC	1-ethyl-3-(3-dimethylaminopropyl)carbodiimide
EDTA	ethylenediaminetetraacetic acid
EIU	endotoxin-induced uveitis
ELISA	enzyme-linked immunosorbent assay
FPLC	fast protein liquid chromatography
HA or HyA	hyaluronic acid
IL	interleukin
ITV	intravitreal
kDa	kilodaltons
MDa	megadaltons
MES	2-(N-morpholino)ethanesulfonic acid
MVP	multivalent protein
MW	molecular weight
MWCO	molecular weight cutoff
NHS	N-hydroxysuccinimide
PBS	phosphate buffered saline
RPM	revolutions per minute
RT	room temperature
SEC	size-exclusion chromatography
SEC MALS	size-exclusion chromatography multi-angle light scattering
TCEP	tris(2-carboxyethyl)phosphine
TNF α	tumor necrosis factor alpha

Example 1. Preparation of Peptides

[0232] Biologically active peptides were prepared optionally with a C-terminal peptide linker for attachment to the polymer.

TABLE 2

Peptides		
SEQ ID NO:	Protein	Type
101	Hu_anti-TNF α -rigid	VHH
102	Hu_anti-TNF α -aH	VHH
103	mu_anti-TNF α -aH	VHH
104	mu_anti-TNF α _3Mut-aH	VHH
105	anti-TNF α	affibody
106	anti-IL-1 β -rigid	scFv
107	Hu_anti-TNF α	VHH
108	Hu_anti-TNF α	VHH
109	Hu_anti-TNF α	VHH

Example 2. Preparation of Purified Thiol Reactive Hyaluronic Acid Conjugate Intermediates

[0233] Hyaluronic acid (HA, 830 kDa) was suspended in water or 0.1 M 2-(N-morpholino)ethanesulfonic acid buffer

pH 5.7 at 4 mg/mL by gentle rotation or mixing with nutation overnight at RT. To 3 mg (3.6 nmol, amount will vary based on polymer composition and MW) of HA in solution is added hydroxybenzotriazole (HOBt) hydrate as a ~5-100 mg/mL stock solution in DMSO, thiol reactive linker agent (e.g., hydrazide-X-thiol-reactive-group or amine-X-thiol-reactive-group, for example, MP2H or EMCH) in 10-100% DMSO (10-100 mg/mL stock), and a coupling agent (1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) or 4-(4,6-Dimethoxy-1,3,5-triazin-2-yl)-4-methylmorpholinium chloride) as a ~1-0.05 g/mL stock in water or 0.1 M MES buffer pH 5.7. The molar equivalents for each reactant per mole of HA and per carboxylate for different methods of performing the reaction, and example methods are described in the table below:

TABLE 3

Relative Ratios of Coupling Agent, Catalyst, and Linkers in Methods		
Reactant	Method 1	Method 2
Coupling agent (EDC)	500-750	9500-12000
HOBt	3000	50
Linker agent	3000	500-1000

[0234] Solution was mixed with gentle pipetting between each reagent addition and the final reaction volume was raised to 1 mL with buffer. The final mixture was allowed to react at room temperature for 45 min to 2 h with nutating mixer depending on Method. After the reaction, the thiol reactive biopolymer was purified using 7 kDa MWCO 5-10 mL Zeba desalting spin column equilibrated with 10% v/v glycerol pH 6.5 DPBS, and 0.01% v/v polysorbate 20 (optional), loaded with crude reaction at 20% volume of resin. The desired intermediate was eluted into clean conical tube using centrifuge at RT, elution time ~25-60 minutes. The intermediate was used immediately for reaction with thiol or aliquotted and flash frozen on dry ice. Maleimide concentration and number of modifications per polymer was determined using UV absorbance, NMR, or a modified Ellman's reaction assay.

[0235] Alternatively, reaction pH or equivalents of hydrazide linker, catalyst, and coupling agent (EDC) were altered higher or lower to increase or decrease the number of thiol reactive small molecule linkers covalently linked per biopolymer (valency).

[0236] Alternative coupling reagents can be used in place of EDC and HOBt such as DMTMM or oxyma. Activated biopolymer intermediate can also be purified away from reactants using size exclusion chromatography, other desalting columns, tangential flow filtration, ion exchange chromatography, dialysis, or alcohol/acetone precipitation.

Example 3. Conjugate Preparation

[0237] A fixed concentration of peptide was combined with the polymer at various defined feed ratios in PBS and allowed to react at either 4° C. or ambient temperature for at least 4 and 2 hours respectively with rotation or nutating mixing (most reactions are ran at RT to improve solubility). Before the conjugation reaction, 10-100 equivalents of a reducing agent such as DTT or TCEP HCl were added per protein equivalent to reduce any disulfide bridging between

peptides. This was removed from the protein solution prior to conjugation by a desalting column or buffer exchange or was added to the conjugation reaction directly in the form of TCEP immobilized on polymeric beads. During the conjugation reaction, one or more of the following was added to improve the reaction efficiency: 0.5-10 mM EDTA to minimize free thiol oxidation, tween20, carbohydrate, or glycerol to stabilize protein and/or help reduce non-specific interactions between protein and activated biopolymer, increased or decreased salt concentration to stabilize protein and/or help reduce non-specific interactions between protein and activated biopolymer. Unreacted peptide was removed from the peptide-polymer conjugates by one or more of the following methods: dialysis with 50-1000 kDa MWCO against an appropriate buffer (pH should be >1 unit above or below the pI of peptide) for two times for 4 hours each and once for at least 4 hours at 4° C.-room temperature, tangential flow filtration against DPBS pH 6-8, or 50 mM tris 150 mM NaCl pH 8-8.5 with EDTA and tween or other additives like trehalose, depending on peptide, FPLC polishing using a

size exclusion column, FPLC polishing with an affinity chromatography column designed to bind the polymer component of the conjugate, or selective precipitation of the conjugates. If reaction efficiency was high enough (<4% unreacted protein present) purification may not be necessary. [0238] Alternatively, to each solution of activated polymer, the peptide was added at a suitable peptide:polymer molar feed ratio and Tween-20 to a final concentration of 0.01%-0.03% (optional). The solution was allowed to react for 2 hours to overnight while agitating by rotation (~5 RPM) or nutation at ambient temperatures. Unreacted peptides were removed by dialysis using 100-1000 kDa MWCO membranes against phosphate buffered saline or equivalent citrate or succinate buffered saline (pH and buffer salt used depends on peptide) with 0.01-0.03% Tween-20 (optional) for three to five times for 4-18 hours each at 4° C.-room temperature. Alternative methods include tangential flow filtration against appropriate buffer or FPLC polishing using a size exclusion column. Additives like tween20, EDTA, and carbohydrates were optionally added to enhance protein stability, depending on peptide.

TABLE 4

Reaction conditions for hyaluronic acid-protein conjugates						
Conjugate ID	Method	Reactive Thiol (μM)	Peptide	Polymer (μM)	Peptide (μM)	% unconjugated peptide after reaction
1	2	—	anti-TNFα VHH (human) - rigid (SEQ ID NO: 101)	—	—	—
2	2	75.85	anti-TNFα VHH (human) - rigid (SEQ ID NO: 101)	0.54	38.02	—
3	1	299.21	anti-TNFα VHH (human) - aH (SEQ ID NO: 102)	1.85	329.20	—
4	1	166.59	anti-TNFα VHH (human) - aH (SEQ ID NO: 102)	1.23	333.17	1.8
5	1	190.59	anti-TNFα VHH (mouse) - aH (SEQ ID NO: 103)	1.15	209.60	<LOD
6	1	412.78	anti-TNFα 3MUT VHH (mouse) - aH (SEQ ID NO: 104)	3.03	454.06	—
7	1	303.91	anti-TNFα 3MUT VHH (mouse) - aH (SEQ ID NO: 104)	2.22	334.32	13.3
8	2	—	anti-TNFα Affibody (SEQ ID NO: 105)	1.11	64.35	—
9	2	—	anti-IL-1β scFv - rigid (SEQ ID NO: 106)	0.09	5.36	—
10	1	515.02	anti-TNFα 3MUT VHH (mouse) - aH (SEQ ID NO: 104)	3.05	566.53	12.2
11	1	646.13	anti-TNFα VHH (human) - aH (SEQ ID NO: 102)	3.20	710.74	3.8
12	2	610.67	Hu αTNFα 3MUT aH_CYS (SEQ ID NO: 149)	3.03	671.73	4.7
13	2	610.52	Hu αTNFα 5MUT aH_CYS (SEQ ID NO: 150)	3.03	671.57	3.2
14	2	430.36	Hu αTNFα 7MUT aH_CYS (SEQ ID NO: 151)	2.13	473.40	3.1

TABLE 4-continued

Reaction conditions for hyaluronic acid-protein conjugates						
Conjugate ID	Method	Reactive Thiol (μM)	Peptide	Polymer (μM)	Peptide (μM)	% unconjugated peptide after reaction

<LOD = below limit of detection

[0239] The conjugates in the following table were generated using hyaluronic acid (830 kDa or 850 kDa lots). After purification, the products of the conjugation reactions were analyzed by SDS-PAGE separation to confirm that <20% of the peptide monomer had entered the resolving gel and that >90% of the peptide was present as a macromolecular conjugate at the top of the stacking gel (FIGS. 1A-1F). The reaction products were further analyzed for protein concentration, percent unconjugated peptide, conjugated peptide, valency (molar ratio of conjugated peptide to polymer), and hydrodynamic radius (Rh). Protein concentration was determined based on spectrophotometry at A280, percent unconjugated protein was determined by densitometric analysis of the SDS-PAGE gels, and hydrodynamic radius was measured using dynamic light scattering (DLS).

TABLE 5

Conjugates								
conjugate #	SEQ ID NO:	Method #	Organic linker	Protein Valency (UV)	Protein conc. (mg/mL)	K _d (nM, BLI)	R _h (nm, DLS)	R _h (% PD)
1	101	2	EMCH	9	0.094	—	—	—
2	101	2	MP2H	55	0.39 ± 0.02	≤0.001	—	—
3	102	1	MP2H	65	1.24 ± 0.01	≤0.001	87.09 ± 9.87	54 ± 8
4	102	1	MP2H	120	2.46 ± 0.02	—	—	—
5	103	1	MP2H	121	2.25 ± 0.08	≤0.001	—	—
6	104	1	MP2H	51	4.77 ± 0.02	≤0.001	76.8 ± 1.58	—
7	104	1	MP2H	98	2.67 ± 0.02	≤0.001	76.42 ± 5.89	—
8	105	2	EMCH	21	0.15 ± 0.01	0.152	—	—
9	106	2	EMCH	15	0.030 ± 0.00	0.353	—	—
10	104	1	MP2H	96.5	3.80 ± 0.07	≤0.001	99.14 ± 5.04	71.04 ± 13.13
11	102	1	MP2H	131	6.80 ± 0.17	≤0.001	135.80 ± 10.77	78.26 ± 21.97
12	149	2	MP2H	99	4.33 ± 0.01	≤0.001	70.12 ± 4.98	48.74 ± 14.98
13	150	2	MP2H	102	6.23 ± 0.23	≤0.001	71.64 ± 7.64	41.65 ± 14.67
14	151	2	MP2H	100	3.02 ± 0.31	0.0115	96.10 ± 9.45	65.02 ± 18.70

Example 4. Potency of MVPs Comprising Anti-Inflammatory Proteins

[0240] As one example, we engineered anti-inflammatory proteins containing a peptide linker and thiol linker for conjugation. These antibodies were conjugated to HyA to generate multivalent conjugates at a range of valencies and on different polymer backbones and sizes. Biolayer interferometry (BLI) was performed to quantify binding kinetics

for purified MVP as an assessment of bioactivity using a GatorPrime (Gator Bio) or similar instrument and either streptavidin coated probes (Cat #160002) for AVI-tagged ligands or anti-Human Fc (Cat #160003) coated probes for Fc-tagged ligands. All analytes and ligands were diluted in BLI Buffer (1xDPBS, 0.1% w/v BSA and 0.1% v/v polysorbate 20, 0.2 μm filtered). The appropriate ligand for each analyte as noted in Table 6 was first resuspended and stored for long term use according to the manufacturer's directions.

The unconjugated analytes were diluted to a top concentration in the range of 5 μ M to 1 nM. The multivalent conjugates were diluted to a top concentration of 50-1.0 nM based on the entire multivalent conjugate molecular weight ((protein MW \times valency)+polymer mw). The concentration range for each ligand-analyte pair is what demonstrates dose dependence binding affinity in pilot range-finding experiments over a wide titration of concentrations from 10 μ M-1 nM.

[0241] All reagents were equilibrated to room temperature before use for at least 30 minutes. Two probes per sample (one for kinetic assay and one for ligand free control) were equilibrated in 250 μ L BLI buffer (PBS pH 7.4, 0.2% Tween and 0.2% BSA filtered at 0.2 μ m) for at minimum 10 min in a Gator Bio Max plate. Ligands were diluted to a fixed concentration of 25-100 nM based on performance in pilot reactions in BLI buffer. Analytes were prepared at the top concentration determined in pilot reactions in BLI buffer and serially diluted 1:3 two to five more times using BLI buffer (Table 6). Black flat-bottom non-coated 96 well plates (Greiner Bio One Cat #655209 or similar) were loaded column-wise with 200 L of ligand, analyte dilutions and one column of BLI buffer for each column of ligand and analyte. One well in each column of analyte was BLI buffer to be used as a blank for reference subtraction. The sample plate was placed in the Gator on a tilted platform set to 25° C. Gator K assay loading and kinetic steps were set up using double reference and step times shown in Table 7. Ligand was loaded until signal reaches between 0.4 and 0.6 nm then returned to buffer column for a baseline measurement for 60-90 s. Next, the kinetic reads were started using the step parameters. When kinetic reads were complete with ligand-loaded probes, a ligand free control was run using new probes that were not loaded with the ligand. The same kinetic assay timing and same sample wells were used that were analyzed with ligand loaded probes. This data was used to correct for any non-specific interactions between the sample and probe.

[0242] When kinetic assay was complete, data was analyzed using the Gator software. The raw data was corrected to include the association time after 1 second to 180 seconds. The Y-axis was aligned to the beginning of the association step and interstep correction was used. Savitzky-Golay filtering of data was used. The samples were set for a double reference by denoting which probes and wells were buffer references in the software. Then, the reference subtraction formula for each assay was edited so that for each assay it was a double reference with the equation of (Kinetic Assay well–Ligand Free Assay well)–(Kinetic Assay buffer reference well–Ligand free assay buffer well). All titrations of the same MVP were grouped by color and the parameters adjusted to a 1:1 binding model that included both association and dissociation with global, Rmax unlinked fitting. The window of interest was moved to include only 100 seconds of dissociation. The binding curve was fitted and checked that the residuals did not vary from the actual curve more than 10%, that the full R² is >0.98 and the Full X² is <3.0. The kinetics and variables K_D, K_{on} and response were noted.

TABLE 6

BLI Ligands and analyte pairings				
BLI Ligand	Tags	Analyte	Supplier	Catalog Number
Human TNF α	Avi, His	anti-TNF α	Acro Biosystems	TNA-H82E3
Human IL-1 β	Fc	Anti-IL-1 β	Acro Biosystems	ILA-H525c

TABLE 7

BLI method parameters and results specifications for kinetic quantitation		
Parameter	Wells Used	Step time (s) or info
Probe equilibration	Buffer in Max Plate	>600
Basic Parameters		5 Hz, 30 s equilibration, 1000 rpm shaking
Buffer		PBS pH 7, 0.2% Tween and 0.2% BSA filtered at 0.2 μ m
BLI Experiment Parameters		
Baseline	Buffer Column 1	60
Ligand loading	100-25 nM Ligand	When loading signal is 0.4-0.6
Baseline	Buffer Column 1	90
Association	MVP Sample(s)	180
Dissociation	Buffer Column 2	300
Ligand free control (with blank probes)		
Baseline	Buffer Column 2	90
Association	MVP Sample(s)	180
Dissociation	Buffer Column 2	300

Example 5. Hydrodynamic Radius of MVPs

[0243] Anti-inflammatory agents containing a peptide linker and thiol linker for conjugation were engineered. These agents were conjugated to HyA to generate multivalent conjugates at a range of valencies and on different polymer backbones and sizes. Dynamic light scattering (DLS) was performed to quantify the hydrodynamic radius (Rh) for purified unconjugated protein or MVP as an assessment of size using either a Wyatt Dynapro single cuvette Nanostar, plate reader or similar instrument.

[0244] Samples were equilibrated to room temperature for at least 30 minutes. The solution was diluted in 0.1 μ m filtered formulation buffer without polysorbate 20 to a final concentration of 100 nM in 100 μ L (typically a 1:10 dilution) and mixed by gentle trituration in a 1.5 mL centrifuge tube or up to 30 minutes on a neutator. Large aggregates and dust particles could be removed by spinning the tubes at 5000 g for 5 minutes in a centrifuge. For single cuvette measurements in a NanoStar, a 40 μ L sample of the sample solution was loaded into a Wyatt Technology disposable microcuvette (Wyatt Cat #WNDMC) with cap, tapped to remove bubbles, and placed into the instrument for analysis. For multiple readings using the plate reader, 25-35 μ L of sample was added to a clear bottomed black well 384 well plate (Corning Cat #P8802-384 or similar). Bubbles in the sample wells were removed. Instrument settings for this and

the other sample analyses by DLS in this document are presented in Table 8. DLS acquisition parameters are shown in Table 8 and results specifications are in Table 9. Representative DLS intensity plot for purified, filtered MVP is shown in FIG. 3.

TABLE 8

DLS Acquisition Parameters	
Instrument	
Laser wavelength	664 nm
Laser power	100%
Auto attenuation	On
Temp	25° C.
Acquisition time (s)	5-10
Acquisition #	5
Fixed Parameters	
Correlation function low cutoff	1.5 μ s
Correlation function high cutoff (μ s)	103000 μ s
Peak radius low cutoff	0 nm
Peak radius high cutoff	1E6 nm
Analysis type	Dynals
Measurement time limit factor	5
Auto-attenuation time limit	60
Sample	
Mw-R model	Globular proteins
Solvent	PBS
dn/dc	0.185
Rg model	Sphere

TABLE 9

DLS results specification for Multivalent Protein Conjugates		
Peak	Unconjugated Proteins	Multivalent Protein Conjugates
0-25 nm (unconjugated)	>80% intensity	<13% intensity
25-1000 nm	>13% intensity	>80% intensity
>1000 nm	<2% intensity	<2% intensity

Example 6. Stability of MVPs in a Vitreous Mimetic Buffer

[0245] Thermal stability was used as a surrogate to evaluate anti-inflammatory MVPs that may serve as a long term therapeutic and to compare relative stabilities of different constructs. Unconjugated antibodies were diluted to 1.0 mg/mL and anti-inflammatory MVPs to 0.5 mg peptide/mL in formulation buffer. 3×30 μ L of each sample was placed in a UV-VIS compatible 384 well plate (Greiner Bio-One Cat #781801 or similar), bubbles were removed, and the plate sealed with UV transparent sealing tape (Greiner Bio-One Cat #676070 or similar). A plate reader with temperature control (Biotek Synergy HTX plate reader with UV/VIS capabilities or similar) was used, and the temperature was increased from 25° C. to 37° C. The plate was incubated for 15 minutes, and the absorbance at 280 nm measured in each well at each step. This program continued until the instrument reached 50° C., where the samples were held for 60 minutes total, measuring the absorbance at 280 nm every 15 minutes. The formulation buffer reference absorbance at 280 nm (A280) value was subtracted from the sample measurement A280 value at each temperature and then they were

normalized to the measurement at 37° C. and plotted. These thermal stability plots were compared across different antibodies, mutants and peptide linkers to determine the anti-inflammatory constructs that would be the most resistant to thermal changes and therefore also more likely to be stable enough for a long-term intraocular therapy.

[0246] We then used the top performers from the thermal stability experiments in a long-term 37° C. stability studies. The MVPs were synthesized under sterile conditions and diluted to around 0.4 mg/mL in a sterile filtered human vitreous mimetic buffer (see Table 10) or remained in formulation buffer. The samples were either filtered using sterile 0.2 or 5 μ m spin filters before use or mixed with 0.010% sodium azide as an anti-microbial agent. Then, several 100-150 μ L aliquots of each sample were added to wells of a sterile 96 well plate with one day 0 aliquot reserved at 4° C. The remaining wells were filled with a sterile filtered human vitreous buffer+0.01% sodium azide to minimize evaporation. The plate was incubated in a standard tissue culture incubator at 37° C. with 5% CO₂. At discrete timepoints, one aliquot from each sample was removed from the plate under sterile conditions and analyzed. First, the UV-VIS spectrum of the sample was taken from 200-600 nm in 10 nm steps to monitor any dramatic changes in sample composition. Then, the protein concentration was measured to adjust for any differences in volume that may have occurred. The binding affinity to the appropriate ligand was measured using BLT methods described above using 5-10 nM of MVP as the top concentration. The change in K_{on} (association constant) over time was used to assess relative stability over time. To monitor changes in radius over time, the samples were spun for 5 minutes at 5000 g to remove any large aggregates or dust particles and the Rh was measured using DLS methods described above except that the instrument is at 37° C. and without any sample dilution.

TABLE 10

Vitreous Mimetic Buffer Composition	
Component	mg/mL
NaCl	7.14
KCl	0.38
CaCl ₂ 2H ₂ O	0.154
MgCl ₂ 6H ₂ O	0.2
dibasic NaPhosphate (NaH ₂ PO ₄)	0.42
NaHCO ₃	2.1
Dextrose	0.92
lactic acid	0.358
CuSO ₄	8.28 × 10 ⁻⁵
ZnSO ₄ heptahydrate	0.000561
FeCl ₂ tetrahydrate	0.000618
Transferrin (2 Fe binding sites)	0.0878
Reduced Glutathione (GSH)	0.0154

Example 7. Peptide-Polymer Conjugates Showed Intravitreal Retention

[0247] An extended intravitreal retention time of the conjugates was shown in a well-established pharmacokinetics model. New Zealand White rabbits (n=9) were divided into 3 groups randomized by weight. All animals received a 50- μ L ITV injections of hu_{anti}-TNF α -aH MVP in the left eye and the unconjugated VHH in the right eye using a 31 G insulin syringe. Both eyes received an equivalent molar dose of antibody. At 1 hour, 5 days and 10 or days post

injection, one group of three rabbits are sacrificed, and their eyes enucleated for analysis of intravitreal VHH. Both eyes were flash frozen, and the vitreous, retina, and aqueous humor were isolated from the frozen eye. Each tissue sample was then homogenized with a bead beater. After homogenization, the VHH concentrations were quantified either using ELISA or by digesting the peptide using trypsin and subjecting the samples to LC/mass spectrometry, or a similar method. Representative results for the extended intravitreal half-life in rabbit eyes after bioconjugation are shown in FIG. 5.

Example 8. MVP Efficacy in a Rat Model of Uveitis

[0248] The efficacy of mu_anti-TNF α _aH_CYS MVPs were validated to provide a treatment effect that can sufficiently reduce the symptoms of uveitis. A rat model of experimental autoimmune uveoretinitis (EAU) was used as a model of chronic posterior uveitis in humans. This model was induced by systemic immunization with the uveitogenic interphotoreceptor retinoid-binding protein (IRBP), and symptoms of uveitis appeared after 9-11 days. Rats were treated intravitreally with 12.5 μ g of mu_anti-TNF α _aH_CYS MVP (Conjugate #7). As a positive control, dexamethasone was used.

[0249] Male Lewis rats were divided into 4 groups (n=8) and randomized by weight. The groups received either mu_anti-TNF α _aH_CYS MVP (Conjugate #7) (using anti-mouse TNF α VHH) at 12.5 g, dexamethasone (40 μ g), or vehicle control. On Day 1, rats were immunized by a subcutaneous injection at the base of the tail and in each thigh with 30 μ g of bovine IRBP peptide R16 in 0.2 mL of Freund's adjuvant. On days 8 and 10, rat eyes were treated bilaterally with a 5 μ L ITV injection of either dose of their assigned treatment. Prior to the start of the study and on days 7, 9, 11, 14, we assessed ocular inflammation by slit lamp microscopy and assigned a clinical EAU score of 0-4 based on the appearance of inflammation. On day 14, animals were euthanized, one eye from each animal was processed for histopathology, and assigned a score of 0-4 based on the appearance of inflammation and cell infiltration. EAU and histopathology were scored based on published standardized scoring systems. The study results are summarized in FIG. 6.

Example 9. MVP Efficacy in a Second Rat Model of Uveitis

[0250] The efficacy of conjugate #10 was validated to provide a treatment effect that can sufficiently reduce the symptoms of uveitis. Male Lewis rats were divided into 4 groups (n=12 for EAU induced groups and n=8 for uninduced control) and randomized by weight. The groups received either conjugate #10 at 19 μ g, triamcinolone (40 μ g), or vehicle control. On Day 1, rats in the induced groups were immunized by a subcutaneous injection in each flank with 25 μ g of interphotoreceptor retinoid-binding protein (IRBP) peptide R16 in 0.1 mL of complete Freund's adjuvant for 50 μ g total. On days 4 and 8, rat eyes were treated bilaterally with a 5 μ L intravitreal (ITV) injection of conjugate #10 or vehicle control, or 1 μ L ITV injection of triamcinolone based on their assigned treatment. Prior to the start of the study and on days 3, 6, 10, 12 and 14, ocular inflammation was accessed by slit lamp microscopy and assigned a clinical EAU score of 0-4 based on the appearance of inflammation. FIGS. 7A-7B show the effect of conjugate #10 was comparable to triamcinolone in reducing ocular inflammation by slit lamp and as measured by inflammatory cytokine or inflammatory regulator levels.

[0251] On day 14, animals were euthanized and one eye from each animal was dissected into vitreous and aqueous humor to perform for cytokine analysis. After dissection the aqueous humor from each group was pooled and all tissues were flash frozen. A multiplex Rat Cytokine/Chemokine magnetic bead panel (Millipore Cat #RECYMAG65K27PMX) was used to assess the relative cytokine concentrations in the tissues according to the manufacturer's protocol. Briefly, the ocular tissues were thawed on ice and the sample volume was measured. Then, assay buffer was added to the samples to achieve a final volume of 55 μ L for duplicate readings and mixed well. Next, 25 μ L of samples, standards or controls was added to the assay plate, mixed with 25 μ L of beads and incubated at room temperature for 2 hours. The wells were washed on a magnetic plate washer and incubated with 25 μ L of detection antibodies for one hour and then 25 μ L of Streptavidin-Phycoerythrin for 30 minutes. Wells were washed on a magnetic plate washer and 125 μ L of Sheath Fluid Plus was added per well and then read on the Luminex. The amount of cytokine recovered from each sample was normalized to the volume of tissue recovered and plotted. A graphical analysis of key pro-inflammatory cytokines and inflammatory regulators is shown in FIG. 7B. Select cytokine concentrations are provided in Table 11. No statistical significance was observed in cytokine levels between triamcinolone and conjugate #10 treated rats.

TABLE 11

Cytokine concentration in the vitreous of rat eyes 14 days after uveitis induction				
Cytokine	Concentration (ng/mL) \pm SD			Statistical Significance [§]
	Vehicle Control	Triamcinolone	Conjugate #10	Vehicle Control vs. Conjugate #10
TNF α	0.169 \pm 0.097	0.046 \pm 0.051	0.058 \pm 0.038	*
IL-1 β	2.558 \pm 1.769	0.721 \pm 0.445	0.667 \pm 0.271	**
IL-1 α	5.532 \pm 3.527	1.321 \pm 0.844	1.511 \pm 0.709	**
IL-6	75.857 \pm 48.960	19.936 \pm 17.156	25.565 \pm 15.702	*
IFN γ	8.048 \pm 4.287	2.540 \pm 1.804	3.116 \pm 1.436	**
MCP-1	9.432 \pm 5.231	2.705 \pm 2.640	3.843 \pm 2.499	*

[§]Shown as the p-value of Tukey tests post hoc to ANOVA

Example 10. MVP Efficacy in a Rabbit
TNF- α -Induced Model of Uveitis

[0252] An anti-TNF α MVP was evaluated in the TNF- α -induced uveitis (EIU) model in rabbits, which involved an ITV injection of human TNF- α that elevated other inflammatory cytokines and induced ocular inflammation characteristic of non-infectious uveitis (NIU) in humans.

[0253] Male New Zealand white (NZW) rabbits were divided into 9 groups randomized by weight (n=3). On day zero, the groups received either the 0.26 mg of hu_anti-TNF α _aH MVP (conjugate #11) (4 groups) or vehicle control (4 groups) administered by bilateral 50- μ L ITV injections or no injection (1 group). One day after ITV drug delivery, ocular inflammation was induced by delivering 7.5, 5.0 or 2.5 μ g of human TNF α or PBS vehicle control by a unilateral 50- μ L ITV injection to the left eye. One uninduced group received no intravitreal injections. Prior to TNF α injection, and at 6, 24 and 48 hours after administering TNF α , inflammation severity was assessed by ocular examination and intraocular pressure was measured using a rebound tonometer. Clinical scores were assigned to each eye based on a published scale. The rabbits were euthanized 48 hours post TNF- α injection. Results are shown in FIGS. 8A-8B.

[0254] The left eyes can be dissected into the aqueous and vitreous humor and the vitreous humor can be processed for inflammatory cytokine analysis. Briefly, the dissected vitreous humor is thawed on ice and weighed. The vitreous is gently mixed in a homogenization buffer of PBS 0.05% v/v Tween-20, 1% w/v casein and 0.01% v/v Protease inhibitor cocktail set III (Sigma Catalog number 535140) at a concentration of 500 mg vitreous/mL. Then a solution of bovine testes hyaluronidase (MP Biochemicals catalog #37326-33-3) in PBS with 50 μ M MgCl₂ and 100 μ M CaCl₂ is added to the vitreous tissue in homogenization buffer at a final concentration of 0.04 mg hyaluronidase/g vitreous tissue. The tissue homogenization reactions are incubated at RT for 1 hour and then 4° C. overnight. Lastly, the homogenized vitreous tissue is spun at 5000 g for 5 minutes to pellet any debris and used for cytokine analysis.

[0255] A Milliplex Bovine Cytokine/Chemokine magnetic bead panel (Millipore Cat #BCYT1-33K-12) can be used to assess the relative cytokine concentrations in the tissues according to the manufacturer's protocol. Dissected aqueous humor is thawed and used as is, whereas vitreous humor is homogenized in hyaluronidase as described above. First, 25 μ L of samples, standards or controls is added to the assay plate, mixed with 25 μ L of beads and incubated at room temperature for 2 hours. The wells are washed on a magnetic plate washer and incubated with 25 μ L of detection antibodies for one hour and then 25 μ L of Streptavidin-Phycoerythrin for 30 minutes. Wells are washed on a magnetic plate washer and 125 μ L of Sheath Fluid Plus is added per well and then read on the Luminex. The amount of cytokine recovered from each sample is normalized to the volume of tissue recovered and plotted.

Example 11. MVP Efficacy in a Rabbit
Endotoxin-Induced Model of Uveitis

[0256] hu_anti-TNF α _aH MVP MVPs are evaluated in the endotoxin-induced uveitis (EIU) model in rabbits, which

involves an ITV injection of lipopolysaccharide (LPS) that elevates TNF α levels and induces ocular inflammation characteristic of NIU in humans.

[0257] NZW rabbits are divided into 4 groups randomized by weight (n=7, 3M/3F, one random). The groups receive either the hu_anti-TNF α _aH MVP, a positive control of either adalimumab or triamcinolone, or vehicle control administered by bilateral 50- μ L ITV injections. Two groups receive anti-TNF α MVP and one group receives adalimumab at an equivalent molar dose of antigen-binding epitope per eye: 225 μ g of total VHH antibody, 1 mg of adalimumab, or 1 mg triamcinolone.

[0258] Fifteen days after ITV drug delivery, EIU is induced with 10 g of LPS in 50- μ L ITV injections into the left eye of each animal except one of the anti-TNF α MVP groups (durability cohort). 60 days after ITV drug delivery, EIU is induced in the durability cohort using the same method. Prior to LPS injection, and at 6 and 24 hours after administering LPS, inflammation and EIU severity are assessed by ocular examination. EIU clinical scores will be assigned to each eye based on a published scale. The rabbits are euthanized 24 hours post LPS injection. LPS-induced eyes are processed for aqueous humor cell infiltration, inflammatory cytokine analysis, and histopathology to quantify cellular infiltrates. The uninduced right eyes are flash frozen and the anti-TNF α concentrations in the vitreous and aqueous humor are measured.

Example 12. MVP Durability in a Rabbit Model of
TNF α Uveitis

[0259] NZW rabbits are divided into 4 groups randomized by weight (n=7, 3M/3F, one random). The groups receive either the hu_anti-TNF α _aH MVP (0.25 mg), triamcinolone (1 mg), or vehicle control administered by bilateral 50- μ L ITV injections. Two groups receive anti-TNF α MVP and one group receives triamcinolone.

[0260] 1-30 days after ITV drug delivery, ocular inflammation is induced by delivering 7.5 μ g of human TNF α or PBS vehicle control by a unilateral 50- μ L ITV injection to the left eye. One uninduced group received no intravitreal injections. 60 days after ITV drug delivery, EIU is induced in the durability cohort using the same method. Prior to TNF α injection, and at 6, 24 and 48 hours after administering TNF α , inflammation severity is assessed by ocular examination. Intraocular pressure is also measured using a rebound tonometer on a daily basis. Clinical scores are assigned to each eye based on a published scale. The rabbits are euthanized 48 hours post TNF α injection.

[0261] The left eyes are dissected into the aqueous and vitreous humor, and the vitreous humor is processed for inflammatory cytokine analysis. Briefly, the dissected vitreous humor is thawed on ice and weighed. The vitreous is gently mixed in a homogenization buffer of PBS 0.05% v/v Tween-20, 1% w/v casein and 0.01% v/v Protease inhibitor cocktail set III (Sigma Catalog number 535140) at a concentration of 500 mg vitreous/mL. Then a solution of bovine testes hyaluronidase (MP Biochemicals catalog #37326-33-3) in PBS with 50 μ M MgCl₂ and 100 μ M CaCl₂ is added to the vitreous tissue in homogenization buffer at a final concentration of 0.04 mg hyaluronidase/g vitreous tissue. The tissue homogenization reactions are incubated at RT for 1 hour and then 4C overnight. Lastly, the homogenized vitreous tissue is spun at 5000 g for 5 minutes to pellet any debris and used for cytokine analysis.

[0262] A Milliplex Bovine Cytokine/Chemokine magnetic bead panel (Millipore Cat#BCYT1-33K-12) is used to assess the relative cytokine concentrations in the tissues according to the manufacturer's protocol. Dissected aqueous humor is thawed and used and vitreous humor is homogenized in hyaluronidase as described above. First, 25 μ L of samples, standards or controls is added to the assay plate, mixed with 25 μ L of beads and incubated at room tempera-

ture for 2 hours. The wells are washed on a magnetic plate washer and incubated with 25 μ L of detection antibodies for one hour and then 25 μ L of Streptavidin-Phycoerythrin for 30 minutes. Wells are washed on a magnetic plate washer and 125 μ L of Sheath Fluid Plus is added per well and then read on the Luminex. The amount of cytokine recovered from each sample is normalized to the volume of tissue recovered and plotted.

TABLE 12

Sequences	
Name	Sequence
blank	SEQ ID NO: 1-4
framework region 1	QVQLVESGGGLVQPGGSLRLSCAASG (SEQ ID NO: 5)
framework region 2	MGWFRQAPGKEREFVAAI (SEQ ID NO: 6)
framework region 3	YADSVKGRFTISRDN SKNTVY LQMNSLRPEDTAVYYCAA (SEQ ID NO: 7)
framework region 4	YWGQGT LVT VSS (SEQ ID NO: 8)
Nb42 CDR1	FAYSTYS (SEQ ID NO: 9)
Nb42 CDR2	NSGTFRLW (SEQ ID NO: 10)
Nb42 CDR3	RAWSPYSSTVDAGDFR (SEQ ID NO: 11)
blank	SEQ ID NO: 12-14
aTNFa-mu CDR1	GTFSSII (SEQ ID NO: 15)
aTNFa-mu CDR2	SWSGGTTV (SEQ ID NO: 16)
aTNFa-mu CDR3	RPYQKYNWASASYNV (SEQ ID NO: 17)
E1-1 CDR1	GGSDAGT (SEQ ID NO: 18)
E1-1 CDR2	SWAGTAWR (SEQ ID NO: 19)
E1-1 CDR3	LGSYEMDHH (SEQ ID NO: 20)
aH linker	AEAAAKEAAAKEAAKAGC (SEQ ID NO: 21)
AE3K2R(2) linker	AEEEKRKAEEEKRKAEEEAGC (SEQ ID NO: 22)
AE3K2R(3) linker	AEEEKRKAEEEKRKAEEEKRKAEEEAGC (SEQ ID NO: 23)
E4K4(2) linker	AEEEEKKKKKEEEEKKKAGC (SEQ ID NO: 24)
EA3K(2) linker	AEAAAKEAAKAGC (SEQ ID NO: 25)
Alfa linker	PSRLEELRRRLTEGC (SEQ ID NO: 26)
MyosinVI linker	AEEEEKKKQEEEAERLRRIQEEMEKERKREDEERRRKEEEERMK LEMEAKRKQEEERKKREDDEKRKKKAGC (SEQ ID NO: 27)
Spot linker	PDRVRAVSHWSSC (SEQ ID NO: 28)
GT9 linker	GTGTGTGTGTGTGTGTGC (SEQ ID NO: 29)

TABLE 12-continued

Sequences	
Name	Sequence
Modified Rigid linker	TPTTPTPTPTPGTPPGGC (SEQ ID NO: 30)
blank	SEQ ID NO: 31-60
Nb42	QVQLQESGGGLVQAGASLRSLSCAASGFAYSTYSMGWFRQVSGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 61)
HuNb42	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 62)
HuNb42 P14A	QVQLVESGGGLVQAGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 63)
HuNb42 T61A	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYADSVKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 64)
HuNb42 S75A	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 65)
HuNb42 L79V	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 66)
HuNb42 A88P	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 67)
HuNb42 L121Q	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 68)
HuNb42 T61A A88P	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYADSVKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 69)
HuNb42 A88P L115Q	QVQLVESGGGLVQPGGSLRSLSCAASGFAYSTYSMGWFRQAPGKERE GVATINSGTFRWLWYTDVSKGRFTISRDNKNTLYLQMNLSLRAEDTAVYY CAARAWSPYSSSTVDAGDFRYWGQGTQVTVSS (SEQ ID NO: 70)
aTNF- α mu	QVQLQDSGGGLVQAGGSLRSLSCAASGGTFSSIIIMAWFRQAPGKEREFV GAVSWGGTTVYADSVLGRFEISRDSARKSVLYLQMNLSLRAEDTAVYYC AARPYQKYNWASASYNVWGQGTQVTVSS (SEQ ID NO: 71)
aTNF- α mu 3MUT	QVQLQESGGGLVQAGGSLRSLSCAASGGTFSSIIIMAWFRQAPGKEREFV GAVSWGGTTVYADSVKGRFTISRDSARKSVLYLQMNLSLRAEDTAVYYC CAARPYQKYNWASASYNVWGQGTQVTVSS (SEQ ID NO: 72)
aTNF- α VHH	QVQLQESGGGLVQPGGSLRSLSCAASGRFSDHSGYTYTIGWFRQAPGK EREFVARIYWSSGNTYYADSVKGRFAISRDIKNTVDLTMMNLEPEDT AVYYCAARDGIPTSRVSVEYNVWGQGTQVTVSS (SEQ ID NO: 73)
blank	SEQ ID NO: 74-80
E1-1	EVQLQASGGGFVQPGGSLRSLSCAASGGGSDAGTMGWFRQAPGKEREF VSAISWAGTAWRYADSVKGRFTISRDNKNTLYLQMNLSLRAEDTAT YYCALGSYEMDHHYWGQGTQVTVSS (SEQ ID NO: 81)
E1-1 F11L	EVQLQASGGGLVQPGGSLRSLSCAASGGGSDAGTMGWFRQAPGKEREF VSAISWAGTAWRYADSVKGRFTISRDNKNTLYLQMNLSLRAEDTAT YYCALGSYEMDHHYWGQGTQVTVSS (SEQ ID NO: 82)
E1-1 S49A	EVQLQASGGGFVQPGGSLRSLSCAASGGGSDAGTMGWFRQAPGKEREF VAAISWAGTAWRYADSVKGRFTISRDNKNTLYLQMNLSLRAEDTAT YYCALGSYEMDHHYWGQGTQVTVSS (SEQ ID NO: 83)
E1-1 F11L	EVQLQASGGGLVQPGGSLRSLSCAASGGGSDAGTMGWFRQAPGKEREF

TABLE 12-continued

Sequences	
Name	Sequence
S49A	VAAISWAGTAWRYYADSVKGRFTISRDN SKNTVYLQMN SLRAEDTAT YYCALGSYEMDHHYWGQGTQVTVSS (SEQ ID NO: 84)
E1-1 CDR	EVQLQASGGGFVQPGGSLRLSCAASGRRFSIEAMGWFRQAPGKEREFV SAIDSGGSTDYADSVKGRFTISRDN SKNTVYLQMN SLRAEDTATYYCA VIGSSWYGRGLDYWGQGTQVTVSS (SEQ ID NO: 85)
blank	SEQ ID NO: 86-90
anti-VEGF VHH	DVQLVESGGGLVQPGGSLRLSCAASGRTFSSYSMGWFRQAPGKEREFV VAISKGGYKYDAVSLEGRFTISRDN AKNTVYLQIN SLRPEDTAVYYCAS SRAYGSSRLRLADTYEYWGQGT LVTVSS (SEQ ID NO: 91)
anti-VEGF DARPIN	GSDLDKKLLEAARAGQDDEVRI LMANGADV NARDSTGWTPLHLAAP WGHPEIVEVLLKNGADVNAADFQGWTPHLHLAAVGHLEIVEVLLKYG ADVNAQDKFGKTAFDISIDNGNEDLAEILQKAAGGSGGGGS (SEQ ID NO: 92)
anti-VEGF HuNb22 2MUT	QVQLVESGGGLVQPGGSLRLSCAASGYAYDTYYMGWFRQAPGKEREG VAGITSLVSGVAYYKYTYDSVKGRFTISRDN SKNTVDLQMN SLRAEDT AVYYCAASRSGLRARLLRPELYEYWGQGT LVTVSS (SEQ ID NO: 93)
anti-VEGF HuNb23 3MUT	QVQLVESGGGLVQPGGSLRLSCVASGDTYSSACMGWFRQAPGKEREG VATICSTSMRTRYADSVKGRFTISRDN SKNTVYLQMN SLRAEDTAV YYCATGHTVGSSWRDPGAWRYWGQGT LVTVSS (SEQ ID NO: 94)
anti-VEGF HuNb35 4MUT	QVQLVESGGGLVQPGGSLRLSCAASGLSYRPGYMGWFRQAPGKEREG VAIITGGVTHYADSVKGRFTISRDN SKNTVYLQMN SLRAEDTAVYYC ALANWVQFPLRVDGYKYWGQGT LVTVSS (SEQ ID NO: 95)
Hu_aVEGF_ VHH_3MUT	QVQLVESGGGLVQPGGSLRLSCAASGRTFSSYSMGWFRQAPGKEREFV AAISKGGYKYDAVSLEGRFTISRDN SKNTVYLQMN SLRPEDTAVYYCA SSRAYGSSRLRLADTYEYWGQGT LVTVSS (SEQ ID NO: 96)
Hu_aVEGF_ VHH_5MUT	QVQLVESGGGLVQPGGSLRLSCAASGRTFSSYSMGWFRQAPGKEREFV AAISKGGYKYDAVSVKGRFTISRDN SKNTVYLQMN SLRPEDTAVYYCA SSRAYGSSRLRLADTYEYWGQGT LVTVSS (SEQ ID NO: 97)
Hu_aVEGF_ VHH_6MUT	QVQLVESGGGLVQPGGSLRLSCAASGRTFSSYSMGWFRQAPGKEREFV AAISKGGYKYAVSVKGRFTISRDN SKNTVYLQMN SLRPEDTAVYYCA SSRAYGSSRLRLADTYEYWGQGT LVTVSS (SEQ ID NO: 98)
blank	SEQ ID NO: 99-100
anti-TNF α VHH (human) - rigid	QVQLQESGGGLVQPGGSLRLSCAASGRTFSDHSGYTYTIGWFRQAPGK EREFVARIYWSSGNTYYADSVKGRFAISRDI AKNTVDLT MNLEPEDT AVYYCAARDGIPTSRSVESYNYWGQGTQVTVSSPSTPPTPSPSTPPGGC DDDDK (SEQ ID NO: 101)
anti-TNF α VHH (human) - aH	QVQLQESGGGLVQPGGSLRLSCAASGRTFSDHSGYTYTIGWFRQAPGK EREFVARIYWSSGNTYYADSVKGRFAISRDI AKNTVDLT MNLEPEDT AVYYCAARDGIPTSRSVESYNYWGQGTQVTVSSAEAAAKEAAAKEAA AKAGC (SEQ ID NO: 102)
anti-TNF α VHH (mouse) - aH	QVQLQDSGGGLVQAGGSLRLSCAASGGTFSSII MAWFRQAPGKEREFV GAVSWSGGTTVYADSVLGRFEISRDSARKSVYLQMN SLKFPEDTAVYY CAARPYQKYNWASASYNVWGQGTQVTVSSAEAAAKEAAAKEAAK AGC (SEQ ID NO: 103)
anti-TNF α 3MUT VHH (mouse) - aH	QVQLQESGGGLVQAGGSLRLSCAASGGTFSSII MAWFRQAPGKEREFV GAVSWSGGTTVYADSVKGRFTISRDSARKSVYLQMN SLKFPEDTAVYY CAARPYQKYNWASASYNVWGQGTQVTVSSAEAAAKEAAAKEAAK AGC (SEQ ID NO: 104)
anti-TNF α affibody	CGGGVDNKNFKEVGWAFGEIGALPNLNLQFRAFIISLWDDPSQSANL LAEAKKLNDQAQPK (SEQ ID NO: 105)

TABLE 12-continued

Sequences	
Name	Sequence
anti-IL-1 β scFv-rigid	EIVMTQSPSTLSASVGDRIITCQASQSIDNWSWYQQKPGKAPKLLIYR ASTLASGVPSRFSGSGSGAEFTLTISLQPDDEFATYYCQNTGGGVSIAPG QGKTLTVLGGGGSGGGSGGGSGGGSEVQLVESGGGLVQPGGSL RLSCTASGFSLSAAMAWVRQAPGKLEWVGIIYDSASTYYASWAKG RFTISRDTSKNTVYLQMNSLR AEDTAVYYCARERAIFSGDFVLWGQGT LVTVSSSPSTPPTPSPSTPPGGC (SEQ ID NO: 106)
Hu_aTNFa Mu_3MUT	QVQLVESGGGLVQPGGSLRLSCAASGGTFSSII MAWFRQAPGKEREFVG AVSWSGGTTVYADSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYC AARPYQKYNWASASYNVWGQGT LVTVSS (SEQ ID NO: 107)
Hu_aTNFa Mu_5MUT	QVQLVESGGGLVQPGGSLRLSCAASGGTFSSII MAWFRQAPGKEREFVA AISWSGGTTVYADSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYCA ARPYQKYNWASASYNVWGQGT LVTVSS (SEQ ID NO: 108)
Hu_aTNFa Hu_7MUT	QVQLVESGGGLVQPGGSLRLSCAASGRTFSDHSGYTYTMGWFRQAPG KEREFVARIYWSSGNTYYADSVKGRFTISRDN SKNTVYLQMNSLRPED TAVYYCAARDGIPTSRSVESYNYWGQGT LVTVSS (SEQ ID NO: 109)
blank	SEQ ID NO: 110
Hu_aEGFR 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGRTSR SYGMGWFRQAPGKEREFV AGISWRGDS TGYADSVKGRFTISRDN SKNTVDLQMNSLRPEDTAVYYC AAAAGSAWYGTLYEYDYWGQGT LVTVSS (SEQ ID NO: 111)
Hu_aHer2_ 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGITFMRYAMGWYRQAPGKQREM VASINSGGTTNYADSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYC NARWVKPQFIDNNYWGQGT LVTVSS (SEQ ID NO: 112)
Hu_aPD1_ 102C3 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGSIFSIHAMGWFRQAPGKEREFVA AITWSGGITYYEDSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYCA ADRAESSWYDYWGQGT LVTVSS (SEQ ID NO: 113)
Hu_aPD1_ 102C12 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGSIASIHAMGWFRQAPGKEREFV AVITWSGGITYYADSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYC AGDKHQSSWYDYWGQGT LVTVSS (SEQ ID NO: 114)
Hu_aPD1_ 102E2 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGSISSIHAMGWFRQAPGKEREFVA AITWSGGITYYADSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYCA ADRAQSSWYDYWGQGT LVTVSS (SEQ ID NO: 115)
Hu_aPD1_ 102E8 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGSIFSIHAMGWFRQAPGKEREFVA LISWSGGSTYYEDSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYCA ADRDVSNWYDYWGQGT LVTVSS (SEQ ID NO: 116)
Hu_aPD1_ 102H12 4MUT	EVQLVESGGGLVQPGGSLRLSCAASGRAFS SGTMGWFRQAPGKEREFV ASIPWSGGRTYYADSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYC AVKERSTGWDFAWGQGT LVTVSS (SEQ ID NO: 117)
Hu aCaffeine 3MUT	EVQLVESGGGLVQPGGSLRLSCAASGRGTGTIYSMAWFRQAPGKEREFV ATIGWSSGITYYMDSVKGRFTISRDN SKNTVYLQMNSLRPEDTAVYYC AATRAYSVGYDYWGQGT LVTVSS (SEQ ID NO: 118)
blank	SEQ ID NO: 119-144
HuNb42 A88P aH_CYS	QVQLVESGGGLVQPGGSLRLSCAASGFAYSTYSMGWFRQAPGKEREA VATINSGTFR LWYTDSVKGRFTISRDN SKNTLYLQMNSLRPEDTAVYY CAARAWSPYSS TVDAGDFRYWGQGT LVTVSSA EAAAKEAAKEAAA KAGC (SEQ ID NO: 145)
HuNb42 T61A A88P aH_CYS	QVQLVESGGGLVQPGGSLRLSCAASGFAYSTYSMGWFRQAPGKEREA VATINSGTFR LWYTDSVKGRFTISRDN SKNTLYLQMNSLRPEDTAVYY CAARAWSPYSS TVDAGDFRYWGQGT LVTVSSA EAAAKEAAKEAAA KAGC (SEQ ID NO: 146)
HuNb42 A88P L115Q aH_CYS	QVQLVESGGGLVQPGGSLRLSCAASGFAYSTYSMGWFRQAPGKEREA VATINSGTFR LWYTDSVKGRFTISRDN SKNTLYLQMNSLRPEDTAVYY CAARAWSPYSS TVDAGDFRYWGQGT QVTVSSA EAAAKEAAKEAAA KAGC (SEQ ID NO: 147)

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SEQUENCE: 5		
QVQLVESGGG LVQPGGSLRL SCAASG		26
SEQ ID NO: 6	moltype = AA length = 18	
FEATURE	Location/Qualifiers	
source	1..18	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..18	
	note = framework region 2	
SEQUENCE: 6		
MGWFRQAPGK EREFVAAI		18
SEQ ID NO: 7	moltype = AA length = 39	
FEATURE	Location/Qualifiers	
source	1..39	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..39	
	note = framework region 3	
SEQUENCE: 7		
YADSVKGRFT ISRDNKNTV YLQMNSLRPE DTAVYYCAA		39
SEQ ID NO: 8	moltype = AA length = 12	
FEATURE	Location/Qualifiers	
source	1..12	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..12	
	note = framework region 4	
SEQUENCE: 8		
YWQQGTLVTV SS		12
SEQ ID NO: 9	moltype = AA length = 7	
FEATURE	Location/Qualifiers	
source	1..7	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..7	
	note = Nb42 CDR1	
SEQUENCE: 9		
FAYSTYS		7
SEQ ID NO: 10	moltype = AA length = 8	
FEATURE	Location/Qualifiers	
source	1..8	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..8	
	note = Nb42 CDR2	
SEQUENCE: 10		
NSGTFRLW		8
SEQ ID NO: 11	moltype = AA length = 16	
FEATURE	Location/Qualifiers	
source	1..16	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..16	
	note = Nb42 CDR3	
SEQUENCE: 11		
RAWSPYSSTV DAGDFR		16
SEQ ID NO: 12	moltype = length =	
SEQUENCE: 12		
000		
SEQ ID NO: 13	moltype = length =	
SEQUENCE: 13		
000		
SEQ ID NO: 14	moltype = length =	
SEQUENCE: 14		
000		
SEQ ID NO: 15	moltype = AA length = 7	

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FEATURE	Location/Qualifiers	
source	1..7	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..7	
	note = aTNFa-mu CDR1	
SEQUENCE: 15		
GTFSSII		7
SEQ ID NO: 16	moltype = AA length = 8	
FEATURE	Location/Qualifiers	
source	1..8	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..8	
	note = aTNFa-mu CDR2	
SEQUENCE: 16		
SWSGGTTV		8
SEQ ID NO: 17	moltype = AA length = 15	
FEATURE	Location/Qualifiers	
source	1..15	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..15	
	note = aTNFa-mu CDR3	
SEQUENCE: 17		
RPYQKYNWAS ASYNV		15
SEQ ID NO: 18	moltype = AA length = 7	
FEATURE	Location/Qualifiers	
source	1..7	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..7	
	note = E1-1 CDR1	
SEQUENCE: 18		
GGSDAGT		7
SEQ ID NO: 19	moltype = AA length = 8	
FEATURE	Location/Qualifiers	
source	1..8	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..8	
	note = E1-1 CDR2	
SEQUENCE: 19		
SWAGTAWR		8
SEQ ID NO: 20	moltype = AA length = 9	
FEATURE	Location/Qualifiers	
source	1..9	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..9	
	note = E1-1 CDR3	
SEQUENCE: 20		
LGSYEMDHH		9
SEQ ID NO: 21	moltype = AA length = 19	
FEATURE	Location/Qualifiers	
source	1..19	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..19	
	note = aH linker	
SEQUENCE: 21		
AEAAAKEAAA KEAAAKAGC		19
SEQ ID NO: 22	moltype = AA length = 21	
FEATURE	Location/Qualifiers	
source	1..21	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..21	
	note = AE3K2R(2) linker	

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SEQUENCE: 22		
AEEEEKKAEE EKRKAEEEAG C		21
SEQ ID NO: 23	moltype = AA length = 28	
FEATURE	Location/Qualifiers	
source	1..28	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..28	
	note = AE3K2R(3) linker	
SEQUENCE: 23		
AEEEEKKAEE EKRKAEEEKR KAEEEAGC		28
SEQ ID NO: 24	moltype = AA length = 20	
FEATURE	Location/Qualifiers	
source	1..20	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..20	
	note = E4K4(2) linker	
SEQUENCE: 24		
AEEEEKKKKE EEEKKKKAGC		20
SEQ ID NO: 25	moltype = AA length = 14	
FEATURE	Location/Qualifiers	
source	1..14	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..14	
	note = EA3K(2) linker	
SEQUENCE: 25		
AEAAAKEAAA KAGC		14
SEQ ID NO: 26	moltype = AA length = 16	
FEATURE	Location/Qualifiers	
source	1..16	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..16	
	note = Alfa linker	
SEQUENCE: 26		
PSRLEELRR RLTEGC		16
SEQ ID NO: 27	moltype = AA length = 77	
FEATURE	Location/Qualifiers	
source	1..77	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..77	
	note = MyosinVI linker	
SEQUENCE: 27		
AEEEEKKKQQ EEEAERLRR QEEEMEKRRK REEDEERRRK EEEERRMKLE MEAKRKQEEE		60
ERKKREDEK RKKKAGC		77
SEQ ID NO: 28	moltype = AA length = 13	
FEATURE	Location/Qualifiers	
source	1..13	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..13	
	note = Spot linker	
SEQUENCE: 28		
PDRVRAVSHW SSC		13
SEQ ID NO: 29	moltype = AA length = 20	
FEATURE	Location/Qualifiers	
source	1..20	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..20	
	note = GT9 linker	
SEQUENCE: 29		
GTGTGTGTGT GTGTGTGTGC		20
SEQ ID NO: 30	moltype = AA length = 17	
FEATURE	Location/Qualifiers	

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source	1..17	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..17	
	note = Modified Rigid linker	
SEQUENCE: 30		
TPPTPTPTPTP GTPPGGC		17
SEQ ID NO: 31	moltype =	length =
SEQUENCE: 31		
000		
SEQ ID NO: 32	moltype =	length =
SEQUENCE: 32		
000		
SEQ ID NO: 33	moltype =	length =
SEQUENCE: 33		
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SEQ ID NO: 34	moltype =	length =
SEQUENCE: 34		
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SEQ ID NO: 35	moltype =	length =
SEQUENCE: 35		
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SEQ ID NO: 36	moltype =	length =
SEQUENCE: 36		
000		
SEQ ID NO: 37	moltype =	length =
SEQUENCE: 37		
000		
SEQ ID NO: 38	moltype =	length =
SEQUENCE: 38		
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SEQ ID NO: 39	moltype =	length =
SEQUENCE: 39		
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SEQ ID NO: 40	moltype =	length =
SEQUENCE: 40		
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SEQ ID NO: 41	moltype =	length =
SEQUENCE: 41		
000		
SEQ ID NO: 42	moltype =	length =
SEQUENCE: 42		
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SEQ ID NO: 43	moltype =	length =
SEQUENCE: 43		
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SEQ ID NO: 44	moltype =	length =
SEQUENCE: 44		
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SEQ ID NO: 45	moltype =	length =
SEQUENCE: 45		
000		
SEQ ID NO: 46	moltype =	length =
SEQUENCE: 46		
000		
SEQ ID NO: 47	moltype =	length =
SEQUENCE: 47		
000		

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SEQ ID NO: 48	moltype =	length =	
SEQUENCE: 48			
000			
SEQ ID NO: 49	moltype =	length =	
SEQUENCE: 49			
000			
SEQ ID NO: 50	moltype =	length =	
SEQUENCE: 50			
000			
SEQ ID NO: 51	moltype =	length =	
SEQUENCE: 51			
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SEQ ID NO: 52	moltype =	length =	
SEQUENCE: 52			
000			
SEQ ID NO: 53	moltype =	length =	
SEQUENCE: 53			
000			
SEQ ID NO: 54	moltype =	length =	
SEQUENCE: 54			
000			
SEQ ID NO: 55	moltype =	length =	
SEQUENCE: 55			
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SEQ ID NO: 56	moltype =	length =	
SEQUENCE: 56			
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SEQ ID NO: 57	moltype =	length =	
SEQUENCE: 57			
000			
SEQ ID NO: 58	moltype =	length =	
SEQUENCE: 58			
000			
SEQ ID NO: 59	moltype =	length =	
SEQUENCE: 59			
000			
SEQ ID NO: 60	moltype =	length =	
SEQUENCE: 60			
000			
SEQ ID NO: 61	moltype = AA	length = 126	
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = Nb42		
SEQUENCE: 61			
QVQLQESGGG SLQAGASLRL SCAASGFAYS TYSMGWFRQV SGKEREQVAT INSGTFRLWY		60	
TDSVKGSFTI SRDNAKNMLY LQMNSLKPED TAIYYCAARA WSPYSSTVDA GDFRYWGQGT		120	
QVTVSS		126	
SEQ ID NO: 62	moltype = AA	length = 126	
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42		
SEQUENCE: 62			
QVQLVESGGG LVQPGGSLRL SCAASGFAYS TYSMGWFRQA PGKEREAVAT INSGTFRLWY		60	
TDSVKGRFTI SRDNSKNTLY LQMNSLRAED TAVYYCAARA WSPYSSTVDA GDFRYWGQGT		120	
LVTVSS		126	

SEQ ID NO: 63	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42 P14A		
SEQUENCE: 63			
QVQLVESGGG LVQAGGSLRL	SCAASGFAYS	TYSMGWFRQA	PGKEREA VAT INSGTFRLWY 60
TDSVKGRFTI SRDNSKNTLY	LQMNSLRAED	TAVYYCAARA	WSPYSSTVDA GDFRYWGQGT 120
LVTVSS			126
SEQ ID NO: 64	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42 T61A		
SEQUENCE: 64			
QVQLVESGGG LVQPGGSLRL	SCAASGFAYS	TYSMGWFRQA	PGKEREA VAT INSGTFRLWY 60
ADSVKGRFTI SRDNSKNTLY	LQMNSLRAED	TAVYYCAARA	WSPYSSTVDA GDFRYWGQGT 120
LVTVSS			126
SEQ ID NO: 65	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42 S75A		
SEQUENCE: 65			
QVQLVESGGG LVQPGGSLRL	SCAASGFAYS	TYSMGWFRQA	PGKEREA VAT INSGTFRLWY 60
TDSVKGRFTI SRD NAKNTLY	LQMNSLRAED	TAVYYCAARA	WSPYSSTVDA GDFRYWGQGT 120
LVTVSS			126
SEQ ID NO: 66	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42 L79V		
SEQUENCE: 66			
QVQLVESGGG LVQPGGSLRL	SCAASGFAYS	TYSMGWFRQA	PGKEREA VAT INSGTFRLWY 60
TDSVKGRFTI SRDNSKNTLY	LQMNSLRAED	TAVYYCAARA	WSPYSSTVDA GDFRYWGQGT 120
LVTVSS			126
SEQ ID NO: 67	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42 A88P		
SEQUENCE: 67			
QVQLVESGGG LVQPGGSLRL	SCAASGFAYS	TYSMGWFRQA	PGKEREA VAT INSGTFRLWY 60
TDSVKGRFTI SRDNSKNTLY	LQMNSLRPED	TAVYYCAARA	WSPYSSTVDA GDFRYWGQGT 120
LVTVSS			126
SEQ ID NO: 68	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..126		
	note = HuNb42 L121Q		
SEQUENCE: 68			
QVQLVESGGG LVQPGGSLRL	SCAASGFAYS	TYSMGWFRQA	PGKEREA VAT INSGTFRLWY 60
TDSVKGRFTI SRDNSKNTLY	LQMNSLRAED	TAVYYCAARA	WSPYSSTVDA GDFRYWGQGT 120
QVT VSS			126
SEQ ID NO: 69	moltype = AA length = 126		
FEATURE	Location/Qualifiers		
source	1..126		
	mol type = protein		

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REGION                organism = synthetic construct
                      1..126
                      note = HuNb42 T61A A88P

SEQUENCE: 69
QVQLVESGGG LVQPGGSLRL SCAASGFAYS TYSMGWFRQA PGKEREAVAT INSGTFRLWY 60
ADSVKGRFTI SRDNSKNTLY LQMNSLRPED TAVYYCAARA WSPYSSTVDA GDFRYWGQGT 120
LVTVSS                                           126

SEQ ID NO: 70        moltype = AA length = 126
FEATURE              Location/Qualifiers
source               1..126
                    mol_type = protein
                    organism = synthetic construct

REGION              1..126
                    note = HuNb42 A88P L115Q

SEQUENCE: 70
QVQLVESGGG LVQPGGSLRL SCAASGFAYS TYSMGWFRQA PGKEREAVAT INSGTFRLWY 60
TDSVKGRFTI SRDNSKNTLY LQMNSLRPED TAVYYCAARA WSPYSSTVDA GDFRYWGQGT 120
QVTVSS                                           126

SEQ ID NO: 71        moltype = AA length = 124
FEATURE              Location/Qualifiers
source               1..124
                    mol_type = protein
                    organism = synthetic construct

REGION              1..124
                    note = aTNF-a mu

SEQUENCE: 71
QVQLQDSGGG LVQAGGSLRL SCAASGGTFS SIIMAWFRQA PGKEREAVGA VWSGGGTVY 60
ADSVLGRFEI SRDSARKSVY LQMNSLKPED TAVYYCAARP YQKYNWASAS YNVWGQGTQV 120
TVSS                                           124

SEQ ID NO: 72        moltype = AA length = 124
FEATURE              Location/Qualifiers
source               1..124
                    mol_type = protein
                    organism = synthetic construct

REGION              1..124
                    note = aTNF-a mu 3MUT

SEQUENCE: 72
QVQLQESGGG LVQAGGSLRL SCAASGGTFS SIIMAWFRQA PGKEREAVGA VWSGGGTVY 60
ADSVKGRFTI SRDSARKSVY LQMNSLKPED TAVYYCAARP YQKYNWASAS YNVWGQGTQV 120
TVSS                                           124

SEQ ID NO: 73        moltype = AA length = 129
FEATURE              Location/Qualifiers
source               1..129
                    mol_type = protein
                    organism = synthetic construct

REGION              1..129
                    note = aTNF-a VHH

SEQUENCE: 73
QVQLQESGGG LVQPGGSLRL SCAASGRTFS DHSGYTYTIG WFRQAPGKER EFVARIYWSS 60
GNTYYADSVK GRFAISRDLA KNTVDLTMMN LEPEDTAVYY CAARDGIPTS RSVESYNYWG 120
QGTQVTVSS                                           129

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

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

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

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

SEQ ID NO: 81 moltype = AA length = 119
FEATURE Location/Qualifiers
source 1..119
 mol_type = protein
 organism = synthetic construct
REGION 1..119
 note = E1-1
SEQUENCE: 81
EVQLQASGGG FVQPGGSLRL SCAASGGGSD AGTMGWFRQA PGKEREFVSA ISWAGTAWRY 60
YADSVKGRFT ISRDNSKNTV YLQMNSLRAE DTATYYCALG SYEMDHHYWG QGTQVTVSS 119

SEQ ID NO: 82 moltype = AA length = 119
FEATURE Location/Qualifiers
source 1..119
 mol_type = protein
 organism = synthetic construct
REGION 1..119
 note = E1-1 F11L
SEQUENCE: 82
EVQLQASGGG FVQPGGSLRL SCAASGGGSD AGTMGWFRQA PGKEREFVSA ISWAGTAWRY 60
YADSVKGRFT ISRDNSKNTV YLQMNSLRAE DTATYYCALG SYEMDHHYWG QGTQVTVSS 119

SEQ ID NO: 83 moltype = AA length = 119
FEATURE Location/Qualifiers
source 1..119
 mol_type = protein
 organism = synthetic construct
REGION 1..119
 note = E1-1 S49A
SEQUENCE: 83
EVQLQASGGG FVQPGGSLRL SCAASGGGSD AGTMGWFRQA PGKEREFVAA ISWAGTAWRY 60
YADSVKGRFT ISRDNSKNTV YLQMNSLRAE DTATYYCALG SYEMDHHYWG QGTQVTVSS 119

SEQ ID NO: 84 moltype = AA length = 119
FEATURE Location/Qualifiers
source 1..119
 mol_type = protein
 organism = synthetic construct
REGION 1..119
 note = E1-1 F11L S49A
SEQUENCE: 84
EVQLQASGGG FVQPGGSLRL SCAASGGGSD AGTMGWFRQA PGKEREFVAA ISWAGTAWRY 60
YADSVKGRFT ISRDNSKNTV YLQMNSLRAE DTATYYCALG SYEMDHHYWG QGTQVTVSS 119

SEQ ID NO: 85 moltype = AA length = 120
FEATURE Location/Qualifiers
source 1..120
 mol_type = protein
 organism = synthetic construct
REGION 1..120
 note = E1-1 CDR
SEQUENCE: 85
EVQLQASGGG FVQPGGSLRL SCAASGRRFS IEAMGWFRQA PGKEREFVSA IDSGGSTDYA 60
DSVKGRFTIS RDNSKNTVYL QMNSLRAEDT ATYYCAVIGS SWYGRGLDYW GGTQVTVSS 120

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 = AA	length = 125			
FEATURE	Location/Qualifiers				
source	1..125				
	mol_type = protein				
	organism = synthetic construct				
REGION	1..125				
	note = anti-VEGF VHH				
SEQUENCE: 91					
DVQLVESGGG LVQPGGSLRL	SCAASGRFTS	SYSMGWFRQA	PGKEREFVVA	ISKGGYKYDA	60
VSLEGRFTIS	RDNAKNTVYL	QINSLRPEDT	AVYYCASSRA	YGSSRLRLAD	120
VTVSS					125
SEQ ID NO: 92	moltype = AA	length = 134			
FEATURE	Location/Qualifiers				
source	1..134				
	mol_type = protein				
	organism = synthetic construct				
REGION	1..134				
	note = anti-VEGF DARPIN				
SEQUENCE: 92					
GSDLDKKLLE AARAGQDDEV	RILMANGADV	NARDSTGWTP	LHLAAPWGHF	EIVEVLLKNG	60
ADVNAADFQG WTPLHLAAAV	GHLEIVEVLL	KYGADVNAQD	KFGKTAFDIS	IDNGNEDLAE	120
ILQKAAGGGS GGGG					134
SEQ ID NO: 93	moltype = AA	length = 130			
FEATURE	Location/Qualifiers				
source	1..130				
	mol_type = protein				
	organism = synthetic construct				
REGION	1..130				
	note = anti-VEGF HuNb22 2MUT				
SEQUENCE: 93					
QVQLVESGGG LVQPGGSLRL	SCAASGYAYD	TYIMGWFRQA	PGKEREGVAG	ITSLVSGVAY	60
YKYYTDSVKG RFTISRDNK	NTVDLQMNSL	RAEDTAVYYC	AASRSLRLAR	LLRPELYEYV	120
QGGLTVTVSS					130
SEQ ID NO: 94	moltype = AA	length = 127			
FEATURE	Location/Qualifiers				
source	1..127				
	mol_type = protein				
	organism = synthetic construct				
REGION	1..127				
	note = anti-VEGF HuNb23 3MUT				
SEQUENCE: 94					
QVQLVESGGG LVQPGGSLRL	SCVASGDTYS	SACMGWFRQA	PGKEREGVAT	ICTSTSMRTR	60
YYADSVKGRF TISRDNKNT	VYLQMNSLRA	EDTAVYYCAT	GHTVGSSSWRD	PGAWRYWGQG	120
TLTVTVSS					127
SEQ ID NO: 95	moltype = AA	length = 123			
FEATURE	Location/Qualifiers				
source	1..123				
	mol_type = protein				
	organism = synthetic construct				
REGION	1..123				
	note = anti-VEGF HuNb35 4MUT				
SEQUENCE: 95					
QVQLVESGGG LVQPGGSLRL	SCAASGLSYR	PGYMGWFRQA	PGKEREGVAI	ITTGGVTHYA	60
DSVKGRFTIS RDNSKNTVYL	QMNSLRAEDT	AVYYCALANW	VQPLPLRVDGY	KYWGQGTLVLT	120
VSS					123
SEQ ID NO: 96	moltype = AA	length = 125			
FEATURE	Location/Qualifiers				
source	1..125				
	mol_type = protein				
	organism = synthetic construct				
REGION	1..125				
	note = Hu_aVEGF_VHH_3MUT				
SEQUENCE: 96					
QVQLVESGGG LVQPGGSLRL	SCAASGRFTS	SYSMGWFRQA	PGKEREFVAA	ISKGGYKYDA	60

VSLEGRFTIS	RDNSKNTVYL	QMNSLRPEDT	AVYYCASSRA	YGSSRLRLAD	TYEYWGQGTL	120
VTVSS						125
SEQ ID NO: 97	moltype = AA length = 125					
FEATURE	Location/Qualifiers					
source	1..125					
	mol_type = protein					
	organism = synthetic construct					
REGION	1..125					
	note = Hu_aVEGF_VHH_5MUT					
SEQUENCE: 97						
QVQLVESGGG	LVQPGGSLRL	SCAASGRTFS	SYSMGWFRQA	PGKEREFVAA	ISKGGYKYDA	60
VSVKGRFTIS	RDNSKNTVYL	QMNSLRPEDT	AVYYCASSRA	YGSSRLRLAD	TYEYWGQGTL	120
VTVSS						125
SEQ ID NO: 98	moltype = AA length = 125					
FEATURE	Location/Qualifiers					
source	1..125					
	mol_type = protein					
	organism = synthetic construct					
REGION	1..125					
	note = Hu_aVEGF_VHH_6MUT					
SEQUENCE: 98						
QVQLVESGGG	LVQPGGSLRL	SCAASGRTFS	SYSMGWFRQA	PGKEREFVAA	ISKGGYKYAA	60
VSVKGRFTIS	RDNSKNTVYL	QMNSLRPEDT	AVYYCASSRA	YGSSRLRLAD	TYEYWGQGTL	120
VTVSS						125
SEQ ID NO: 99	moltype = length =					
SEQUENCE: 99						
000						
SEQ ID NO: 100	moltype = length =					
SEQUENCE: 100						
000						
SEQ ID NO: 101	moltype = AA length = 150					
FEATURE	Location/Qualifiers					
source	1..150					
	mol_type = protein					
	organism = synthetic construct					
REGION	1..150					
	note = anti-TNFalpha VHH (human) - rigid					
SEQUENCE: 101						
QVQLQESGGG	LVQPGGSLRL	SCAASGRTFS	DHSGYTYTIG	WFRQAPGKER	EFVARIYWSS	60
GNTYYADSVK	GRFAISRDLA	KNTVDLTMMN	LEPEDTAVYY	CAARDGIPTS	RSVESYNYWG	120
QGTQVTVSSP	STPTPTSPST	PPGGCDDDDK				150
SEQ ID NO: 102	moltype = AA length = 148					
FEATURE	Location/Qualifiers					
source	1..148					
	mol_type = protein					
	organism = synthetic construct					
REGION	1..148					
	note = anti-TNFalpha VHH (human)- aH					
SEQUENCE: 102						
QVQLQESGGG	LVQPGGSLRL	SCAASGRTFS	DHSGYTYTIG	WFRQAPGKER	EFVARIYWSS	60
GNTYYADSVK	GRFAISRDLA	KNTVDLTMMN	LEPEDTAVYY	CAARDGIPTS	RSVESYNYWG	120
QGTQVTVSSA	EAAAKEAAAK	EAAAKAGC				148
SEQ ID NO: 103	moltype = AA length = 144					
FEATURE	Location/Qualifiers					
source	1..144					
	mol_type = protein					
	organism = synthetic construct					
REGION	1..144					
	note = anti-TNFalpha VHH (mouse)- aH					
SEQUENCE: 103						
QVQLQDSGGG	LVQAGGSLRL	SCAASGGTFS	SIIMAWFRQA	PGKEREFVGA	VWSGGTTTVY	60
ADSVLGRFEI	SRDSARKSVY	LQMNSLKFPF	DTAVIYCAAR	PYQKYNWASA	SYNVWVGQGTQ	120
VTVSSAEAAA	KEAAAKEAAA	KAGC				144
SEQ ID NO: 104	moltype = AA length = 143					
FEATURE	Location/Qualifiers					
source	1..143					
	mol_type = protein					
	organism = synthetic construct					

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REGION 1..143
note = anti-TNFalpha 3MUT VHH (mouse)- aH

SEQUENCE: 104
QVQLQESGGG LVQAGGSLRL SCAASGGTFS SIIMAWFRQA PGKEREFGVA VWSGGGTTVY 60
ADSVKGRFTI SRDSARKSVY LQMNSLKPED TAVYYCAARP YQKYNWASAS YNVWGQGTQV 120
TVSSAEAAAK EAAAKEAAAK AGC 143

SEQ ID NO: 105 moltype = AA length = 62
FEATURE Location/Qualifiers
source 1..62
mol_type = protein
organism = synthetic construct

REGION 1..62
note = anti-TNFalpha affibody

SEQUENCE: 105
CGGGVDNKFN KEVGWAFGEI GALPNLNALQ FRAFIISLWD DPSQSANLLA EAKKLNDQA 60
PK 62

SEQ ID NO: 106 moltype = AA length = 265
FEATURE Location/Qualifiers
source 1..265
mol_type = protein
organism = synthetic construct

REGION 1..265
note = anti-IL-1beta scFv-rigid

SEQUENCE: 106
EIVMTQSPST LSASVGDRLV ITCQASQSID NWLSWYQQKP GKAPKLLIYR ASTLASGVPS 60
RFSGSGSGAE FTLTISSLQP DDFATYYCQN TGGGVSIAPG QGTKLTVLGG GGGSGGGGSG 120
GGGSGGGGSE VQLVESGGGL VQPGGSLRLS CTASGFSLSA AAMAWVRQAP GKGLEWVGII 180
YDSASTYYAS WAKGRFTISR DTSKNTVYLQ MNSLRAEDTA VYCARERA FSGDFVLWGQ 240
GTLTVTSSSP STPTPSPST PPGGC 265

SEQ ID NO: 107 moltype = AA length = 124
FEATURE Location/Qualifiers
source 1..124
mol_type = protein
organism = synthetic construct

REGION 1..124
note = Hu_aTNFaMu_3MUT

SEQUENCE: 107
QVQLVESGGG LVQPGGSLRL SCAASGGTFS SIIMAWFRQA PGKEREFGVA VWSGGGTTVY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAARP YQKYNWASAS YNVWGQGTLV 120
TVSS 124

SEQ ID NO: 108 moltype = AA length = 124
FEATURE Location/Qualifiers
source 1..124
mol_type = protein
organism = synthetic construct

REGION 1..124
note = Hu_aTNFaMu_5MUT

SEQUENCE: 108
QVQLVESGGG LVQPGGSLRL SCAASGGTFS SIIMAWFRQA PGKEREFGVA ISWSGGGTTVY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAARP YQKYNWASAS YNVWGQGTLV 120
TVSS 124

SEQ ID NO: 109 moltype = AA length = 129
FEATURE Location/Qualifiers
source 1..129
mol_type = protein
organism = synthetic construct

REGION 1..129
note = Hu_aTNFaHu_7MUT

SEQUENCE: 109
QVQLVESGGG LVQPGGSLRL SCAASGRTFS DHSGYTYTMG WFRQAPGKER EFVARIYWSS 60
GNTYYADSVK GRFTISRDN S KNTVYLMQNS LRPEDTAVYY CAARDGIPTS RSVESYNYWG 120
QGTLTVTSS 129

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

SEQ ID NO: 111 moltype = AA length = 124
FEATURE Location/Qualifiers
source 1..124
mol_type = protein

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REGION                organism = synthetic construct
                      1..124
                      note = Hu_aEGFR_3MUT

SEQUENCE: 111
EVQLVESGGG LVQPGGSLRL SCAASGRTSR SYGMGWFRQA PGKEREFVAG ISWRGDSTGY 60
ADSVKGRFTI SRDNSKNTVD LQMNSLRPED TAVYYCAAAA GSAWYGTLYE YDYWGQGLTV 120
TVSS                                         124

SEQ ID NO: 112        moltype = AA length = 120
FEATURE              Location/Qualifiers
source               1..120
                    mol_type = protein
                    organism = synthetic construct

REGION              1..120
                    note = Hu_aHer2_3MUT

SEQUENCE: 112
EVQLVESGGG LVQPGGSLRL SCAASGITFM RYAMGWYRQA PGKQREMVAS INSGGTTNYA 60
DSVKGRFTIS RDNSKNTVYL QMNSLRPEDT AVYYCNARWV KPQFIDNNYW GQGLVTVSS 120

SEQ ID NO: 113        moltype = AA length = 119
FEATURE              Location/Qualifiers
source               1..119
                    mol_type = protein
                    organism = synthetic construct

REGION              1..119
                    note = Hu_aPD1_102C3 3MUT

SEQUENCE: 113
EVQLVESGGG LVQPGGSLRL SCAASGSIFS IHAMGWFRQA PGKEREFVAA ITWSGGITYY 60
EDSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAADR AESSWYDYWG QGLVTVSS 119

SEQ ID NO: 114        moltype = AA length = 119
FEATURE              Location/Qualifiers
source               1..119
                    mol_type = protein
                    organism = synthetic construct

REGION              1..119
                    note = Hu_aPD1_102C12 3MUT

SEQUENCE: 114
EVQLVESGGG LVQPGGSLRL SCAASGSIAS IHAMGWFRQA PGKEREFVAV ITWSGGITYY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAGDK HQSSWYDYWG QGLVTVSS 119

SEQ ID NO: 115        moltype = AA length = 119
FEATURE              Location/Qualifiers
source               1..119
                    mol_type = protein
                    organism = synthetic construct

REGION              1..119
                    note = Hu_aPD1_102E2 3MUT

SEQUENCE: 115
EVQLVESGGG LVQPGGSLRL SCAASGSISS IHAMGWFRQA PGKEREFVAA ITWSGGITYY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAADR AQSSWYDYWG QGLVTVSS 119

SEQ ID NO: 116        moltype = AA length = 119
FEATURE              Location/Qualifiers
source               1..119
                    mol_type = protein
                    organism = synthetic construct

REGION              1..119
                    note = Hu_aPD1_102E8 3MUT

SEQUENCE: 116
EVQLVESGGG LVQPGGSLRL SCAASGSIFS INAMAWFRQA PGKEREFVAL ISWGGSTYY 60
EDSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAADR VDSNWDYWG QGLVTVSS 119

SEQ ID NO: 117        moltype = AA length = 119
FEATURE              Location/Qualifiers
source               1..119
                    mol_type = protein
                    organism = synthetic construct

REGION              1..119
                    note = Hu_aPD1_102H12 4MUT

SEQUENCE: 117
EVQLVESGGG LVQPGGSLRL SCAASGRAFS SGTMGWFRQA PGKEREFVAS IPWGGGRITYY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAVKE RSTGWDFAWG QGLVTVSS 119

SEQ ID NO: 118        moltype = AA length = 119
FEATURE              Location/Qualifiers

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source	1..119	
	mol_type = protein	
	organism = synthetic construct	
REGION	1..119	
	note = Hu aCaffeine 3MUT	
SEQUENCE: 118		
EVQLVESGGG LVQPGGSLRL SCAASGRTGT IYSMAWFRQA PGKEREFPLAT IGWSSGITYY	60	
MDSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAATR AYSVGVDYWG QGTLVTVSS	119	
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		
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SEQ ID NO: 125	moltype =	length =
SEQUENCE: 125		
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SEQ ID NO: 126	moltype =	length =
SEQUENCE: 126		
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SEQ ID NO: 127	moltype =	length =
SEQUENCE: 127		
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SEQ ID NO: 128	moltype =	length =
SEQUENCE: 128		
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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		
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SEQ ID NO: 132	moltype =	length =
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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		

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SEQ ID NO: 136	moltype =	length =	
SEQUENCE: 136			
000			
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 = AA	length = 145	
FEATURE	Location/Qualifiers		
source	1..145		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..145		
	note = HuNb42 A88P aH_CYS		
SEQUENCE: 145			
QVQLVESGGG LVQPGGSLRL SCAASGFAYS TYSMGWFRQA PGKEREAVAT INSGTFRLWY		60	
TDSVKGRFTI SRDNSKNTLY LQMNSLRPED TAVYYCAARA WSPYSSTVDA GDFRYWGQGT		120	
LVTVSSAEAA AKEAAAKEAA AKAGC		145	
SEQ ID NO: 146	moltype = AA	length = 145	
FEATURE	Location/Qualifiers		
source	1..145		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..145		
	note = HuNb42 T61A A88P aH_CYS		
SEQUENCE: 146			
QVQLVESGGG LVQPGGSLRL SCAASGFAYS TYSMGWFRQA PGKEREAVAT INSGTFRLWY		60	
ADSVKGRFTI SRDNSKNTLY LQMNSLRPED TAVYYCAARA WSPYSSTVDA GDFRYWGQGT		120	
LVTVSSAEAA AKEAAAKEAA AKAGC		145	
SEQ ID NO: 147	moltype = AA	length = 145	
FEATURE	Location/Qualifiers		
source	1..145		
	mol_type = protein		
	organism = synthetic construct		
REGION	1..145		
	note = HuNb42 A88P L115Q aH_CYS		
SEQUENCE: 147			
QVQLVESGGG LVQPGGSLRL SCAASGFAYS TYSMGWFRQA PGKEREAVAT INSGTFRLWY		60	
TDSVKGRFTI SRDNSKNTLY LQMNSLRPED TAVYYCAARA WSPYSSTVDA GDFRYWGQGT		120	
QVTVSSAEAA AKEAAAKEAA AKAGC		145	
SEQ ID NO: 148	moltype = AA	length = 65	
FEATURE	Location/Qualifiers		
source	1..65		

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                                mol_type = protein
                                organism = synthetic construct
REGION                          1..65
                                note = anti-TNFalpha affibody Gly
SEQUENCE: 148
CGGGVDNKFN KEVGWAFGEI GALPNLNALQ FRAFIISLWD DPSQSANLLA EAKKLNDQA 60
PKGKG                                           65

SEQ ID NO: 149                  moltype = AA length = 143
FEATURE                         Location/Qualifiers
source                          1..143
                                mol_type = protein
                                organism = synthetic construct
REGION                          1..143
                                note = Hu_aTNFaMu_3MUT aH_CYS
SEQUENCE: 149
QVQLVESGGG LVQPGGSLRL SCAASGGTFS SIIMAWFRQA PGKREFVGA VWSGGTTVY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAARP YQKYNWASAS YNVWGQGLV 120
TVSSAEAAAK EAAAKEAAAK AGC                                           143

SEQ ID NO: 150                  moltype = AA length = 143
FEATURE                         Location/Qualifiers
source                          1..143
                                mol_type = protein
                                organism = synthetic construct
REGION                          1..143
                                note = Hu_aTNFaMu_5MUT aH_CYS
SEQUENCE: 150
QVQLVESGGG LVQPGGSLRL SCAASGGTFS SIIMAWFRQA PGKREFVAA ISWSGGTTVY 60
ADSVKGRFTI SRDNSKNTVY LQMNSLRPED TAVYYCAARP YQKYNWASAS YNVWGQGLV 120
TVSSAEAAAK EAAAKEAAAK AGC                                           143

SEQ ID NO: 151                  moltype = AA length = 148
FEATURE                         Location/Qualifiers
source                          1..148
                                mol_type = protein
                                organism = synthetic construct
REGION                          1..148
                                note = Hu_aTNFaHu_7MUT aH_CYS
SEQUENCE: 151
QVQLVESGGG LVQPGGSLRL SCAASGRTFS DHSGYTYTMG WFRQAPGKER EFVARIYWSS 60
GNTYYADSVK GRFTISRDN S KNTVYLMQNS LRPEDTAVYY CAARDGIPTS RSVESYNYWG 120
QGTLLVTVSSA EAAAKEAAAK EAAAKAGC                                     148

SEQ ID NO: 152                  moltype = AA length = 152
FEATURE                         Location/Qualifiers
source                          1..152
                                mol_type = protein
                                organism = synthetic construct
REGION                          1..152
                                note = IL-2_C125S aH_CYS
SEQUENCE: 152
APTSSSTKKT QLQLEHLLLD LQMILNGINN YKNPKLTRML TFKFYMPKKA TELKHLQCLE 60
EELKPLEEVL NLAQSKNFHL RPRDLISNIN VIVLELKGSE TTFMCEYADE TATIVEFLNR 120
WITFSQSIIS TLTAEAAAKE AAAKEAAKA GC                                   152

SEQ ID NO: 153                  moltype = AA length = 133
FEATURE                         Location/Qualifiers
source                          1..133
                                mol_type = protein
                                organism = synthetic construct
REGION                          1..133
                                note = IL-15_5MUT aH_CYS
SEQUENCE: 153
NWNVISDLK KIEDLIQSMH IDATLYTESD VHPSCKVTAM QCFLSELQVI SLESGDASIH 60
DTVENLTILA NNSLSSNGYV TSGCKECEEE LEAKNIKEFL QSFVHVIVQMF INTSAEAAAK 120
EAAAKEAAAK AGC                                           133

SEQ ID NO: 154                  moltype = AA length = 166
FEATURE                         Location/Qualifiers
source                          1..166
                                mol_type = protein
                                organism = synthetic construct
REGION                          1..166
                                note = aTNFa_DARPin G3S_CYS
SEQUENCE: 154

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DLGKKLLEVA	RAGQDDEVRI	LMANGADVNA	ADHQSFTHPLH	LYAIFGHLEI	VEVLLKNGAD	60
VNASDWHGNT	PLHLAAWIGH	LEIVEVLLKY	GADVNTDHS	GSTPLHLAAT	LGHLEIVEVL	120
LKYGADVNAQ	DKFGKTAFDI	SIDNGNEDLA	EILQKAAGGG	SGGGSC		166

What is claimed is:

1. A method for treating uveitis in a subject in need thereof, comprising administering to the subject a therapeutically effective amount of a conjugate of Formula V:



wherein

each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

each Y is an organic linker;

Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and subscript n is an integer of from 1 to 1000.

2. The method of claim 1, wherein each X is independently an anti-TNF- α peptide or an anti-interleukin-1 β peptide.

3. The method of claim 1 or 2, wherein each X is a monoclonal IgG, an IgG fragment, single chain scFv, single-domain heavy-chain VHH, adnectin, affibody, anticalin, DARPin, or an engineered Kunitz-type inhibitor.

4. The method of any one of claims 1 to 3, wherein each X is a peptide having an amino acid sequence comprising any one of SEQ ID NOS: 61-73, 81-85, 91-98, 101-109, 111-118, and 145-154.

5. The method of any one of claims 1 to 3, wherein each X is a peptide having an amino acid sequence comprising:

(SEQ ID NO: 101)

QVQLQES GGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNNLEPEDTA
VYYCAARDGI PTERSSESYN YWGQGTQVTV SSPSTPPTPS PSTPPGGCDD
DDK,

(SEQ ID NO: 102)

QVQLQES GGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNNLEPEDTA
VYYCAARDGI PTERSSESYN YWGQGTQVTV SSAEAAAKEA AAKEAAKAG
C,

(SEQ ID NO: 103)

QVQLQDS GGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKREF
VGAVSWSGGT TVYADSVLGR FEISRDSARK SVYLMNSLK PETAIVYYCA
ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAAKEA AAKAGC,

(SEQ ID NO: 104)

QVQLQES GGLVQAGGS LRLSCAASGG TFSSIIMAWF RQAPGKREF
VGAVSWSGGT TVYADSVKGR FTISRDSARK SVYLMNSLK PETAIVYYCA
ARPYQKYNWA SASYNVWGQG TQVTVSSAEA AAKEAAAKEA AAKAGC,

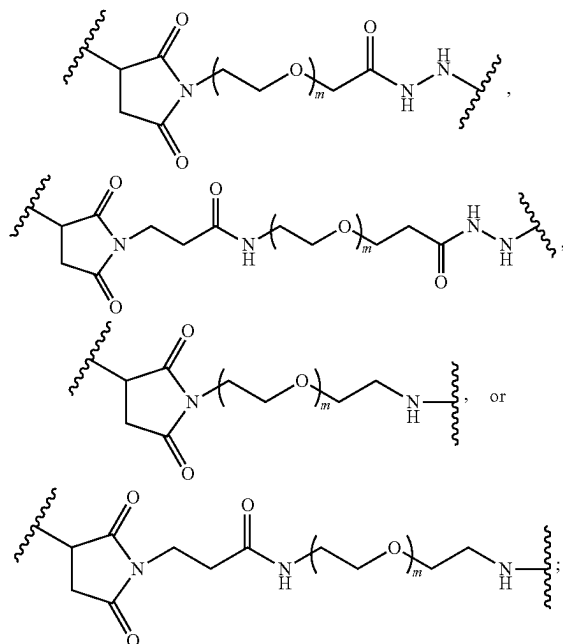
(SEQ ID NO: 105)

CGGGVDNKFN KEVGWAFGEI GALPNLNLQ FRAFIISLWD DPSQSANLLA
EAKKLNDQA PK,
or

(SEQ ID NO: 106)

EIVMTQS PSTLSASVGD RVIITCQASQ SIDNWLSWYQ QKPGKAPKLL
IYRASTLASG VPSRFSGSGS GAFTLTIS LQPDFFATYY CQNTGGGVSI
AFGQGTSLTV LGGGGSGSGG GSGGGSGSGG GSEVQLVESG GGLVQPGGSL
RLSCTASGFS LSSAAMAWVR QAPGKGLEWV GIIYDSASTY YASWAKGRFT
ISRDTSKNTV YLQMNSLRAE DTAVYYCARE RAIFSGDFVL WGQGTSLVTVS
SSPSTPPTPS PSTPPGGC.

6. The method of any one of claims 1 to 5, wherein each Y is an organic linker having the structure:



and

subscript m is an integer of from 1 to 300.

7. The method of any one of claims 1 to 5, wherein Z has a molecular weight of from about 0.4 MDa to about 2 MDa.

8. The method of any one of claims 1 to 7, wherein Z has a molecular weight of from about 0.7 MDa to about 1.5 MDa.

9. The method of any one of claims 1 to 8, wherein Z has a molecular weight of about 0.8 MDa.

10. The method of any one of claims 1 to 9, wherein conjugate of Formula V has the structure of Formula Va:

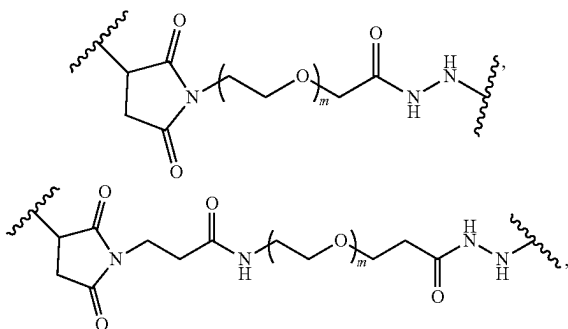


wherein

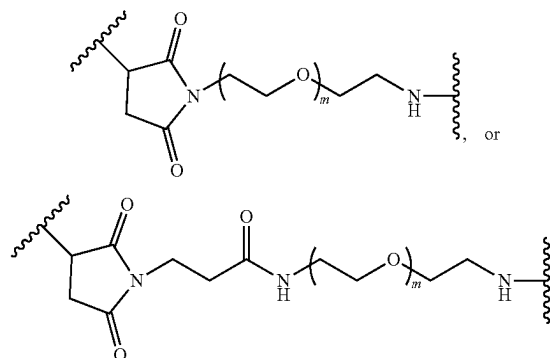
each X^1 is an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

each X^2 is a peptide linker that comprises an alpha-helix;

each Y is an organic linker having the structure:



-continued



Z is a hyaluronic acid polymer having a molecular weight of from about 0.1 MDa to about 3 MDa; and

subscript m is an integer of from 1 to 300.

11. The method of claim 10, wherein

each X^2 is a peptide linker having an amino acid sequence comprising:

(SEQ ID NO: 21)
AEAAAKEAAAKEAAKAGC,

(SEQ ID NO: 22)
AEEKKRKAEEKKRKAEEEGAGC,

(SEQ ID NO: 23)
AEEKKRKAEEKKRKAEEKKRKAEEEGAGC,

(SEQ ID NO: 24)
AEEEEKKKKKEEEKKKKKAGC,

(SEQ ID NO: 25)
AEAAAKEAAKAGC,

(SEQ ID NO: 26)
PSRLEELRRRLTEGC,
or

(SEQ ID NO: 27)
AEEEEKKKQEEEAERLRRIQEEMEKERKRREDEERRRKEEEER
RMKLEMEAKRKQEEERKKREDDEKRRKKAGC.

12. The method of any one of claims 1 to 11, wherein the conjugate of Formula V is a random polymer of Formula VI:



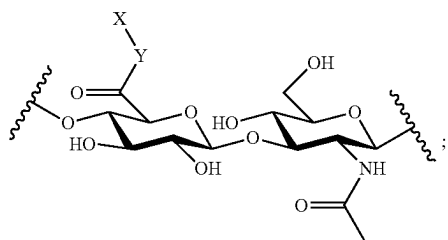
having a molecular weight of from about 0.1 MDa to about 3 MDa;

wherein

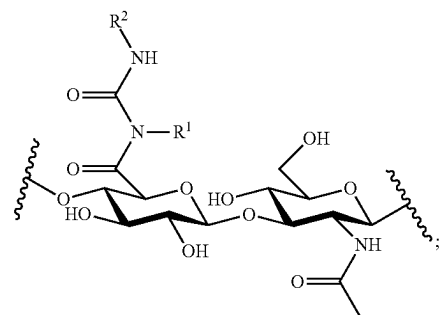
each X is independently an anti-inflammatory peptide having a molecular weight of from about 5 kDa to about 200 kDa;

each Y is an organic linker;

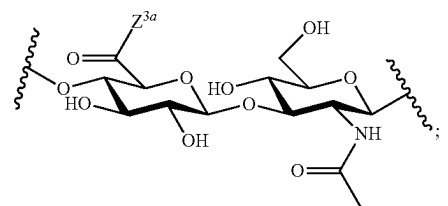
each $X-Y-Z^1$ moiety has the structure:



each Z^2 has the structure:



each Z^3 independently has the structure:



each R^1 and R^2 is independently C_1 - C_6 alkyl, $-(C_1$ - C_6 alkyl)- NR^3R^4 , or C_5 - C_8 cycloalkyl;

each R^3 and R^4 is independently H or C_1 - C_6 alkyl;

each Z^3a is independently OH or Y' ;

each Y' is an unreacted organic linker;

subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;

subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 100 to 10000.

13. The method of claim **12**, wherein each R^1 and R^2 is independently C_1 - C_3 alkyl or $-(C_1$ - C_3 alkyl)- NR^3R^4 .

14. The method of claim **12** or **13**, wherein each R^3 and R^4 is independently C_1 - C_3 alkyl.

15. The method of any one of claims **12** to **14**, wherein subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 100 to 10000.

16. The method of any one of claims **12** to **15**, wherein

subscript n is an integer of from 1 to 1000 and less than about 10% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 800 and less than about 8% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 100 to 10000.

17. The method of any one of claims **12** to **16**, wherein subscript n is an integer of from 10 to 450 and less than about 15% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 300 and less than about 10% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 1000 to 3000.

18. The method of any one of claims **12** to **17**, wherein subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 240 and less than about 8% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 1000 to 3000.

19. The method of any one of claims **12** to **18**, wherein subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 60 and less than about 2% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 1000 to 3000.

20. The method of any one of claims **12** to **19**, wherein subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 30 and less than about 1% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 1000 to 3000.

21. The method of any one of claims **12** to **20**, wherein subscript n is an integer of from 10 to 300 and less than about 10% of the sum of subscripts n, p, and q;

subscript p is an integer of from 1 to 15 and less than about 0.5% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 1000 to 3000.

22. The method of any one of claims **1** to **21**, wherein the uveitis is chronic uveitis.

23. The method of any one of claims **1** to **22**, wherein the uveitis is chronic non-infectious uveitis.

24. The method of any one of claims **1** to **23**, comprising intravitreal administration.

25. The method of any one of claims **1** to **24**, comprising multiple administrations of the conjugate.

26. The method of claim **25**, comprising administering the conjugate every month, every two months, or every three months.

27. The method of claim **25**, comprising administering the conjugate twice or three times yearly.

28. The method of claim **25**, comprising administering the conjugate yearly.

29. A conjugate that is a random polymer of Formula VI:



having a molecular weight of from about 0.1 MDa to about 3 MDa;

wherein
each X is independently an anti-TNF- α or anti-IL-1 β
peptide comprising:

(SEQ ID NO: 101)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
VYYCAARDGI PSTRSVESYN YWGQGTQVTV SSPSTPPTPS PSTPPGGCDD
DDK,

(SEQ ID NO: 102)

QVQLQES GGGLVQPGGS LRLSCAASGR TFSHSGYTY TIGWFRQAPG
KEREFVARIY WSSGNTYYAD SVKGRFAISR DIAKNTVDLT MNLEPEDTA
VYYCAARDGI PSTRSVESYN YWGQGTQVTV SSAEAAAKEA AAKEAAKAG
C,

(SEQ ID NO: 103)

QVQLQDS GGGLVQAGGS LRLSCAASGG TFSSIMAWF RQAPGKEREF
VGAVSWSGGT TVYADSVLGR FEISRDSARK SVYLMNSLK PEDTAVYYCA
ARPYQKYNWA SASYNVWGQG TQVTSSAEA AAKEAAAKEA AAKAGC,

(SEQ ID NO: 104)

QVQLQES GGGLVQAGGS LRLSCAASGG TFSSIMAWF RQAPGKEREF
VGAVSWSGGT TVYADSVKGR FTISRDSARK SVYLMNSLK PEDTAVYYCA
ARPYQKYNWA SASYNVWGQG TQVTSSAEA AAKEAAAKEA AAKAGC,

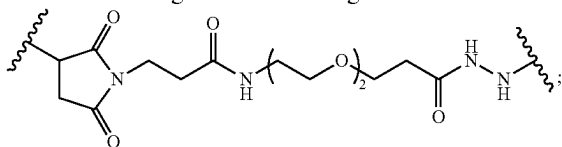
(SEQ ID NO: 105)

CGGGVDNKEN KEVGWAFGEI GALPNLNLQ FRAFIISLWD DPSQSANLLA
EAKKLNDQA PK,
or

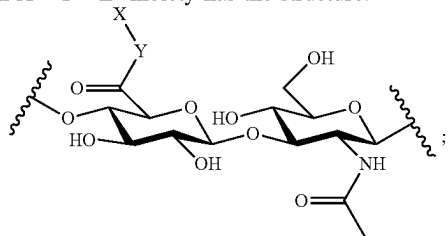
(SEQ ID NO: 106)

EIVMTQS PSTLSASVD RVIITCQASQ SIDNWLWYQ QKPGKAPKLL
IYRASTLASG VPSRFGSGS GAEFTLTIS LQPDDEFATY CQNTGGGVSI
AFGQGTCLTV LGGGGGSGG GSGGGGSGG GSEVQLVESG GGLVQPGGSL
RLSCTASGFS LSSAAMAVR QAPGKLEWV GIIYDSASTY YASWAKGRFT
ISRDTSKNTV YLMNSLRAE DTAVYYCARE RAIFSGDFVL WGQGTCLTVS
SSPSTPPTPS PSTPPGGC;

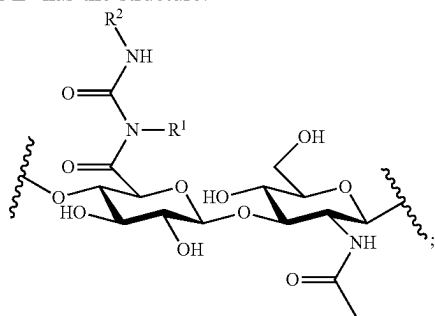
each Y is an organic linker having the structure:



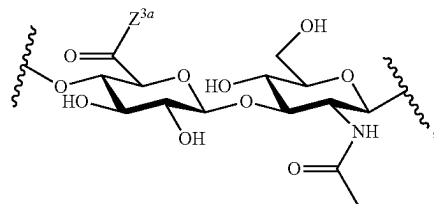
each X—Y—Z¹ moiety has the structure:



each Z² has the structure:



each Z³ independently has the structure:



each R¹ and R² is independently C₁-C₆ alkyl, —(C₁-C₆ alkyl)-NR³R⁴, or C₅-C₈ cycloalkyl;

each R³ and R⁴ is independently H or C₁-C₆ alkyl;

each Z^{3a} is independently OH or Y';

each Y' is an unreacted organic linker;

subscript n is an integer of from 1 to 1500 and less than about 15% of the sum of subscripts n, p, and q;

subscript p is an integer of from 0 to 1000 and less than about 10% of the sum of subscripts n, p, and q; and

subscript q is an integer of from 100 to 10000.

* * * * *