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# Compositions and methods for chimeric ligand receptor (CLR)-mediated conditional gene expression

#### **Abstract**

Disclosed are composition comprising (a) an inducible transgene construct, comprising a sequence encoding an inducible promoter and a sequence encoding a transgene, and (b) a receptor construct, comprising a sequence encoding a constitutive promoter and a sequence encoding an exogenous receptor, wherein, upon integration of the construct of (a) and the construct of (b) into a genomic sequence of a cell, the exogenous reporter is expressed, and wherein the exogenous reporter, upon binding a ligand, transduces an intracellular signal that targets the inducible promoter of (a) to modify gene expression. Methods for introducing compositions into cells and the use of the resultant cells in adoptive cell therapies are also provided.

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## **Field of Classification Search**

**USPC:** None

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## **Background/Summary**

CROSS-REFERENCE TO RELATED APPLICATIONS (1) This application is a U.S. National Phase Application, filed under 35 U.S.C. § 371 of International Patent Application No. PCT/2018/050288, filed Sep. 10, 2018, which claims the benefit of provisional application U.S. Ser. No. 62/556,310, filed Sep. 8, 2017. The contents of each of these applications are herein incorporated by reference in their entirety.

## INCORPORATION OF SEQUENCE LISTING

(1) The contents of the text file named "POTH-027-N01US\_SequenceListing\_R.txt" which was created on Jan. 11, 2021 and is 55,448 KB in size, are hereby incorporated by reference it their entirety.

#### FIELD OF THE DISCLOSURE

(2) The disclosure is directed to molecular biology, and more, specifically, to compositions and methods for use in a conditional gene expression system responsive to a chimeric ligand receptor (CLR)-mediated signal.

#### **BACKGROUND**

(3) There has been a long-felt but unmet need in the art for a method of controlling gene expression in genetically modified cells for the long-term delivery of therapeutic agents. The disclosure provides a solution by genetically modified cells that conditionally express genes upon activation of a cell-surface receptor.

#### **SUMMARY**

- (4) The disclosure provides a composition comprising (a) an inducible transgene construct, comprising a sequence encoding an inducible promoter and a sequence encoding a transgene, and (b) a receptor construct, comprising a sequence encoding a constitutive promoter and a sequence encoding an exogenous receptor, wherein, upon integration of the construct of (a) and the construct of (b) into a genomic sequence of a cell, the exogenous reporter is expressed, and wherein the exogenous reporter, upon binding a ligand, transduces an intracellular signal that targets the inducible promoter of (a) to modify gene expression. In certain embodiments, the composition modifies gene expression by increasing gene expression. In certain embodiments, the composition modifies gene expression by transiently modifying gene expression (e.g. for the duration of binding of the ligand to the exogenous receptor). In certain embodiments, the composition modifies gene expression acutely (e.g. the ligand reversibly binds to the exogenous receptor). In certain embodiments, the composition modifies gene expression chronically (e.g. the ligand irreversibly binds to the exogenous receptor).
- (5) In certain embodiments of the compositions of the disclosure, the cell may be a prokaryotic cell. Prokaryotic cells of the disclosure include, but are not limited to, bacteria and archaea. For example, bacteria of the disclosure include, but are not limited to, *Listeria monocytogenes*.
- (6) In certain embodiments of the compositions of the disclosure, the cell may be a eukaryotic cell. Eukaryotic cells of the disclosure include, but are not limited to, yeast, plants, algae, insects, mammals, amphibians, birds, reptiles, marsupials, rodents, and humans. Preferred eukaryotic cells of the disclosure include, but are not limited to, human cells. Exemplary human cells of the disclosure include but are not limited to, immune cells (e.g. T cells), myeloid cells and bone marrow cells (e.g. hematopoietic stem cells (HSCs)).
- (7) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises an endogenous receptor with respect to the genomic sequence of the cell. Exemplary receptors include, but are not limited to, intracellular receptors, cell-surface receptors, transmembrane receptors, ligand-gated ion channels, and G-protein coupled receptors.
- (8) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the non-naturally occurring receptor is a synthetic, modified, recombinant, mutant or chimeric receptor. In certain embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a T-cell receptor (TCR). In certain embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a scaffold protein. In certain embodiments, including those wherein the non-naturally occurring receptor does not comprise a transmembrane domain, the non-naturally occurring receptor interacts with a second transmembrane, membrane-bound and/or an intracellular receptor that, following contact with the non-naturally occurring receptor, transduces an intracellular signal.
- (9) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the non-naturally occurring receptor is a synthetic, modified, recombinant, mutant or chimeric receptor. In certain embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a T-cell receptor (TCR). In certain embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a scaffold protein. In certain embodiments, the non-naturally occurring receptor comprises a transmembrane domain. In certain embodiments, the non-naturally occurring receptor interacts with an intracellular receptor that transduces an intracellular signal. In certain embodiments, the non-naturally

occurring receptor comprises an intracellular signalling domain. In certain embodiments, the non-naturally occurring receptor is a chimeric ligand receptor (CLR). In certain embodiments, the CLR is a chimeric antigen receptor.

(10) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the CLR is a chimeric antigen receptor. In certain embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In certain embodiments, the ectodomain of (a) further comprises a signal peptide. In certain embodiments, the ectodomain of (a) further comprises a hinge between the ligand recognition region and the transmembrane domain. In certain embodiments, the signal peptide comprises a sequence encoding a human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\zeta$ , CD4, CD8 $\alpha$ , CD19, CD28, 4-1BB or GM-CSFR signal peptide. In certain embodiments, the signal peptide comprises a sequence encoding a human CD8a signal peptide. In certain embodiments, the signal peptide comprises an amino acid sequence comprising

MALPVTALLLPLALLLHAARP (SEQ ID NO:17000). In certain embodiments, the signal peptide is encoded by a nucleic acid sequence comprising

aggcactgccagtcaccgccctgctgctctgctgctgctgctgctgcagcagctagacca (SEQ ID NO:17001). In certain embodiments, the transmembrane domain comprises a sequence encoding a human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD4, CD8 $\epsilon$ , CD19, CD28, 4-1BB or GM-CSFR transmembrane domain. In certain embodiments, the transmembrane domain comprises a sequence encoding a human CD8 $\epsilon$  transmembrane domain. In certain embodiments, the transmembrane domain comprises an amino acid sequence comprising IYIWAPLAGTCGVLLLSLVITLYC (SEQ ID NO: 17002). In certain embodiments, the transmembrane domain is encoded by a nucleic acid sequence comprising

atctacatttgggcaccactggccgggacctgtggagtgctgctgctgagcctggtcatcacactgtactgc (SEQ ID NO: 17003). In certain embodiments, the endodomain comprises a human CD3 $\zeta$  endodomain. In certain embodiments, the at least one costimulatory domain comprises a human 4-1BB, CD28, CD3 $\zeta$ , CD40, ICOS, MyD88, OX-40 intracellular segment, or any combination thereof. In certain embodiments, the at least one costimulatory domain comprises a human CD3 $\zeta$  and/or a 4-1BB costimulatory domain. In certain embodiments, the CD3 $\zeta$  costimulatory domain comprises an amino acid sequence comprising

RVKFSRSADAPAYKQGQNQLYNELNLGRREEYDVLDKRRGRDPEMGGKPRRKNPQ EGLYNELQKDKMAEAYSEIGMKGERRRGKGHDGLYQGLSTATKDTYDALHMQALP PR (SEQ ID NO: 17004). In certain embodiments, the CD3 costimulatory domain is encoded by a nucleic acid sequence comprising

(11) TABLE-US-00001 (SEQ ID NO: 17008)

TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFACD.

ACCACAACCCCTGCCCCAGACCTCCCACACCCGCCCCTACCATCGCGAGTCAGC

CCCTGAGTCTGAGACCTGCAGGCCAGCCAGCTGCAGGAGGAGCTGTGCACA CCAGGGGCCTGGACTTCGCCTGCGAC (SEQ ID NO: 17009). In certain embodiments, the at least one protein scaffold specifically binds the ligand.

(12) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the CLR is a chimeric antigen receptor. In certain embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In certain embodiments, the at least one protein scaffold comprises an antibody, an antibody fragment, a single domain antibody, a single chain antibody, an antibody mimetic, or a Centyrin. In certain embodiments, the ligand recognition region comprises one or more of an antibody, an antibody fragment, a single domain antibody, a single chain antibody, an antibody mimetic, and a Centyrin. In certain embodiments, the single domain antibody comprises or consists of a VHH. In certain embodiments, the antibody mimetic comprises or consists of an affibody, an affilin, an affimer, an affitin, an alphabody, an anticalin, an avimer, a DARPin, a Fynomer, a Kunitz domain peptide or a monobody. In certain embodiments, the Centyrin comprises or consists of a consensus sequence of at least one fibronectin type III (FN3) domain. (13) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the CLR is a chimeric antigen receptor. In certain embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In certain embodiments, the Centyrin comprises or consists of a consensus sequence of at least one fibronectin type III (FN3) domain. In certain embodiments, the at least one fibronectin type III (FN3) domain is derived from a human protein. In certain embodiments, the human protein is Tenascin-C. In certain embodiments, the consensus sequence comprises

LPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAINLTVPGSERSYDL TGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT (SEQ ID NO: 17010). In certain embodiments, the consensus sequence comprises

MLPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAINLTVPGSERSYD LTGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT (SEQ ID NO: 17011). In certain embodiments, the consensus sequence is modified at one or more positions within (a) a A-B loop comprising or consisting of the amino acid residues TEDS at positions 13-16 of the consensus sequence; (b) a B-C loop comprising or consisting of the amino acid residues TAPDAAF at positions 22-28 of the consensus sequence; (c) a C-D loop comprising or consisting of the amino acid residues SEKVGE at positions 38-43 of the consensus sequence; (d) a D-E loop comprising or consisting of the amino acid residues GSER at positions 51-54 of the consensus sequence; (e) a E-F loop comprising or consisting of the amino acid residues GLKPG at positions 60-64 of the consensus sequence; (f) a F-G loop comprising or consisting of the amino acid residues KGGHRSN at positions 75-81 of the consensus sequence; or (g) any combination of (a)-(f). In certain embodiments, the Centyrin comprises a consensus sequence of at least 5 fibronectin type IT (FN3) domains. In certain embodiments, the Centyrin comprises a consensus sequence of at least 10 fibronectin type III (FN3) domains. In certain embodiments, the Centyrin comprises a consensus sequence of at least 15 fibronectin type III (FN3) domains. In certain embodiments, the scaffold binds an antigen with at least one affinity selected from a K.sub.D of less than or equal to 10.sup. -9 M, less than or equal to 10.sup. -10M, less than or equal to 10.sup. –1 M, less than or equal to 10.sup. –12 M, less than or equal to 10.sup. –13 M, less than or equal to 10.sup.-14 M, and less than or equal to 10.sup.-15 M. In certain embodiments, the K.sub.D is determined by surface plasmon resonance. In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the CLR is a chimeric antigen receptor. In certain embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least a VHH antibody; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In certain embodiments, the VHH is camelid. Alternatively, or in addition, in certain embodiments, the VHH is humanized. In certain embodiments, the sequence comprises two heavy chain variable regions of an antibody, wherein the complementaritydetermining regions (CDRs) of the VHH are human sequences.

- (14) In certain embodiments of the compositions of the disclosure, the sequence encoding the constitutive promoter of (b) comprises a sequence encoding an EF1 $\alpha$  promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the constitutive promoter of (b) comprises a sequence encoding a CMV promoter, a U6 promoter, a SV40 promoter, a PGK1 promoter, a Ubc promoter, a human beta actin promoter, a CAG promoter, or an EF1 $\alpha$  promoter.
- (15) In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding an NFkB promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding an interferon (IFN) promoter or a sequence encoding an interleukin-2 promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding a nuclear receptor subfamily 4 group A member 1 (NR4A1; also known as NUR77) promoter or a sequence encoding a NR4A1 promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding a T-cell surface glycoprotein CD5 (CD5) promoter or a sequence encoding a CD5 promoter. In certain embodiments, the interferon (IFN) promoter is an IFNy promoter. In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a cytokine or a chemokine. In certain embodiments, the cytokine or chemokine comprises IL2, IL3, IL4, IL5, IL6, IL10, IL12, IL13, IL17A/F, IL21, IL22, IL23, transforming growth factor beta (TGFβ), colony stimulating factor 2 (GM-CSF), interferon gamma (IFNy), Tumor necrosis factor (TNFα), LTα, perforin, Granzyme C (Gzmc), Granzyme B (Gzmb), C-C motif chemokine ligand 5 (CCL5), C-C motif chemokine ligand 4 (CCL4). C-C motif chemokine ligand 3 (CCL3), X-C motif chemokine ligand 1 (XCL1) and LIF interleukin 6 family cytokine (Lif).
- (16) In certain embodiments of the compositions of the disclosure, including those wherein the sequence encoding the inducible promoter of (a) comprises a sequence encoding a NR4A1 promoter or a sequence encoding a NR4A1 promoter, the NR4A1 promoter is activated by T-cell Receptor (TCR) stimulation in T cells and by B-cell Receptor (BCR) stimulation in B cells, therefore, inducing expression of any sequence under control of the NR4A1 promoter upon activation of a T-cell or B-cell of the disclosure through a TCR or BCR, respectively.
- (17) In certain embodiments of the compositions of the disclosure, including those wherein the sequence encoding the inducible promoter of (a) comprises a sequence encoding a CD5 promoter or a sequence encoding a CD5 promoter, the CD5 promoter is activated by T-cell Receptor (TCR) stimulation in T cells, therefore, inducing expression of any sequence under control of the CD5 promoter upon activation of a T-cell of the disclosure through a TCR.
- (18) In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a gene comprising a surface protein involved in cell differention, activation, exhaustion and function. In certain embodiments, the gene comprises CD69, CD71, CTLA4, PD-1, TIGIT, LAG3, TIM-3, GITR, MHCII, COX-2, FASL and 4-1BB.
- (19) In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a gene involved in CD metabolism and differentiation. In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of Nr4a1, Nr4a3, Tnfrsf9 (4-1BB), Sema7a, Zfp3612, Gadd45b, Dusp5, Dusp6 and Neto2. (20) In certain embodiments of the compositions of the disclosure, the transgene comprises a sequence that is endogenous with respect to the genomic sequence of the cell.
- (21) In certain embodiments of the compositions of the disclosure, the transgene comprises a sequence that is exogenous with respect to the genomic sequence of the cell. In certain embodiments, the exogenous sequence is a sequence variant of an endogenous sequence within the genome of the cell. In certain embodiments, the exogenous sequence is a wild type sequence of gene that is entirely or partially absent in the cell, and wherein the gene is entirely present in the genome of a healthy cell. In certain embodiments, the exogenous sequence is a synthetic, modified, recombinant, chimeric or non-naturally occurring sequence with respect to the genome of the cell. In certain embodiments, the transgene encodes a secreted protein. In certain embodiments, the secreted protein is produced and/or secreted from the cell at a level that is therapeutically effective to treat a disease or disorder in a subject in need thereof.

- (22) In certain embodiments of the compositions of the disclosure, a first transposon comprises the inducible transgene construct of (a) and a second transposon comprises the receptor construct of (b). In certain embodiments of the compositions of the disclosure, a first vector comprises the first transposon and a second vector comprises the second transposon. In certain embodiments of the compositions of the disclosure, a vector comprises the first transposon and the second transposon. In certain embodiments, the first transposon and the second transposon are oriented in the same direction. In certain embodiments, the first transposon and the second transposon are oriented in opposite directions. In certain embodiments, the vector is a plasmid. In certain embodiments, the vector is a nanoplasmid.
- (23) In certain embodiments of the compositions of the disclosure, the vector is a viral vector. Viral vectors of the disclosure may comprise a sequence isolated or derived from a retrovirus, a lentivirus, an adenovirus, an adeno-associated virus or any combination thereof. The viral vector may comprise a sequence isolated or derived from an adeno-associated virus (AAV). The viral vector may comprise a recombinant AAV (rAAV). Exemplary adeno-associated viruses and recombinant adeno-associated viruses of the disclosure comprise two or more inverted terminal repeat (ITR) sequences located in cis next to a sequence encoding a construct of the disclosure. Exemplary adeno-associated viruses and recombinant adeno-associated viruses of the disclosure include, but are not limited to all serotypes (e.g. AAV1, AAV2, AAV3, AAV4. AAV5, AAV6, AAV7, AAV8, and AAV9). Exemplary adeno-associated viruses and recombinant adeno-associated viruses of the disclosure include, but are not limited to, self-complementary AAV (scAAV) and AAV hybrids containing the genome of one serotype and the capsid of another serotype (e.g. AAV2/5, AAV-DJ and AAV-DJ8). Exemplary adeno-associated viruses and recombinant adeno-associated viruses of the disclosure include, but are not limited to, rAAV-LK03 and AAVs with the NP-59 and NP-84 capsid variants.
- (24) In certain embodiments of the compositions of the disclosure, the vector is a nanoparticle. Exemplary nanoparticle vectors of the disclosure include, but are not limited to, nucleic acids (e.g. RNA, DNA, synthetic nucleotides, modified nucleotides or any combination thereof), amino acids (L-amino acids, D-amino acids, synthetic amino acids, modified amino acids, or any combination thereof), polymers (e.g. polymersomes), micelles, lipids (e.g. liposomes), organic molecules (e.g. carbon atoms, sheets, fibers, tubes), inorganic molecules (e.g. calcium phosphate or gold) or any combination thereof. A nanoparticle vector may be passively or actively transported across a cell membrane.
- (25) In certain embodiments of the compositions of the disclosure, first transposon or the second transposon is a piggyBac transposon. In certain embodiments, the first transposon and the second transposon is a piggyBac transposon. In certain embodiments, the composition further comprises a plasmid or a nanoplasmid comprising a sequence encoding a transposase enzyme. In certain embodiments, the sequence encoding a transposase enzyme is an mRNA sequence. In certain embodiments, the transposase is a piggyBac transposase. In certain embodiments, the piggyBac transposase comprises an amino acid sequence comprising SEQ ID NO: 1. In certain embodiments, the piggyBac transposase is a hyperactive variant and wherein the hyperactive variant comprises an amino acid substitution at one or more of positions 30, 165, 282 and 538 of SEQ ID NO: 1. In certain embodiments, the amino acid substitution at position 30 of SEQ ID NO: 1 is a substitution of a valine (V) for an isoleucine (I) (130V). In certain embodiments, the amino acid substitution at position 165 of SEQ ID NO: 1 is a substitution of a serine (S) for a glycine (G) (G165S). In certain embodiments, the amino acid substitution at position 282 of SEQ ID NO: 1 is a substitution of a valine (V) for a methionine (M) (M282V). In certain embodiments, the amino acid substitution at position 538 of SEQ ID NO: 1 is a substitution of a lysine (K) for an asparagine (N) (N538K). In certain embodiments, the transposase is a Super piggyBac (SPB) transposase. In certain embodiments, the Super piggyBac (SPB) transposase comprises an amino acid sequence comprising SEQ ID NO: 2.
- (26) In certain embodiments of the disclosure, the transposase enzyme is a piggyBac<sup>™</sup> (PB) transposase enzyme. The piggyBac (PB) transposase enzyme may comprise or consist of an amino acid sequence at least 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (27) TABLE-US-00002 (SEQ ID NO: 17029) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEI SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVI EQPGSSLASN RILTLPQRTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121 PTRMCRNIYD PLDCFKLFFT DEIISEIVKW TNAEISLKRR ESMTGATFRD TNEDEIYAFF

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181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS
IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PQAHLTIDEQ LLGFRQRQPF
RMYIPNKPSK YQIKILMMCD 301 SGYKYMINGM PYLGRGTQTN GVRLGEYYVK
ELSKPVHGSC RNITCDNWFT SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE
VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST
GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWRMALL YGMINIACIN 481
SFIIYSHNVS SKGEKVQSRK KFMRNLYMSL TSSFMRKRLE APTLKRYLRD NISNILPNEV
541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC QSCF.
(28) In certain embodiments of the disclosure, the transposase enzyme is a piggyBac™ (PB) transposase
enzyme that comprises or consists of an amino acid sequence having an amino acid substitution at one or
more of positions 30, 165, 282, or 538 of the sequence:
(29) TABLE-US-00003 (SEQ ID NO: 17029) 1 MGSSLDDEHI LSALLQSDDE
LVGEDSDSEI SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVI
EQPGSSLASN RILTLPQRTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121
PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTGATFRD TNEDEIYAFF
181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS
IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGFRGRCPF
RMYIPNKPSK YGIKILMMCD 301 SGYKYMINGM PYLGRGTQTN GVPLGEYYVK
ELSKPVHGSC RNITCDNWFT SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE
VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST
GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481
SFIIYSHNVS SKGEKVQSRK KFMRNLYMSL TSSFMRKRLE APTLKRYLRD NISNILPNEV
541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC QSCF.
(30) In certain embodiments, the transposase enzyme is a piggyBac<sup>TM</sup> (PB) transposase enzyme that
comprises or consists of an amino acid sequence having an amino acid substitution at two or more of
positions 30, 165, 282, or 538 of the sequence of SEQ ID NO: 1. In certain embodiments, the transposase
enzyme is a piggyBac<sup>TM</sup> (PB) transposase enzyme that comprises or consists of an amino acid sequence
having an amino acid substitution at three or more of positions 30, 165, 282, or 538 of the sequence of
SEQ ID NO: 1. In certain embodiments, the transposase enzyme is a piggyBac™ (PB) transposase
enzyme that comprises or consists of an amino acid sequence having an amino acid substitution at each of
the following positions 30, 165, 282, and 538 of the sequence of SEQ ID NO: 1. In certain embodiments,
the amino acid substitution at position 30 of the sequence of SEQ ID NO: 1 is a substitution of a valine
(V) for an isoleucine (1). In certain embodiments, the amino acid substitution at position 165 of the
sequence of SEQ ID NO: 1 is a substitution of a serine (S) for a glycine (G). In certain embodiments, the
amino acid substitution at position 282 of the sequence of SEQ ID NO: 1 is a substitution of a valine (V)
for a methionine (M). In certain embodiments, the amino acid substitution at position 538 of the sequence
of SEQ ID NO: 1 is a substitution of a lysine (K) for an asparagine (N).
(31) In certain embodiments of the disclosure, the transposase enzyme is a Super piggyBac™ (SPB)
transposase enzyme. In certain embodiments, the Super piggyBac™ (SPB) transposase enzymes of the
disclosure may comprise or consist of the amino acid sequence of the sequence of SEQ ID NO: 1 wherein
the amino acid substitution at position 30 is a substitution of a valine (V) for an isoleucine (I), the amino
acid substitution at position 165 is a substitution of a serine (S) for a glycine (G), the amino acid
substitution at position 282 is a substitution of a valine (V) for a methionine (M), and the amino acid
substitution at position 538 is a substitution of a lysine (K) for an asparagine (N). In certain embodiments,
the Super piggyBac™ (SPB) transposase enzyme may comprise or consist of an amino acid sequence at
least 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
(32) TABLE-US-00004 (SEQ ID NO: 17030) 1 MGSSLDDEHI LSALLQSDDE
LVGEDSDSEV SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVI
EQPGSSLASN RILTLPQRTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121
PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTSATFRD TNEDEIYAFF
181 GILVMTAVRK DNHMSTDDLF DPSLSMVYVS VMSRDRFDFL IRCLRMDDKS
IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGFRGRCPF
RVYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTQTN GVPLGEYYVK
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ELSKPVHGSC RNITCDNWFT SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE
VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST
GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481
                SKGEKVQSRK KFMRNLYMSL TSSFMRKRLE APTLKRYLRD NISNILPKEV
SFIIYSHNVS
541 PGTSDDSTEE PVMKKRTYCT YCRSKIRRKA NASCKKCKKV ICREHNIDMC QSCF.
(33) In certain embodiments of the disclosure, including those embodiments wherein the transposase
comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> or Super
piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more of
positions 3, 46, 82, 103, 119, 125, 177, 180, 185, 187, 200, 207, 209, 226, 235, 240, 241, 243, 258, 296,
298, 311, 315, 319, 327, 328, 340, 421, 436, 456, 470, 486, 503, 552, 570 and 591 of the sequence of SEQ
ID NO: 1 or SEQ ID NO: 2. In certain embodiments, including those embodiments wherein the
transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac™
or Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more
of positions 46, 119, 125, 177, 180, 185, 187, 200, 207, 209, 226, 235, 240, 241, 243, 296, 298, 311, 315,
319, 327, 328, 340, 421, 436, 456, 470, 485, 503, 552 and 570. In certain embodiments, the amino acid
substitution at position 3 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of an asparagine (N) for a
serine (S). In certain embodiments, the amino acid substitution at position 46 of SEQ ID NO: 1 or SEQ ID
NO: 2 is a substitution of a serine (S) for an alanine (A). In certain embodiments, the amino acid
substitution at position 46 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a threonine (T) for an
alanine (A). In certain embodiments, the amino acid substitution at position 82 of SEO ID NO: 1 or SEO
ID NO: 2 is a substitution of a tryptophan (W) for an isoleucine (I). In certain embodiments, the amino
acid substitution at position 103 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a proline (P) for a
serine (S). In certain embodiments, the amino acid substitution at position 119 of SEQ ID NO: 1 or SEQ
ID NO: 2 is a substitution of a proline (P) for an arginine (R). In certain embodiments, the amino acid
substitution at position 125 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of an alanine (A) a
cysteine (C). In certain embodiments, the amino acid substitution at position 125 of SEQ ID NO: 1 or SEQ
ID NO: 2 is a substitution of a leucine (L) for a cysteine (C). In certain embodiments, the amino acid
substitution at position 177 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a lysine (K) for a
tyrosine (Y). In certain embodiments, the amino acid substitution at position 177 of SEQ ID NO: 1 or SEQ
ID NO: 2 is a substitution of a histidine (H) for a tyrosine (Y). In certain embodiments, the amino acid
substitution at position 180 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a leucine (L) for a
phenylalanine (F). In certain embodiments, the amino acid substitution at position 180 of SEQ ID NO: 1
or SEQ ID NO: 2 is a substitution of an isoleucine (I) for a phenylalanine (F). In certain embodiments, the
amino acid substitution at position 180 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a valine (V)
for a phenylalanine (F). In certain embodiments, the amino acid substitution at position 185 of SEQ ID
NO: 1 or SEQ ID NO: 2 is a substitution of a leucine (L) for a methionine (M). In certain embodiments,
the amino acid substitution at position 187 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a
glycine (G) for an alanine (A). In certain embodiments, the amino acid substitution at position 200 of SEQ
ID NO: 1 or SEQ ID NO: 2 is a substitution of a tryptophan (W) for a phenylalanine (F). In certain
embodiments, the amino acid substitution at position 207 of SEQ ID NO: 1 or SEQ ID NO: 2 is a
substitution of a proline (P) for a valine (V). In certain embodiments, the amino acid substitution at
position 209 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a phenylalanine (F) for a valine (V).
In certain embodiments, the amino acid substitution at position 226 of SEQ ID NO: 1 or SEQ ID NO: 2 is
a substitution of a phenylalanine (F) for a methionine (M). In certain embodiments, the amino acid
substitution at position 235 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of an arginine (R) for a
leucine (L). In certain embodiments, the amino acid substitution at position 240 of SEQ ID NO: 1 or SEQ
ID NO: 1 is a substitution of a lysine (K) for a valine (V). In certain embodiments, the amino acid
substitution at position 241 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a leucine (L) for a
phenylalanine (F). In certain embodiments, the amino acid substitution at position 243 of SEQ ID NO: 1
or SEQ ID NO: 2 is a substitution of a lysine (K) for a proline (P). In certain embodiments, the amino acid
substitution at position 258 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a serine (S) for an
asparagine (N). In certain embodiments, the amino acid substitution at position 296 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of a tryptophan (W) for a leucine (L). In certain embodiments, the amino
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acid substitution at position 296 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a tyrosine (Y) for a
leucine (L). In certain embodiments, the amino acid substitution at position 296 of SEQ ID NO: 1 or SEQ
ID NO: 2 is a substitution of a phenylalanine (F) for a leucine (L). In certain embodiments, the amino acid
substitution at position 298 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a leucine (L) for a
methionine (M). In certain embodiments, the amino acid substitution at position 298 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of an alanine (A) for a methionine (M). In certain embodiments, the amino
acid substitution at position 298 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a valine (V) for a
methionine (M). In certain embodiments, the amino acid substitution at position 311 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of an isoleucine (I) for a proline (P). In certain embodiments, the amino
acid substitution at position 311 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a valine for a
proline (P). In certain embodiments, the amino acid substitution at position 315 of SEQ ID NO: 1 or SEQ
ID NO: 2 is a substitution of a lysine (K) for an arginine (R). In certain embodiments, the amino acid
substitution at position 319 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a glycine (G) for a
threonine (T). In certain embodiments, the amino acid substitution at position 327 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of an arginine (R) for a tyrosine (Y). In certain embodiments, the amino
acid substitution at position 328 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a valine (V) for a
tyrosine (Y). In certain embodiments, the amino acid substitution at position 340 of SEQ ID NO: 1 or SEQ
ID NO: 2 is a substitution of a glycine (G) for a cysteine (C). In certain embodiments, the amino acid
substitution at position 340 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a leucine (L) for a
cysteine (C). In certain embodiments, the amino acid substitution at position 421 of SEO ID NO: 1 or SEO
ID NO: 2 is a substitution of a histidine (H) for the aspartic acid (D). In certain embodiments, the amino
acid substitution at position 436 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of an isoleucine (I)
for a valine (V). In certain embodiments, the amino acid substitution at position 456 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of a tyrosine (Y) for a methionine (M). In certain embodiments, the amino
acid substitution at position 470 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a phenylalanine
(F) for a leucine (L). In certain embodiments, the amino acid substitution at position 485 of SEQ ID NO: 1
or SEQ ID NO: 2 is a substitution of a lysine (K) for a serine (S). In certain embodiments, the amino acid
substitution at position 503 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a leucine (L) for a
methionine (M). In certain embodiments, the amino acid substitution at position 503 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of an isoleucine (I) for a methionine (M). In certain embodiments, the
amino acid substitution at position 552 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a lysine (K)
for a valine (V). In certain embodiments, the amino acid substitution at position 570 of SEO ID NO: 1 or
SEQ ID NO: 2 is a substitution of a threonine (T) for an alanine (A). In certain embodiments, the amino
acid substitution at position 591 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a proline (P) for a
glutamine (Q). In certain embodiments, the amino acid substitution at position 591 of SEQ ID NO: 1 or
SEQ ID NO: 2 is a substitution of an arginine (R) for a glutamine (Q).
(34) In certain embodiments of the disclosure, including those embodiments wherein the transposase
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comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac™ transposase enzyme may comprise or the Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more of positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 1 or SEQ ID NO: 2. In certain embodiments of the methods of the disclosure, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> transposase enzyme may comprise or the Super piggyBac<sup>™</sup> transposase enzyme may further comprise an amino acid substitution at two, three, four, five, six or more of positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 1 or SEQ ID NO: 2. In certain embodiments, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> transposase enzyme may comprise or the Super piggyBac<sup>™</sup> transposase enzyme may further comprise an amino acid substitution at positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 1 or SEQ ID NO: 2. In certain embodiments, the amino acid substitution at position 103 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a proline (P) for a serine (S). In certain embodiments, the amino acid substitution at position 194 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a valine (V) for a methionine (M). In certain embodiments, the amino acid substitution at position 372 of SEQ ID NO: 1 or SEQ ID NO: 2 is a

substitution of an alanine (A) for an arginine (R). In certain embodiments, the amino acid substitution at position 375 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of an alanine (A) for a lysine (K). In certain embodiments, the amino acid substitution at position 450 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of an asparagine (N) for an aspartic acid (D). In certain embodiments, the amino acid substitution at position 509 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a glycine (G) for a serine (S). In certain embodiments, the amino acid substitution at position 570 of SEQ ID NO: 1 or SEQ ID NO: 2 is a substitution of a serine (S) for an asparagine (N). In certain embodiments, the piggyBac™ transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 1. In certain embodiments, including those embodiments wherein the piggyBac™ transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 1, the piggyBac<sup>™</sup> transposase enzyme may further comprise an amino acid substitution at positions 372, 375 and 450 of the sequence of SEQ ID NO: 1 or SEQ ID NO: 2. In certain embodiments, the piggyBac<sup>TM</sup> transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 1, a substitution of an alanine (A) for an arginine (R) at position 372 of SEQ ID NO: 1, and a substitution of an alanine (A) for a lysine (K) at position 375 of SEQ ID NO: 1. In certain embodiments, the piggyBac<sup>TM</sup> transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 1, a substitution of an alanine (A) for an arginine (R) at position 372 of SEQ ID NO: 1, a substitution of an alanine (A) for a lysine (K) at position 375 of SEQ ID NO: 1 and a substitution of an asparagine (N) for an aspartic acid (D) at position 450 of SEQ ID NO: 1. (35) In certain embodiments of the disclosure, the transposase enzyme is a Sleeping Beauty transposase enzyme (see, for example, U.S. Pat. No. 9,228,180, the contents of which are incorporated herein in their entirety). In certain embodiments, the Sleeping Beauty transposase is a hyperactive Sleeping Beauty (SB100X) transposase. In certain embodiments, the Sleeping Beauty transposase enzyme comprises an amino acid sequence at least 75% identical to:

(36) TABLE-US-00005 (SEQ ID NO: 17031)

MGKSKEISQDLRKKIVDLHKSGSSLGAISKRLKVPRSSVQTIVRKYKHHG
TTQPSYRSGRRRYLSPRDERTLVRKVQINPRTTAKDLVKMLEETGTKVSI
STVKRVLYRHNLKGRSARKKPLLQNRHKKARLRFATAHGDKDRTFWRNVL
WSDETKIELFGHNDHRYVWRKKGEACKPKNTIPTVKHGGGSIMLWGCFAA
GGTGALHKIDGIMRKENYVDILKQHLKTSVRKLKLGRKWVFQMDNDPKHT
SKVVAKWLKDNKVKVLEWPSQSPDLNPIENLWAELKKRVRARRPTNLTQL
HQLCQEEWAKIHPTYCGKLVEGYPKRLTQVKQFKGNATKY.

- (37) In certain embodiments, including those wherein the Sleeping Beauty transposase is a hyperactive Sleeping Beauty (SB100X) transposase, the Sleeping Beauty transposase enzyme comprises an amino acid sequence at least 75% identical to:
- (38) TABLE-US-00006 (SEQ ID NO. 17032)

MGKSKEISQDLRKRIVDLHKSGSSLGAISKRLAVPRSSVQTIVRKYKHHG
TTQPSYRSGRRRYLSPRDERTLVRKVQINPRTTAKDLVKMLEETGTKVSI
STVKRVLYRHNLKGHSARKKPLLQNRHKKARLRFATAHGDKDRTFWRNVL
WSDETKIELFGHNDHRYVWRKKGEACKPKNTIPTVKHGGGSIMLWGCFAA
GGTGALHKIDGIMDAVQYVDILKQHLKTSVRKLKLGRKWVFQHDNDPKHT
SKVVAKWLKDNKVKVLEWPSQSPDLNPIENLWAELKKRVRARRPTNLTQL
HQLCQEEWAKIHPNYCGKLVEGYPKRLTQVKQFKGNATKY.

(39) In certain embodiments of the compositions of the disclosure, the first transposon and/or the second transposon further comprises a selection gene. In certain embodiments, the selection gene comprises neo, DHFR (Dihydrofolate Reductase), TYMS (Thymidylate Synthetase), MGMT (O(6)-methylguanine-DNA methyltransferase), multidrug resistance gene (MDR1), ALDH1 (Aldehyde dehydrogenase 1 family, member A1), FRANCF, RAD51C (RAD51 Paralog C), GCS (glucosylceramide synthase), NKX2.2 (NK2 Homeobox 2) or any combination thereof. In certain embodiments, the selection gene comprises DHFR (40) In certain embodiments of the compositions of the disclosure, the first transposon and or the second transposon comprises an inducible caspase polypeptide comprising (a) a ligand binding region, (b) a linker, and (c) a truncated caspase 9 polypeptide, wherein the inducible caspase polypeptide does not comprise a non-human sequence. In certain embodiments, the non-human sequence is a restriction site. In

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certain embodiments, the ligand binding region inducible caspase polypeptide comprises a FK506 binding
protein 12 (FKBP12) polypeptide. In certain embodiments, the amino acid sequence of the FK506 binding
protein 12 (FKBP12) polypeptide comprises a modification at position 36 of the sequence. In certain
embodiments, the modification is a substitution of valine (V) for phenylalanine (F) at position 36 (F36V).
In certain embodiments, the FKBP12 polypeptide is encoded by an amino acid sequence comprising
GVQVETISPGDGRTFPKRGQTCVVHYTGMLEDGKKVDSSRDRNKPFKFMLGKQEVI
RGWEEGVAQMSVGQRAKLTISPDYAYGATGHPGIIPPHATLVFDVELLKLE (SEQ ID NO: 17012).
In certain embodiments, the FKBP12 polypeptide is encoded by a nucleic acid sequence comprising
(41) TABLE-US-00007 (SEQ ID NO: 17013)
GGGGTCCAGGTCGAGACTATTTCACCAGGGGATGGGCGAACATTTCCAAA
AAGGGGCCAGACTTGCGTCGTGCATTACACCGGGATGCTGGAGGACGGGA
AGAAAGTGGACAGCTCCAGGGATCGCAACAAGCCCTTCAAGTTCATGCTG
GGAAAGCAGGAAGTGATCCGAGGATGGGAGGAAGGCGTGGCACAGATGTC
AGTCGGCCAGCGGCCAAACTGACCATTAGCCCTGACTACGCTTATGGAG
CAACAGGCCACCCAGGGATCATTCCCCCTCATGCCACCCTGGTCTTCGAT
GTGGAACTGCTGAAGCTGGAG.
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- (42) In certain embodiments, the linker region of the inducible proapoptotic polypeptide is encoded by an amino acid comprising GGGS (SEQ ID NO: 17014). In certain embodiments, the linker region of the inducible proapoptotic polypeptide is encoded by a nucleic acid sequence comprising GGAGGAGGAGGATCC (SEO ID NO: 17015).
- (43) In certain embodiments, the truncated caspase 9 polypeptide of the inducible proapoptotic polypeptide is encoded by an amino acid sequence that does not comprise an arginine (R) at position 87 of the sequence. In certain embodiments, the truncated caspase 9 polypeptide of the inducible proapoptotic polypeptide is encoded by an amino acid sequence that does not comprise an alanine (A) at position 282 the sequence. In certain embodiments, the truncated caspase 9 polypeptide of the inducible proapoptotic polypeptide is encoded by an amino acid comprising

GFGDVGALESLRGNADLAYILSMEPCGHCLIINNVNFCRESGLRTRTGSNIDCEKLRR RFSSLHFMVEVKGDLTAKKMVLALLELAQQDHGALDCCVVVILSHGCQASHLQFPG AVYGTDGCPVSVEKIVNIFNGTSCPSLGGKPKLFFIQACGGEQKDHGFEVASTSPEDE SPGSNPEPDATPFQEGLRTFDQLDAISSLPTPSDIFVSYSTFPGFVSWRDPKSGSWYVE TLDDIFEQWAHSEDLQSLLLRVANAVSVKGIYKQMPGCFNFLRKKLFFKTS (SEQ ID NO: 17016). In certain embodiments, the truncated caspase 9 poly peptide of the inducible proapoptotic polypeptide is encoded by a nucleic acid sequence comprising

(44) TABLE-US-00008 (SEQ ID NO: 17017)

TTTGGGGACGTGGGGCCCTGGAGTCTCTGCGAGGAAATGCCGATCTGGC TTACATCCTGAGCATGGAACCCTGCGGCCACTGTCTGATCATTAACAATG TGAACTTCTGCAGAGAAAGCGGACTGCGAACACGGACTGGCTCCAATATT GACTGTGAGAAGCTGCGGAGAAGGTTCTCTAGTCTGCACTTTATGGTCGA AGTGAAAGGGGATCTGACCGCCAAGAAAATGGTGCTGGCCCTGCTGGAGC TGGCTCAGCAGGACCATGGAGCTCTGGATTGCTGCGTGGTCGTGATCCTG TCCCACGGGTGCCAGGCTTCTCATCTGCAGTTCCCCGGAGCAGTGTACGG AACAGACGGCTGTCCTGTCAGCGTGGAGAAGATCGTCAACATCTTCAACG GCACTTCTTGCCCTAGTCTGGGGGGAAAGCCAAAACTGTTCTTTATCCAG GCCTGTGGCGGGAACAGAAAGATCACGGCTTCGAGGTGGCCAGCACCAG CCCTGAGGACGAATCACCAGGGAGCAACCCTGAACCAGATGCAACTCCAT TCCAGGAGGACTGAGGACCTTTGACCAGCTGGATGCTATCTCAAGCCTG CCCACTCCTAGTGACATTTTCGTGTCTTACAGTACCTTCCCAGGCTTTGT CTCATGGCGCGATCCCAAGTCAGGGAGCTGGTACGTGGAGACACTGGACG ACATCTTTGAACAGTGGGCCCATTCAGAGGACCTGCAGAGCCTGCTGCTG CGAGTGGCAAACGCTGTCTCTGTGAAGGGCATCTACAAACAGATGCCCGG GTGCTTCAATTTTCTGAGAAAGAAACTGTTCTTTAAGACTTCC.

(45) In certain embodiments, the inducible proapoptotic polypeptide is encoded by an amino acid sequence comprising

GVQVETISPGDGRTFPKRGQTCVVHYTGMLEDGKKVDSSRDRNKPFKFMLGKQEVI RGWEEGVAQMSVGQRAKLTISPDYAYGATGHPGIIPPHATLVFDVELLKLEGGGGS GFGDVGALESLRGNADLAYILSMEPCGHCLIINNVNFCRESGLRTRTGSNIDCEKLRR RFSSLHFMVEVKGDTTAKKMVLALLELAQQDHGALDCCVVVILSHGCQASHLQFPG AVYGTDGCPVSVEKIVNIFNGTSCPSLGGKPKLFFIQACGGEQKDHGFEVASTSPEDE SPGSNPEPDATPFQEGLRTFDQLDAISSLPTPSDIFVSYSTFPGFVSWRDPKSGSWYVE TLDDIFEQWAHSEDLQSLLLRVANAVSVKGIYKQMPGCFNFLRKKLFFKTS (SEQ ID NO: 17018) In certain embodiments, the inducible proapoptotic polypeptide is encoded by a nucleic acid sequence comprising

(46) TABLE-US-00009 (SEQ ID NO: 17019) Ggggtccaggtcgagactatttcaccaggggatgggcgaacatttccaaa aaggggccagacttgcgtcgtgcattacaccgggatgctggaggacggga

agaaagtggacagctccagggatcgcaacaagcccttcaagttcatgctg

ggaaagcaggaagtgatccgaggatgggaggaaggcgtggcacagatgtc

agtcggccagcgggccaaactgaccattagccctgactacgcttatggag caacaggccacccagggatcattccccctcatgccaccctggtcttcgat gtggaactgctgaagctggaggaggaggaggatccggatttggggacgt

gggggccctggagtctctgcgaggaaatgccgatctggcttacatcctga gcatggaaccctgcggccactgtctgatcattaacaatgtgaacttctgc agagaaagcggactgcgaacacggactggctccaatattgactgtgagaa

gctgcggagaaggttctctagtctgcactttatggtcgaagtgaaagggg

atctgaccgccaagaaaatggtgctggccctgctggagctggctcagcag

gaccatggagctctggattgctgcgtggtcgtgatcctgtcccacgggtg ccaggcttctcatctgcagttccccggagcagtgtacggaacagacggct gtcctgtcagcgtggagaagatcgtcaacatcttcaacggcacttcttgc cctagtctggggggaaagccaaaactgttctttatccaggcctgtggcgg ggaacagaaagatcacggcttcgaggtggccagcaccagccctgaggacg

aatcaccagggagcaaccctgaaccagatgcaactccattccaggaggga

ctgaggacctttgaccagctggatgctatctcaagcctgcccactcctag tgacattttcgtgtcttacagtaccttcccaggctttgtctcatggcgcgatcccaagtcagggagctggtacgtggagacactggacgacatctttgaa

cagtgggcccattcagaggacctgcagagcctgctgctgcgagtggcaaa

cgctgtctctgtgaagggcatctacaaacagatgcccgggtgcttcaatt ttctgagaaagaaactgttctttaagacttcc.

- (47) In certain embodiments of the compositions of the disclosure, the first transposon and/or the second transposon comprises at least one self-cleaving peptide. In certain embodiments, the at least one selfcleaving peptide comprises a T2A peptide, a GSG-T2A peptide, an E2A peptide, a GSG-E2A peptide, an F2A peptide, a GSG-F2A peptide, a P2A peptide, or a GSG-P2A peptide. In certain embodiments, the at least one self-cleaving peptide comprises a T2A peptide. In certain embodiments, the T2A peptide comprises an amino acid sequence comprising EGRGSLLTCGDVEENPGP (SEQ ID NO: 17020). In certain embodiments, the GSG-T2A peptide comprises an amino acid sequence comprising GSGEGRGSLLTCGDVEENPGP (SEQ ID NO: 17021). In certain embodiments, the E2A peptide comprises an amino acid sequence comprising QCTNYALLKLAGDVESNPGP (SEQ ID NO: 17022). In certain embodiments, the GSG-E2A peptide comprises an amino acid sequence comprising GSGQCTNYALLKLAGDVESNPGP (SEQ ID NO: 17023). In certain embodiments, the F2A peptide comprises an amino acid sequence comprising VKQTLNFDLLKLAGDVESNPGP (SEQ ID NO: 17024). In certain embodiments, the GSG-F2A peptide comprises an amino acid sequence comprising GSGVKQTLNFDLLKLAGDVESNPGP (SEQ ID NO: 17025). In certain embodiments, the P2A peptide comprises an amino acid sequence comprising ATNFSLLKQAGDVEENPGP (SEQ ID NO: 17026). In certain embodiments, the GSG-P2A peptide comprises an amino acid sequence comprising GSGATNFSLLKQAGDVEENPGP (SEQ ID NO: 17027). In certain embodiments, the at least one selfcleaving peptide is positioned between (a) the selection gene and the inducible transgene construct or (b) the inducible transgene construct and the inducible caspase polypeptide. In certain embodiments, the at least one self-cleaving peptide is positioned between (a) the selection gene and the reporter construct or (b) the reporter construct and the inducible caspase polypeptide.
- (48) The disclosure provides a cell comprising the composition of the disclosure.
- (49) The disclosure provides a method of inducing conditional gene expression in a cell comprising (a) contacting the cell with a composition of the disclosure, under conditions suitable to allow for integration of the inducible transgene construct into the genome of the cell and for the expression of the exogenous reporter and (b) contacting the exogenous receptor and a ligand that specifically binds thereto, to transduce

an intracellular signal that targets the inducible promoter, thereby modifying gene expression. In certain embodiments, the cell is in vivo, ex vivo, in vitro or in situ. In certain embodiments, the cell is an immune cell. In certain embodiments, the immune cell is a T-cell, a Natural Killer (NK) cell, a Natural Killer (NK)-like cell, a hematopoeitic progenitor cell, a peripheral blood (PB) derived T cell or an umbilical cord blood (UCB) derived T-cell. In certain embodiments, the immune cell is a T-cell. In certain embodiments, the cell is allogeneic.

- (50) The disclosure provides a method of treating a disease or disorder in a subject in need thereof, comprising administering to the subject a composition of the disclosure, under conditions suitable to allow for integration of the inducible transgene construct into the genome of the cell and for the expression of the exogenous reporter, and administering a ligand to which the exogenous receptor selectively binds, wherein the binding of the ligand to the exogenous receptor transduces an intracellular signal to target the inducible promoter controlling the transgene, wherein the transgene is expressed, and wherein the product of the transgene is therapeutically-effective for treating the disease or disorder. In certain embodiments, the product of the transgene is a secreted protein. In certain embodiments, the secreted protein is a clotting factor. In certain embodiments, the disease or disorder is a clotting disorder.
- (51) In certain embodiments of the methods of the disclosure, conditions suitable to allow for integration of the inducible transgene construct into the genome of the cell and for the expression of the exogenous reporter comprise in vivo conditions. In certain embodiments, conditions suitable to allow for integration of the inducible transgene construct into the genome of the cell and for the expression of the exogenous reporter comprise a temperature substantially similar to an internal temperature of a human body, a CO.sub.2 level substantially similar to an internal CO.sub.2 levels of a human body, an O.sub.2 level substantially similar to an internal O.sub.2 levels of a human body, an aqueous or saline environment with a level of electrolytes substantially similar to a level of electrolytes of an interior of a human body. (52) In certain embodiments of the compositions and methods of the disclosure, the ligand to which the exogenous receptor specifically binds is non-naturally occurring. In certain embodiments, the ligand is a nucleic acid, an amino acid, a polymer, an organic small molecule, an inorganic small molecule, or a combination thereof. Exemplary ligands include, but are not limited to, synthetic, modified, recombinant, mutant, chimeric, endogenous or non-naturally occurring, proteins (soluble or membrane-bound), steroid hormones, gas particles, nucleic acids, growth factors, neurotransmitters, vitamins, and minerals. (53) The disclosure provides a composition comprising (a) an inducible transgene construct, comprising a sequence encoding an inducible promoter and a sequence encoding a transgene, and (b) a ligand construct, comprising a sequence encoding a constitutive promoter and a sequence encoding an exogenous ligand, wherein, upon integration of the construct of (a) and the construct of (b) into a genomic sequence of a cell, the exogenous ligand is expressed, and wherein the exogenous ligand, upon binding a receptor, transduces an intracellular signal that targets the inducible promoter of (a) to modify gene expression. In certain embodiments, the ligand comprises a non-natural or synthetic sequence. In certain embodiments, the ligand comprises a fusion protein. In certain embodiments, the ligand is bound to the surface of the cell. In certain embodiments, the ligand comprises an intracellular domain. In certain embodiments, the intracellular domain transduces a signal in the cell expressing the ligand. In certain embodiments, the structure of the ligand is substantially similar to the structure of the receptor of the compositions of the disclosure. In certain embodiments, the signal transduced by the ligand and the signal transduced by the receptor comprise a bi-directional signal.

## Description

#### BRIEF DESCRIPTION OF THE DRAWINGS

(1) FIG. **1**A-B is a pair of schematic diagrams depicting NF-KB inducible vectors for expression in T-cells. Two T cell activation NF-KB inducible vectors were developed; one with the gene expression system (GES) in the forward orientation (A) and the other in the complementary direction (B), both preceding the constitutive EF1a promoter. These vectors also direct expression of a CAR molecule and a DHFR selection gene, separated by a T2A sequence. Both the conditional NF-KB inducible system and the EF1a directed genes are a part of a piggyBac transposon which can be permanently integrated into T

cells using electroporation (EP). Once integrated into the genome, the T cells will constitutively express the CAR on the membrane surface and the DHFR within the cell, while expression of the NF-KB inducible gene, GFP, will be expressed to the highest level only upon T cell activation.

- (2) FIG. **2** is a pair of graphs depicting NF-KB inducible expression of GFP in activated T cells. T cells were nucleofected with a piggyBac vector expressing an anti-BCMA CAR and a DHFR mutein gene under control of an EF1a promoter along with the absence (No GES control) or presence of an NF-KB inducible expression system driving GFP expression in either the forward (pNFKB-GFP forward) or reverse orientation (pNFKB-GFP reverse). Cells were cultured in the presence of methotrexate selection until the cells were almost completely resting (Day 19) and GFP expression was assessed at Day 5 and Day 19. At Day 5, all T cells are proliferating and highly stimulated, with cells harboring the NF-KB inducible expression cassette producing high levels of GFP due to strong NFκB activity. The No GES control cells did not express detectable levels of GFP. By Day 19, the GES T cells were almost fully resting and GFP expression was significantly lower than Day 5 (\*\*MFI), since NFκB activity is lower. GFP expression is still observed at Day 19, which may due to the long half-life of GFP protein (\*30 hr), or, basal level of NFκB activity through, for example, a TCR, a CAR, a cytokine receptor, or a growth factor receptor signal.
- (3) FIG. **3** is a series of graphs depicting anti-BCMA CAR-mediated activation of NF-KB inducible expression of GFP in presence of BCMA+ tumor cells. T cells were either unmodified (Mock T cells) or nucleofected with a piggyBac vector expressing an anti-BCMA CAR and a DHFR mutein gene under control of an EF1a promoter along with the absence (No GES control) or presence of an NF-KB inducible expression system driving GFP expression in either the forward (pNFKB-GFP forward) or reverse orientation (pNFKB-GFP reverse). All cells were cultured for 22 days, either with or without methotrexate selection (Mock T cells), until the cells were almost completely resting. Cells were then stimulated for 3 days in the absence (No stimulation) or presence of BCMA- (K562), BMCA+ (RPMI 8226), or positive control anti-CD3 anti-CD28 activation reagent (CD3/28 stimulation). GFP expression was undetectable under all conditions with the No GES control or Mock T cells. However, while pNFKB-GFP forward- and reverse-transposed cells exhibited little GFP expression over the No stimulation control when cultured with BCMA- K562 cells, they both demonstrated dramatic upregulation of gene expression either in the presence of BCMA+ tumor cells or under positive control conditions. Little difference in GFP expression was observed between the pNFKB-GFP forward- and reverse-transposed cells that were cocultured with BCMA+ tumor cells.
- (4) FIG. 4 is a series of graphs demonstrating that the Expression level of inducible gene can be regulated by number of response elements preceding the promoter T cells were nucleofected with a piggyBac vector encoding an anti-BCMA CARTyrin followed by a selection gene, both under control of a human EF1a promoter. Further, vectors either additionally encoded the conditional NF-KB inducible gene expression system driving expression of a truncated CD19 protein (dCD19) and included a number of NFKB response elements (RE) varying from 0-5, no GES (No GES), or received an electroporation pulse but no piggyBac nucleic acid (Mock). Data are shown for only the GES in the reverse (opposite) direction/orientation. All cells were cultured for 18 days and included selection for piggyBac-modified T cells using methotrexate addition. Cells were then stimulated for 3 days using anti-CD3 anti-CD28 bead activation reagent and dCD19 surface expression was assessed by FACS at Days 0, 3 and 18, and data are shown as FACS histograms and MF of target protein staining. Surface dCD19 expression was detected at low levels at Day 0 in all T cells transposed with vectors encoding the GES. At 3 days post-stimulation, dramatic upregulation of dCD19 expression was observed for all T cells expressing the GES, with a greater fold increase in surface expression in those with higher numbers of REs. Thus, surface dCD19 expression was directly proportional with the number of REs encoded in the GES. No dCD19 was detected on the surface of T cells that did not harbor the GES: No GES and Mock controls. (5) FIG. **5** is a schematic diagram showing the human coagulation pathway leading to blood clotting. Contact activation, for example by damaging an endothelium, activates an intrinsic clotting pathway. Tissue factors activate an extrinsic clotting pathway, for example following trauma Both pathways converge onto the conversion of Prothrombin into Thrombin, which catalyzes the conversion of fibrinogen into fibrin. Polymerized fibrin together with platelets forms a clot. In the absence of Factor IX (circled), clotting is defective. Factor VII (FVIII) deficiency leads to development of Hemophilia A. Factor IX (FIX)

deficiency leads to development of Hemophilia B. Prior to the compositions and methods of the disclosure, the standard treatment for hemophilia B involved an infusion of recombinant FIX every 2 to 3 days, at an expense of approximately \$250,000 per year. In sharp contrast to this standard treatment option, T cells of the disclosure are maintained in humans for several decades.

- (6) FIG. **6** is a series of Fluorescence-Activated Cell Sorting (FACS plots) depicting FIX-secreting T cells. T cells encoding a human Factor IX transgene showed a T-cell phenotype in approximately 80% of cells. The 6 panels are described in order from left to right. (1) Forward scatter (FSC) on the x-axis versus side scatter (SSC) on the y-axis. The x-axis is from 0 to 250 thousand (abbreviated k) in increments of 50k, the y-axis is for 0 to 250k, in increments of 50k. (2) FSC on the x-axis versus the cell viability marker 7 aminoactinomycin D (7AAD). The x-axis is labeled from 0 to 250k in increments of 50k. The y-axis reads, from top to bottom, -10.sup.3, 0, 10.sup.3, 10.sup.4, 10.sup.5. (3) On the x-axis is shown anti-CD56-APC conjugated to a Cy7 dye (CDCl56-APC-Cy7), units from 0 to 10.sup.5 incrementing in powers of 10. On the y-axis is shown anti-CD3 conjugated to phycoerythrin (PE), units from 0 to 10.sup.5 incrementing in powers of 10. (4) On the x-axis is shown anti-CD8 conjugated to fluorescein isothiocyanate (FITC), units from 0 to 10.sup.5 incrementing in powers of 10. On the y-axis is shown anti-CD4 conjugated to Brilliant Violet 650 dye (BV650), units from 0 to 10.sup.5 incrementing in powers of 10. (5) On the x-axis is shown an anti CD62L antibody conjugated to a Brilliant Violet 421 dye (BV421), units from 0 to 10.sup.5 incrementing in powers of 10 On the y-axis is shown an anti-CD45RA antibody conjugated to PE and Cy7, units from 0 to 10.sup.5 incrementing in powers of 10. This panel is boxed. (6) On the x-axis is shown an anti-CCR7 antibody conjugated to Brilliant Violet 786 (BV786), units from 0 to 10 incrementing in powers of 10. On the y-axis is shown anti-CD45RA conjugated to PE and Cy7, units from 0 to 10.sup.5 incrementing in powers of 10.
- (7) FIG. 7A is a graph showing human Factor IX secretion during production of modified T cells of the disclosure. On the y-axis, Factor IX concentration in nanograms (ng) per milliliter (mL) from 0 to 80 in increments of 20. On the x-axis are shown 9 day and 12 day T cells.
- (8) FIG. **7**B is a graph showing the clotting activity of the secreted Factor IX produced by the T cells. On the y-axis is shown percent Factor IX activity relative to human plasma, from 0 to 8 in increments of 2. On the x-axis are 9 and 12 day T cells.
- (9) FIG. **8** is a series of graphs demonstrating that the expression level of inducible gene can be regulated by number of response elements preceding the promoter in CD4 positive cells. Truncated CD19 (dCD19) expressing CAR-T cells were stimulated by BCMA+ H929 multiple myeloma cells at 2:1 CAR-T:H929 ratio. The expression of dCD19 was driven by the minimal promoter that enhanced by 0, 1, 2, 3, 4 or 5 repeats of the NF-kB response element. The expression of BCMA CAR was driven by human elongation factor- $\alpha$  (EF-1 $\alpha$ ) promoter, a constitutive promoter that is commonly used for gene expression in human T cells. Before tumor cell stimulation, the expression of CAR and dCD19 were both at basal levels compared to mock T cell control. The expression levels of CAR and dCD19 were both upregulated upon tumor stimulation (day 3) and then subsequently downregulated (day 9, 14) and eventually reached their respective basal levels when the cells resume a fully rested status again (day 20). However, CAR surface expression was equivalently up- or down-regulated in all the CAR-T cell samples during cell activation and resting process, while the expression levels of dCD19 were directly proportional to the number of NFκB response elements (day 3, 9, 14). Data are shown as FACS histograms and MFI of target protein staining. Thus, surface dCD19 expression was directly proportional with the number of REs encoded in the GES. No dCD19 was detected on the surface of T cells that did not harbor the GES: No GES and Mock controls.
- (10) FIG. **9** is a series of graphs demonstrating that the expression level of inducible gene can be regulated by number of response elements preceding the promoter in CD8 positive cells. Truncated CD19 (dCD19) expressing CAR-T cells were stimulated by BCMA+ H929 multiple myeloma cells at 2:1 CAR-T:H929 ratio. The expression of dCD19 was driven by the minimal promoter that enhanced by 0, 1, 2, 3, 4 or 5 repeats of the NF-kB response element. The expression of BCMA CAR was driven by human elongation factor- $1\alpha$  (EF- $1\alpha$ ) promoter, a constitutive promoter that is commonly used for gene expression in human T cells. Before tumor cell stimulation, the expression of CAR and dCD19 were both at basal levels compared to mock T cell control. The expression levels of CAR and dCD19 were both upregulated upon tumor stimulation (day 3) and then subsequently downregulated (day 9, 14) and eventually reached their

respective basal levels when the cells resume a fully rested status again (day 20). However, CAR surface expression was equivalently up- or down-regulated in all the CAR-T cell samples during cell activation and resting process, while the expression levels of dCD19 were directly proportional to the number of NF-  $\kappa$ B response elements (day 3, 9, 14). Data are shown as FACS histograms and MFI of target protein staining. Thus, surface dCD19 expression was directly proportional with the number of REs encoded in the GES. No dCD19 was detected on the surface of T cells that did not harbor the GES: No GES and Mock controls.

- (11) FIG. **10** is a bar graph depicting the knock out efficiency of targeting various checkpoint signaling proteins that could be used to armor T-cells. Cas-CLOVER was used to knockout the checkpoint receptors. PD-1, TGFBR2, LAG-3, TIM-3 and CTLA-4 in resting primary human pan T cells. Percent knock-out is shown on the y-axis. Gene editing resulted in 30-70% loss of protein expression at the cell surface as measured by flow cytometry.
- (12) FIG. **11** is a series of schematic diagrams of wildtype, null and switch receptors and their effects on intracellular signaling, either inhibitory or stimulatory, in primary T-cells. Binding of the wildtype inhibitory receptor expressed endogenously on a T-cell with its endogenous ligand results in transmission of an inhibitory signal which, in part, reduces T-cell effector function. However, mutation (Mutated null) or deletion (Truncated null) of the intracellular domain (ICD) of a checkpoint receptor protein, such as PD1 (top panel) or TGFBRII (bottom panel), reduces or eliminates its signaling capability when cognate ligand(s) is bound. Thus, expression of engineered mutated or truncated null receptors on the surface of modified T cells results in a competition with endogenously-expressed wildtype receptors for binding of the free endogenous ligand(s), effectively reducing or eliminating delivery of inhibitory signals by endogenously-expressed wildtype receptors. Specifically, any binding by a mutated or null receptor sequesters the endogenous ligand(s) from binding the wildtype receptor and results in dilution of the overall level of checkpoint signaling effectively delivered to the modified T-cell, thereby reducing or blocking checkpoint inhibition and functional exhaustion of the modified T cells. A switch receptor is created by replacement of the wildtype ICD with an ICD from either a co-stimulatory molecule (such as CD3z, CD28, 4-1BB) or a different inhibitory molecule (such as CTLA4, PD1, Lag3). In the former case, binding of the endogenous ligand(s) by the modified switch receptor results in the delivery of a positive signal to the T-cells, thereby helping to enhance stimulation of the modified T cell and potentially enhance target tumor cell killing. In the latter case, binding of the endogenous ligand(s) by the modified switch receptor results in the delivery of a negative signal to the T-cells, thereby eliminating stimulation of the modified T cell and potentially reducing target tumor cell killing. The signal peptide (purple arrow), extracellular domain (ECD) (bright green), transmembrane domain (yellow), intracellular signaling domain (ICD)(orange), and replacement ICD (green) are displayed in the receptor diagrams. "\*" indicates a mutated ICD. "+" indicates the presence of a checkpoint signal. "-" indicates the absence of a checkpoint signal.
- (13) FIG. **12** is a schematic diagram showing an example of the design of null receptors with specific alterations that may help to increase expression of the receptor on the surface of modified T cells. Examples are shown for PD1 and TGFBRII null receptors and the signal peptide domain (SP), transmembrane domain (TM) and extracellular domain (ECD) of truncated null receptors for PD1 (top panel) and TGFBRII (bottom panel) are displayed. The first of the top four molecules is the wildtype PD-1 receptor, which encodes the wildtype PD-1 SP and TM. For the PD1 null receptor, replacement of PD1 wildtype SP or TM domain (green; light green) with the SP or TM domain of a human T cell CD8a receptor (red) is depicted. The second molecule encodes the CD8a SP along with the native PD-1 TM, the third encodes the wildtype PD-1 SP and the alternative CD8a TM, and the fourth encodes both the alternative CD8a SP and TM. Similarly, for the null receptor of TGFβRII, replacement of the wildtype TGFBRII SP (pink) with a SP domain of a human T cell CD8a receptor (red). The names of the constructs and the amino acid lengths (aa) of each construct protein is listed on the left of the diagram. (14) FIG. **13** is a series of histograms depicting the expression of the PD1 and TGFBRII null Receptors on
- the surface of modified primary human T cells as determined by flow cytometry. Each of the six truncated null constructs from FIG. 12 were expressed on the surface of primary human T cells. T cells were stained with either anti-PD1 (top; blue histograms) or anti-TGF $\beta$ RII (bottom; blue histograms), or isotype control or secondary only (gray histograms). Cells staining positive for PD-1 or TGF $\beta$ RII expression were gated

(frequency shown above gate) and mean fluorescence intensity (MFI) value is displayed above each positive histogram. The names of the null receptor constructs are depicted above each plot. Both null receptor gene strategies, replacement of the wildtype SP with the alternative CD8 $\alpha$  were successfully expressed. 02.8aSP-PD- and 02.8aSP-TGF $\beta$ RII resulted in the highest level of expression at the T-cell surface. 02.8aSP-PD-1 null receptor exhibited an MFI of 43,680, which is 177-fold higher than endogenous T cell PD-1 expression and 2.8-fold higher than the wildtype PD-1 null receptor. 02.8aSP-TGF $\beta$ RII null receptor exhibited an MFI of 13,809, which is 102-fold higher than endogenous T cell TGF $\beta$ RII expression and 1.8-fold higher than the wildtype TGF $\beta$ RII null receptor. Replacement of wildtype SP with the alternative CD8 $\alpha$  SP for both PD1 and TGRBRII results in enhanced surface expression of the null or Switch receptor, which may help to maximize reduction or blockage of checkpoint inhibition upon binding and sequestration of the endogenous ligand(s).

- (15) FIG. **14** is a schematic depiction of the Csy4-T2A-Clo051-G4Slinker-dCas9 construct map (Embodiment 2).
- (16) FIG. **15** is a schematic depiction of the pRT-Clo051-dCas9 Double NLS construct map (Embodiment 1).
- (17) FIG. **16** is a pair of graphs comparing the efficacy of knocking out expression of either B2M on the surface of Pan T-cells (left) or the  $\alpha$ -chain of the T-cell Receptor on the surface of Jurkat cells (right) for either Embodiment 1 (pRT1-Clo051-dCas9 Double NLS, as shown in FIG. **15**) or Embodiment 2 (Csy4-T2A-Clo051-G4Slinker-dCas9, as shown in FIG. **14**) of a Cas-Clover fusion protein of the disclosure. For the right-hand graph, the fusion protein is provided at either 10  $\mu$ g or 20  $\mu$ g, as indicated. (18) FIG. **17** is a photograph of a gel electrophoresis analysis of mRNA encoding each of Embodiment 1 (Lane 2; pRT1-Clo051-dCas9 Double NLS, as shown in FIG. **15**) or Embodiment 2 (Lane 3; Csy4-T2A-Clo051-G4Slinker-dCas9, as shown in FIG. **14**). In addition, a previous preparation ("old version") of mRNA encoding Embodiment 2 is included (Lane 4) for comparison. As shown, all mRNA samples encoding the two different embodiments migrate as distinct bands within the gel, are of high quality, and are similar in size, as expected.

#### DETAILED DESCRIPTION

- (19) The disclosure provides a composition comprising (a) an inducible transgene construct, comprising a sequence encoding an inducible promoter and a sequence encoding a transgene, and (b) a receptor construct, comprising a sequence encoding a constitutive promoter and a sequence encoding an exogenous receptor, wherein, upon integration of the construct of (a) and the construct of (b) into a genomic sequence of a cell, the exogenous reporter is expressed, and wherein the exogenous reporter, upon binding a ligand, transduces an intracellular signal that targets the inducible promoter of (a) to modify gene expression.
- (20) Exogenous Receptors
- (21) Exogenous receptors of the disclosure may comprise a non-naturally occurring receptor. In certain embodiments, the non-naturally occurring receptor is a synthetic, modified, recombinant, mutant or chimeric receptor. In certain embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a T-cell receptor (TCR). In certain embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a scaffold protein. In certain embodiments, the non-naturally occurring receptor comprises a transmembrane domain. In certain embodiments, the non-naturally occurring receptor that transduces an intracellular signal. In certain embodiments, the non-naturally occurring receptor comprises an intracellular signaling domain. In certain embodiments, the non-naturally occurring receptor is a chimeric ligand receptor (CLR). In certain embodiments, the CLR is a chimeric antigen receptor.
- (22) In certain embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In certain embodiments, the CLR is a chimeric antigen receptor. In certain embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain.
- (23) The disclosure provides chimeric receptors comprising at least one Centyrin. Chimeric ligand/antigen receptors (CLRs/CARs) of the disclosure may comprise more than one Centyrin, referred to herein as a CARTyrin.
- (24) The disclosure provides chimeric receptors comprising at least one VHH. Chimeric ligand/antigen

receptors (CLRs/CARs) of the disclosure may comprise more than one VHH, referred to herein as a VCAR.

- (25) Chimeric receptors of the disclosure may comprise a signal peptide of human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\delta$ , IgG4, and/or CD4. An intracellular domain or endodomain of the disclosure may comprise an intracellular signaling domain of human CD3 $\zeta$  and may further comprise human 4-1BB, CD2 $\delta$ , CD4 $\delta$ , ICOS, MyD8 $\delta$ , OX-40 intracellular segment, or any combination thereof. Exemplary transmembrane domains include, but are not limited to a human CD2, CD3 $\delta$ , CD3 $\delta$ , CD3 $\delta$ , CD4, CD8 $\delta$ , CD19, CD2 $\delta$ , 4-1BB or GM-CSFR transmembrane domain. (26) The disclosure provides genetically modified cells, such as T cells, NK cells, hematopoietic progenitor cells, peripheral blood (PB) derived T cells (including T cells from G-CSF-mobilized peripheral blood), umbilical cord blood (UCB) derived T cells rendered specific for one or more ligands or antigens by introducing to these cells a CLR/CAR. CARTyrin and/or VCAR of the disclosure. Cells of the disclosure may be modified by electrotransfer of a transposane of the disclosure (preferably, the sequence encoding a transposase of the disclosure is an mRNA sequence).
- (27) In some embodiments, the armored T-cell comprises a composition comprising (a) an inducible transgene construct, comprising a sequence encoding an inducible promoter and a sequence encoding a transgene, and (b) a receptor construct, comprising a sequence encoding a constitutive promoter and a sequence encoding an exogenous receptor, such as a CLR or CAR, wherein, upon integration of the construct of (a) and the construct of (b) into a genomic sequence of a cell, the exogenous receptor is expressed, and wherein the exogenous receptor, upon binding a ligand or antigen, transduces an intracellular signal that targets directly or indirectly the inducible promoter regulating expression of the inducible transgene (a) to modify gene expression.
- (28) Chimeric Receptors
- (29) Chimeric antigen receptors (CARs) and/or chimeric ligand receptors (CLRs) of the disclosure may comprise (a) an ectodomain comprising an antigen/ligand recognition region, (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In certain embodiments, the ectodomain may further comprise a signal peptide. Alternatively, or in addition, in certain embodiments, the ectodomain may further comprise a hinge between the antigen/ligand recognition region and the transmembrane domain. In certain embodiments of the CARs of the disclosure, the signal peptide may comprise a sequence encoding a human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD4, CD8 $\epsilon$ , CD19, CD28, 4-1BB or GM-CSFR signal peptide. In certain embodiments of the CARs of the disclosure, the signal peptide may comprise a sequence encoding a human CD8 $\epsilon$ 0 signal peptide. In certain embodiments, the transmembrane domain may comprise a sequence encoding a human CD2, CD3 $\epsilon$ 0, CD3 $\epsilon$ 1, CD3 $\epsilon$ 2, CD3 $\epsilon$ 3, CD3 $\epsilon$ 4, CD8 $\epsilon$ 5, CD3 $\epsilon$ 7, CD3 $\epsilon$ 8, CD3 $\epsilon$ 8, CD3 $\epsilon$ 9, CD3 $\epsilon$
- (30) In certain embodiments of the CARs/CLRs of the disclosure, the at least one costimulatory domain may comprise a human 4-1BB, CD28, CD40, ICOS, MyD88, OX-40 intracellular segment, or any combination thereof. In certain embodiments of the CARs of the disclosure, the at least one costimulatory domain may comprise a CD28 and/or a 4-1BB costimulatory domain. In certain embodiments of the CARs of the disclosure, the hinge may comprise a sequence derived from a human CD8 $\alpha$ , IgG4, and/or CD4 sequence. In certain embodiments of the CARs/CLRs of the disclosure, the hinge may comprise a sequence derived from a human CD8 $\alpha$  sequence.
- (31) The CD28 costimulatory domain may comprise an amino acid sequence comprising RVKFSRSADAPAYKQGQNQLYNELNLGRREEYDVLDKRRGRDPEMGGKPRRKNPQ EGLYNELQKDKMAEAYSEIGMKGERRRGKGHDGLYQGLSIATKDTYDALHMQALP PR (SEQ ID NO: 17004) or a sequence having at least 70%, 80%, 90%, 95%, or 99% identity to the amino acid sequence comprising

RVKFSRSADAPAYKQGQNQLYNELNLGRREEYDVLDKRRGRDPEMGGKPRRKNPQ EGLYNELQKDKMAEAYSEIGMKGERRRGKGHDGLYQGLSTAIKDTYDALHMQALP PR (SEQ ID KRGRKKLLYIFKQPFMRPVQTTQEEDGCSCRFPEEEEGGCEL.

The 4-1BB costimulatory domain may be encoded by the nucleic acid sequence comprising aagaggggggaagaaactgctgtatattttcaaacagcccttcatgcgccccgtgcagactacccaggaggaagacgggtgctcc tgtcgattccctgaggaagaggggggtgtgagctg (SEQ ID NO: 17007) The 4-1BB costimulatory domain may be located between the transmembrane domain and the CD28 costimulatory domain.

(33) In certain embodiments of the CARs/CLRs of the disclosure, the hinge may comprise a sequence derived from a human CD8 $\alpha$ , IgG4, and/or CD4 sequence. In certain embodiments of the CARs/CLRs of the disclosure, the hinge may comprise a sequence derived from a human CD8 $\alpha$  sequence. The hinge may comprise a human CD8 $\alpha$  amino acid sequence comprising

TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFACD (SEQ ID NO: 17008) or a sequence having at least 70%, 80%, 90%, 95%, or 99% identity to the amino acid sequence comprising (34) TABLE-US-00011 (SEQ ID NO: 17008)

TTTPAPRPPTPAPTIASQPLSLRPEACRPAAGGAVHTRGLDFACD.

The human CD8α hinge amino acid sequence may be encoded by the nucleic acid sequence comprising (35) TABLE-US-00012 (SEQ ID NO: 17028) actaccacaccagcacctagaccaccaactccagctccaaccatc gcgagtcagccctgagtctgagacctgaggcctgcaggccagct gcaggaggaggtgtgcacaccaggggcctggacttcgcctgcagc. ScFv

- (36) The disclosure provides single chain variable fragment (scFv) compositions and methods for use of these compositions to recognize and bind to a specific target protein. ScFv compositions comprise a heavy chain variable region and a light chain variable region of an antibody. ScFv compositions may be incorporated into an antigen/ligand recognition region of a CAR or CLR of the disclosure. An antigen/ligand recognition region of a CAR or CLR of the disclosure may comprise an ScFv or an ScFv composition of the disclosure. In some embodiments, ScFvs comprise fusion proteins of the variable regions of the heavy (VH) and light (VL) chains of an immunoglobulin, wherein the VH and VL domains are connected with a linker. ScFvs retain the specificity of the original immunoglobulin, despite removal of the constant regions and the introduction of the linker. An exemplary linker comprises a sequence of GGGGSGGGGGG (SEQ ID NO: 17033).
- (37) Centyrins
- (38) Centyrins of the disclosure specifically bind to an antigen or a ligand of the disclosure. CARs and/or CLRs of the disclosure comprising one or more Centyrins that specifically bind an antigen may be used to direct the specificity of a cell, (e.g. a cytotoxic immune cell) towards a cell expressing the specific antigen. Alternatively or in addition, CLRs of the disclosure comprising a Centyrin that specifically binds a ligand antigen may transduce a signal intracellularly to induce expression of a sequence under the control of an inducible promoter.
- (39) Centyrins of the disclosure may comprise a protein scaffold, wherein the scaffold is capable of specifically binding an antigen or a ligand. Centyrins of the disclosure may comprise a protein scaffold comprising a consensus sequence of at least one fibronectin type III (FN3) domain, wherein the scaffold is capable of specifically binding an antigen or a ligand. The at least one fibronectin type III (FN3) domain may be derived from a human protein. The human protein may be Tenascin-C. The consensus sequence may comprise
- (40) TABLE-US-00013 (SEQ ID NO: 17010)
  LPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAI
  NLTVPGSERSYDLTGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT or (SEQ ID NO: 17011)
  MLPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEA

#### INLTVPGSERSYDLTGLKPGTEYTVSFYGVKGGHRSNPLSAEFTT.

- (41) A Centyrin may comprise an amino sequence having at least 50%, 55%, 60% 65%, 70%, 75%, 80%, 85%, 90%, 95%, 97%, 99% or any percentage in between of identity to the sequence of
- (42) TABLE-US-00014 (SEQ ID NO: 17010)
- LPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAI
- NLTVPGSERSYDLTGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT or (SEQ ID NO: 17011)
- MLPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEA
- INLTVPGSERSYDLTGLKPGTEYTVS1YGVKGGHRSNPLSAEFTT.
- (43) A Centyrin may comprise an amino sequence having at least 74% identity to the sequence of (44) TABLE-US-00015 (SEQ ID NO: 17010)
- LPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAI
- NLTVPGSERSYDLTGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT or (SEQ ID NO: 17011)
- MLPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEA
- INLTVPGSERSYDLTGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT.
- (45) The consensus sequence may encoded by a nucleic acid sequence comprising
- (46) TABLE-US-00016 (SEQ ID NO: 17034) atgctgcctgcaccaaagaacctggtggtgtctcatgtgacagagg atagtgccagactgtcatggactgctcccgacgccgcagccttcgatag ttttatcatcgtgtaccgggagaacatcgaaaccggcgaggccatt gtcctgacagtgccagggtccgaacgctcttatgacctgacagatc tgaagcccggaactgagtactatgtgcagatcgccggcgtcaaagg aggcaatatcagcttccctctgtccgcaatcttcaccaca.
- (47) The consensus sequence may be modified at one or more positions within (a) a A-B loop comprising or consisting of the amino acid residues TEDS (SEQ ID NO: 17035) at positions 13-16 of the consensus sequence; (b) a B-C loop comprising or consisting of the amino acid residues TAPDAAF (SEQ ID NO: 17036) at positions 22-28 of the consensus sequence; (c) a C-D loop comprising or consisting of the amino acid residues SEKVGE (SEQ ID NO: 17037) at positions 38-43 of the consensus sequence; (d) a D-E loop comprising or consisting of the amino acid residues GSER (SEQ ID NO: 17038) at positions 51-54 of the consensus sequence; (e) a E-F loop comprising or consisting of the amino acid residues GLKPG (SEQ ID NO: 17039) at positions 60-64 of the consensus sequence; (f) a F-G loop comprising or consisting of the amino acid residues KGGHRSN (SEQ ID NO: 17040) at positions 75-81 of the consensus sequence; or (g) any combination of (a)-(f). Centyrins of the disclosure may comprise a consensus sequence of at least 5 fibronectin type III (FN3) domains, at least 10 fibronectin type III (FN3) domains or at least 15 fibronectin type III (FN3) domains.
- (48) The Centyrin may bind an antigen or a ligand with at least one affinity selected from a K.sub.D of less than or equal to 10.sup.—9M, less than or equal to 10.sup.—10M, less than or equal to 10.sup.—11M, less than or equal to 10.sup.—12M, less than or equal to 10.sup.—13M, less than or equal to 10.sup.—14M, and less than or equal to 10.sup.—15M. The K.sub.D may be determined by surface plasmon resonance. (49) Antibody Mimetic
- (50) The term "antibody mimetic" is intended to describe an organic compound that specifically binds a target sequence and has a structure distinct from a naturally-occurring antibody. Antibody mimetics may comprise a protein, a nucleic acid, or a small molecule. The target sequence to which an antibody mimetic of the disclosure specifically binds may be an antigen. Antibody mimetics may provide superior properties over antibodies including, but not limited to, superior solubility, tissue penetration, stability towards heat and enzymes (e.g. resistance to enzymatic degradation), and lower production costs. Exemplary antibody mimetics include, but are not limited to, an affibody, an affilin, an affimer, an affitin, an alphabody, an anticalin, and avimer (also known as avidity multimer), a DARPin (Designed Ankyrin Repeat Protein), a Fynomer, a Kunitz domain peptide, and a monobody.
- (51) Affibody molecules of the disclosure comprise a protein scaffold comprising or consisting of one or more alpha helix without any disulfide bridges. Preferably, affibody molecules of the disclosure comprise or consist of three alpha helices. For example, an affibody molecule of the disclosure may comprise an immunoglobulin binding domain. An affibody molecule of the disclosure may comprise the Z domain of protein A.
- (52) Affilin molecules of the disclosure comprise a protein scaffold produced by modification of exposed amino acids of, for example, either gamma-B crystallin or ubiquitin. Affilin molecules functionally mimic an antibody's affinity to antigen, but do not structurally mimic an antibody. In any protein scaffold used to

make an affilin, those amino acids that are accessible to solvent or possible binding partners in a properlyfolded protein molecule are considered exposed amino acids. Any one or more of these exposed amino acids may be modified to specifically bind to a target sequence or antigen.

- (53) Affimer molecules of the disclosure comprise a protein scaffold comprising a highly stable protein engineered to display peptide loops that provide a high affinity binding site for a specific target sequence. Exemplary affimer molecules of the disclosure comprise a protein scaffold based upon a cystatin protein or tertiary structure thereof. Exemplary affimer molecules of the disclosure may share a common tertiary structure of comprising an alpha-helix lying on top of an anti-parallel beta-sheet.
- (54) Affitin molecules of the disclosure comprise an artificial protein scaffold, the structure of which may be derived, for example, from a DNA binding protein (e.g. the DNA binding protein Sac7d). Affitins of the disclosure selectively bind a target sequence, which may be the entirety or part of an antigen. Exemplary affitins of the disclosure are manufactured by randomizing one or more amino acid sequences on the binding surface of a DNA binding protein and subjecting the resultant protein to ribosome display and selection. Target sequences of affitins of the disclosure may be found, for example, in the genome or on the surface of a peptide, protein, virus, or bacteria. In certain embodiments of the disclosure, an affitin molecule may be used as a specific inhibitor of an enzyme. Affitin molecules of the disclosure may include heat-resistant proteins or derivatives thereof.
- (55) Alphabody molecules of the disclosure may also be referred to as Cell-Penetrating Alphabodies (CPAB). Alphabody molecules of the disclosure comprise small proteins (typically of less than 10 kDa) that bind to a variety of target sequences (including antigens). Alphabody molecules are capable of reaching and binding to intracellular target sequences. Structurally, alphabody molecules of the disclosure comprise an artificial sequence forming single chain alpha helix (similar to naturally occurring coiled-coil structures). Alphabody molecules of the disclosure may comprise a protein scaffold comprising one or more amino acids that are modified to specifically bind target proteins. Regardless of the binding specificity of the molecule, alphabody molecules of the disclosure maintain correct folding and thermostability.
- (56) Anticalin molecules of the disclosure comprise artificial proteins that bind to target sequences or sites in either proteins or small molecules. Anticalin molecules of the disclosure may comprise an artificial protein derived from a human lipocalin. Anticalin molecules of the disclosure may be used in place of, for example, monoclonal antibodies or fragments thereof. Anticalin molecules may demonstrate superior tissue penetration and thermostability than monoclonal antibodies or fragments thereof. Exemplary anticalin molecules of the disclosure may comprise about 180 amino acids, having a mass of approximately 20 kDa. Structurally, anticalin molecules of the disclosure comprise a barrel structure comprising antiparallel beta-strands pairwise connected by loops and an attached alpha helix. In preferred embodiments, anticalin molecules of the disclosure comprise a barrel structure comprising eight antiparallel beta-strands pairwise connected by loops and an attached alpha helix.
- (57) Avimer molecules of the disclosure comprise an artificial protein that specifically binds to a target sequence (which may also be an antigen). Avimers of the disclosure may recognize multiple binding sites within the same target or within distinct targets. When an avimer of the disclosure recognize more than one target, the avimer mimics function of a bi-specific antibody. The artificial protein avimer may comprise two or more peptide sequences of approximately 30-35 amino acids each. These peptides may be connected via one or more linker peptides. Amino acid sequences of one or more of the peptides of the avimer may be derived from an A domain of a membrane receptor. Avimers have a rigid structure that may optionally comprise disulfide bonds and/or calcium. Avimers of the disclosure may demonstrate greater heat stability compared to an antibody.
- (58) DARPins (Designed Ankyrin Repeat Proteins) of the disclosure comprise genetically-engineered, recombinant, or chimeric proteins having high specificity and high affinity for a target sequence. In certain embodiments, DARPins of the disclosure are derived from ankyrin proteins and, optionally, comprise at least three repeat motifs (also referred to as repetitive structural units) of the ankyrin protein. Ankyrin proteins mediate high-affinity protein-protein interactions. DARPins of the disclosure comprise a large target interaction surface.
- (59) Fynomers of the disclosure comprise small binding proteins (about 7 kDa) derived from the human Fyn SH3 domain and engineered to bind to target sequences and molecules with equal affinity and equal

specificity as an antibody.

- (60) Kunitz domain peptides of the disclosure comprise a protein scaffold comprising a Kunitz domain. Kunitz domains comprise an active site for inhibiting protease activity. Structurally, Kunitz domains of the disclosure comprise a disulfide-rich alpha+beta fold. This structure is exemplified by the bovine pancreatic trypsin inhibitor. Kunitz domain peptides recognize specific protein structures and serve as competitive protease inhibitors. Kunitz domains of the disclosure may comprise Ecallantide (derived from a human lipoprotein-associated coagulation inhibitor (LACI)).
- (61) Monobodies of the disclosure are small proteins (comprising about 94 amino acids and having a mass of about 10 kDa) comparable in size to a single chain antibody. These genetically engineered proteins specifically bind target sequences including antigens. Monobodies of the disclosure may specifically target one or more distinct proteins or target sequences. In preferred embodiments, monobodies of the disclosure comprise a protein scaffold mimicking the structure of human fibronectin, and more preferably, mimicking the structure of the tenth extracellular type III domain of fibronectin. The tenth extracellular type III domain of fibronectin, as well as a monobody mimetic thereof, contains seven beta sheets forming a barrel and three exposed loops on each side corresponding to the three complementarity determining regions (CDRs) of an antibody. In contrast to the structure of the variable domain of an antibody, a monobody lacks any binding site for metal ions as well as a central disulfide bond. Multispecific monobodies may be optimized by modifying the loops BC and FG. Monobodies of the disclosure may comprise an adnectin. (62) VHH
- (63) In certain embodiments of the compositions and methods of the disclosure, a CAR or a CLR comprises a single domain antibody (SdAb). In certain embodiments, the SdAb is a VHH.
- (64) The disclosure provides a CAR or a CLR comprising an antigen or ligand recognition region, respectively, that comprises at least one VHH (to produce a "VCAR" or "VCLR"). CARs and CLRs of the disclosure may comprise more than one VHH. For example, a bi-specific VCAR or VCLR may comprise two VHHs. In some embodiments of the bi-specific VCAR or VCLR, each VHH specifically binds a distinct antigen.
- (65) VHH proteins of the disclosure specifically bind an antigen or a ligand. CARs of the disclosure comprising one or more VHHs that specifically bind an antigen may be used to direct the specificity of a cell, (e.g. a cytotoxic immune cell) towards a target cell expressing the specific antigen. CLRs of the disclosure comprising one or more VHHs that specifically bind an antigen may transduce an intracellular signal upon binding a ligand of either VHH to activate expression of a sequence under the control of an inducible promoter.
- (66) Sequences encoding a VHH of the disclosure can be altered, added and/or deleted to reduce immunogenicity or reduce, enhance or modify binding, affinity, on-rate, off-rate, avidity, specificity, half-life, stability, solubility or any other suitable characteristic, as known in the art.
- (67) Optionally, VHH proteins can be engineered with retention of high affinity for the antigen or ligand and other favorable biological properties. To achieve this goal, the VHH proteins can be optionally prepared by a process of analysis of the parental sequences and various conceptual engineered products using three-dimensional models of the parental and engineered sequences. Three-dimensional models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate sequences and can measure possible immunogenicity (e.g., Immunofilter program of Xencor, Inc. of Monrovia, Calif.). Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate sequence. i.e., the analysis of residues that influence the ability of the candidate VHH protein to bind its antigen/ligand. In this way, residues can be selected and combined from the parent and reference sequences so that the desired characteristic, such as affinity for the target antigen(s)/ligand(s), is achieved. Alternatively, or in addition to, the above procedures, other suitable methods of engineering can be used.
- (68) VH
- (69) In certain embodiments of the compositions and methods of the disclosure, a CAR or a CLR comprises a single domain antibody (SdAb). In certain embodiments, the SdAb is a VH.
- (70) The disclosure provides CARs/CLRs comprising a single domain antibody (to produce a "VCAR" or a "VCLR", respectively). In certain embodiments, the single domain antibody comprises a VH. In certain

embodiments, the VH is isolated or derived from a human sequence. In certain embodiments, VH comprises a human CDR sequence and/or a human framework sequence and a non-human or humanized sequence (e.g. a rat Fc domain). In certain embodiments, the VH is a fully humanized VH. In certain embodiments, the VH s neither a naturally occurring antibody nor a fragment of a naturally occurring antibody. In certain embodiments, the VH is not a fragment of a monoclonal antibody. In certain embodiments, the VH is a UniDab $^{TM}$  antibody (TeneoBio).

- (71) In certain embodiments, the VH is fully engineered using the UniRat™ (TeneoBio) system and "NGS-based Discovery" to produce the VH. Using this method, the specific VH are not naturallyoccurring and are generated using fully engineered systems. The VH are not derived from naturallyoccurring monoclonal antibodies (mAbs) that were either isolated directly from the host (for example, a mouse, rat or human) or directly from a single clone of cells or cell line (hybridoma). These VHs were not subsequently cloned from said cell lines. Instead, VH sequences are fully-engineered using the UniRat™ system as transgenes that comprise human variable regions (VH domains) with a rat Fc domain, and are thus human/rat chimeras without a light chain and are unlike the standard mAb format. The native rat genes are knocked out and the only antibodies expressed in the rat are from transgenes with VH domains linked to a Rat Fc (UniAbs). These are the exclusive Abs expressed in the UniRat. Next generation sequencing (NGS) and bioinformatics are used to identify the full antigen-specific repertoire of the heavychain antibodies generated by UniRat™ after immunization. Then, a unique gene assembly method is used to convert the antibody repertoire sequence information into large collections of fully-human heavy-chain antibodies that can be screened in vitro for a variety of functions. In certain embodiments, fully humanized VH are generated by fusing the human VH domains with human Fcs in vitro (to generate a non-naturally occurring recombinant VH antibody). In certain embodiments, the VH are fully humanized, but they are expressed in vivo as human/rat chimera (human VH, rat Fc) without a light chain. Fully humanized VHs are expressed in vivo as human/rat chimera (human VH, rat Fc) without a light chain are about 80 kDa (vs 150 kDa).
- (72) VCARs/VCLRs of the disclosure may comprise at least one VH of the disclosure. In certain embodiments, the VH of the disclosure may be modified to remove an Fc domain or a portion thereof. In certain embodiments, a framework sequence of the VH of the disclosure may be modified to, for example, improve expression, decrease immunogenicity or to improve function.
- (73) Transposons/Transposases
- (74) Exemplary transposon/transposase systems of the disclosure include, but are not limited to, piggyBac transposons and transposases, Sleeping Beauty transposons and transposases, Helraiser transposons and transposases and Tol2 transposons and transposases.
- (75) The piggyBac transposase recognizes transposon-specific inverted terminal repeat sequences (ITRs) on the ends of the transposon, and moves the contents between the ITRs into TTAA chromosomal sites. The piggyBac transposon system has no payload limit for the genes of interest that can be included between the ITRs. In certain embodiments, and, in particular, those embodiments wherein the transposon is a piggyBac transposon, the transposase is a piggyBac<sup>TM</sup> or a Super piggyBac<sup>TM</sup> (SPB) transposase. In certain embodiments, and, in particular, those embodiments wherein the transposase is a Super piggyBac<sup>TM</sup> (SPB) transposase, the sequence encoding the transposase is an mRNA sequence. (76) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac<sup>TM</sup> (PB) transposase enzyme. The piggyBac (PB) transposase enzyme may comprise or consist of an amino acid sequence at least 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to: (77) TABLE-US-00017 (SEQ ID NO: 14487) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEI SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVT EQPGSSLASN RILTLPORTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121 PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTGATFRD TNEDEIYAFF 101 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFD IRCLRMDDKS IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLT1DEQ LLGFRGRCPF RMYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTOTN GVPLGEYYVK ELSKPVHGSC RNITCDNWFT SIPIAKNLLQ 361 EPYKLTIVGT VRSNKREIPE VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481

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SFIIYSHNVS SKGEKVQSRK KFMRNLYMSLTSSFMRKRLE APTLKRYLRD NISNILPNEV
541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC
(78) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac<sup>TM</sup>
(PB) transposase enzyme that comprises or consists of an amino acid sequence having an amino acid
substitution at one or more of positions 30, 165, 282, or 538 of the sequence:
(79) TABLE-US-00018 (SEQ ID NO: 14487) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEI
SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVI EQPGSSLASN RILTLPQRTI
RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121 PTRMCRNIYD PLLCFKLFFT
DEIISEIVKW TNAEISLKRR ESMTGATFRD TNEDEIYAFF 181 GILVMTAVRK
DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS IRPTLRENDV 241
FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGFRGRCPF RMYIPNKPSK YGIKILMMCD
301 SGTKYMINGM PYLGRGTQTN GVPLGEYYVK ELSKPVHGSC RNITCDNWFT
SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE VLKNSRSRPV GTSMFCFDGP
LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST GKPQMVMYYN QTKGGVDTLD
QMCSVMTCSR KTNRWPMALL YGMINIACIN 481 SFIIYSHNVS SKGEKVQSRK
KFMRNLYMSL TSSFMRKRLE APTLKRYLRD NISNILPNEV 541 PGTSDDSTEE
PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC OSCF.
(80) In certain embodiments, the transposase enzyme is a piggyBac<sup>TM</sup> (PB) transposase enzyme that
comprises or consists of an amino acid sequence having an amino acid substitution at two or more of
positions 30, 165, 282, or 538 of the sequence of SEO ID NO: 14487. In certain embodiments, the
transposase enzyme is a piggyBac<sup>TM</sup> (PB) transposase enzyme that comprises or consists of an amino acid
sequence having an amino acid substitution at three or more of positions 30, 165, 282, or 538 of the
sequence of SEQ ID NO: 14487. In certain embodiments, the transposase enzyme is a piggyBac™ (PB)
transposase enzyme that comprises or consists of an amino acid sequence having an amino acid
substitution at each of the following positions 30, 165, 282, and 538 of the sequence of SEQ ID NO:
14487. In certain embodiments, the amino acid substitution at position 30 of the sequence of SEQ ID NO:
14487 is a substitution of a valine (V) for an isoleucine (I). In certain embodiments, the amino acid
substitution at position 165 of the sequence of SEQ ID NO: 14487 is a substitution of a serine (S) for a
glycine (G). In certain embodiments, the amino acid substitution at position 282 of the sequence of SEQ
ID NO: 14487 is a substitution of a valine (V) for a methionine (M). In certain embodiments, the amino
acid substitution at position 538 of the sequence of SEQ ID NO: 14487 is a substitution of a lysine (K) for
an asparagine (N).
(81) In certain embodiments of the methods of the disclosure, the transposase enzyme is a Super
piggyBac<sup>™</sup> (SPB) transposase enzyme. In certain embodiments, the Super piggyBac<sup>™</sup> (SPB) transposase
enzymes of the disclosure may comprise or consist of the amino acid sequence of the sequence of SEQ ID
NO: 14487 wherein the amino acid substitution at position 30 is a substitution of a valine (V) for an
isoleucine (1), the amino acid substitution at position 165 is a substitution of a serine (S) for a glycine (G),
the amino acid substitution at position 282 is a substitution of a valine (V) for a methionine (M), and the
amino acid substitution at position 538 is a substitution of a lysine (K) for an asparagine (N). In certain
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embodiments, the Super piggyBac™ (SPB) transposase enzyme may comprise or consist of an amino acid

PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTSATFRD TNEDEIYAFF

sequence at least 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to: (82) TABLE-US-00019 (SEQ ID NO: 14484) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEV SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVI EQPGSSLASN RILTLPQRTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121

181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS

RVYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTQTN GVPLGEYYVK ELSKPVHGSC RNITCDNWFT SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST

GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481

541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC QSCF.

SKGEKVQSRK KFMRNLYMSLTSSFMRKRLE APTLKRYLRD NISNILPKEV

IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGFRGRCPF

SFIIYSHNVS

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(83) In certain embodiments of the methods of the disclosure, including those embodiments wherein the
transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac™
or Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more
of positions 3, 46, 82, 103, 119, 125, 177, 180, 185, 187, 200, 207, 209, 226, 235, 240, 241, 243, 258, 296,
298, 311, 315, 319, 327, 328, 340, 421, 436, 456, 470, 486, 503, 552, 570 and 591 of the sequence of SEQ
ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, including those embodiments wherein the
transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac™
or Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more
of positions 46, 119, 125, 177, 180, 185, 187, 200, 207, 209, 226, 235, 240, 241, 243, 296, 298, 311, 315,
319, 327, 328, 340, 421, 436, 456, 470, 485, 503, 552 and 570. In certain embodiments, the amino acid
substitution at position 3 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an asparagine
(N) for a serine (S). In certain embodiments, the amino acid substitution at position 46 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a serine (S) for an alanine (A). In certain embodiments,
the amino acid substitution at position 46 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of
a threonine (T) for an alanine (A). In certain embodiments, the amino acid substitution at position 82 of
SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a tryptophan (W) for an isoleucine (I). In
certain embodiments, the amino acid substitution at position 103 of SEQ ID NO: 14487 or SEQ ID NO:
14484 is a substitution of a proline (P) for a serine (S). In certain embodiments, the amino acid
substitution at position 119 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P)
for an arginine (R). In certain embodiments, the amino acid substitution at position 125 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of an alanine (A) a cysteine (C). In certain embodiments, the
amino acid substitution at position 125 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
leucine (L) for a cysteine (C). In certain embodiments, the amino acid substitution at position 177 of SEQ
ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a tyrosine (Y). In certain
embodiments, the amino acid substitution at position 177 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a histidine (H) for a tyrosine (Y). In certain embodiments, the amino acid substitution at
position 180 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a
phenylalanine (F). In certain embodiments, the amino acid substitution at position 180 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a phenylalanine (F). In certain
embodiments, the amino acid substitution at position 180 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a valine (V) for a phenylalanine (F). In certain embodiments, the amino acid substitution
at position 185 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a
methionine (M). In certain embodiments, the amino acid substitution at position 187 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a glycine (G) for an alanine (A). In certain embodiments,
the amino acid substitution at position 200 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution
of a tryptophan (W) for a phenylalanine (F). In certain embodiments, the amino acid substitution at
position 207 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P) for a valine
(V). In certain embodiments, the amino acid substitution at position 209 of SEQ ID NO: 14487 or SEQ ID
NO: 14484 is a substitution of a phenylalanine (F) for a valine (V). In certain embodiments, the amino
acid substitution at position 226 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
phenylalanine (F) for a methionine (M). In certain embodiments, the amino acid substitution at position
235 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an arginine (R) for a leucine (L). In
certain embodiments, the amino acid substitution at position 240 of SEQ ID NO: 14487 or SEQ ID NO:
14484 is a substitution of a lysine (K) for a valine (V). In certain embodiments, the amino acid substitution
at position 241 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a
phenylalanine (F). In certain embodiments, the amino acid substitution at position 243 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a proline (P). In certain embodiments, the
amino acid substitution at position 258 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
serine (S) for an asparagine (N). In certain embodiments, the amino acid substitution at position 296 of
SEO ID NO: 14487 or SEO ID NO: 14484 is a substitution of a tryptophan (W) for a leucine (L). In
certain embodiments, the amino acid substitution at position 296 of SEQ ID NO: 14487 or SEQ ID NO:
14484 is a substitution of a tyrosine (Y) for a leucine (L). In certain embodiments, the amino acid
substitution at position 296 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
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phenylalanine (F) for a leucine (I.) In certain embodiments, the amino acid substitution at position 298 of SEQ ID NO 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a methionine (M). In certain embodiments, the amino acid substitution at position 298 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an alanine (A) for a methionine (M). In certain embodiments, the amino acid substitution at position 298 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a valine (V) for a methionine (M). In certain embodiments, the amino acid substitution at position 311 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a proline (P) In certain embodiments, the amino acid substitution at position 311 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a valine for a proline (P). In certain embodiments, the amino acid substitution at position 315 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for an arginine (R). In certain embodiments, the amino acid substitution at position 319 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a glycine (G) for a threonine (T). In certain embodiments, the amino acid substitution at position 327 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an arginine (R) for a tyrosine (Y). In certain embodiments, the amino acid substitution at position 328 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a valine (V) for a tyrosine (Y). In certain embodiments, the amino acid substitution at position 340 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a glycine (G) for a cysteine (C). In certain embodiments, the amino acid substitution at position 340 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a cysteine (C). In certain embodiments, the amino acid substitution at position 421 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a histidine (H) for the aspartic acid (D). In certain embodiments, the amino acid substitution at position 436 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a valine (V). In certain embodiments, the amino acid substitution at position 456 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a tyrosine (Y) for a methionine (M). In certain embodiments, the amino acid substitution at position 470 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a phenylalanine (F) for a leucine (L). In certain embodiments, the amino acid substitution at position 485 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a serine (S). In certain embodiments, the amino acid substitution at position 503 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a methionine (M). In certain embodiments, the amino acid substitution at position 503 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a methionine (M). In certain embodiments, the amino acid substitution at position 552 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a valine (V). In certain embodiments, the amino acid substitution at position 570 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a threonine (T) for an alanine (A). In certain embodiments, the amino acid substitution at position 591 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P) for a glutamine (Q). In certain embodiments, the amino acid substitution at position 591 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an arginine (R) for a glutamine (Q). (84) In certain embodiments of the methods of the disclosure, including those embodiments wherein the

transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> transposase enzyme may comprise or the Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more of positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments of the methods of the disclosure, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>TM</sup> transposase enzyme may comprise or the Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at two, three, four, five, six or more of positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> transposase enzyme may comprise or the Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, the amino acid substitution at position 103 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P) for a serine (S). In certain embodiments, the amino acid substitution at position 194 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a valine (V) for a methionine (M). In certain embodiments, the amino acid substitution at position 372 of

SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an alanine (A) for an arginine (R). In certain embodiments, the amino acid substitution at position 375 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an alanine (A) for a lysine (K). In certain embodiments, the amino acid substitution at position 450 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an asparagine (N) for an aspartic acid (D). In certain embodiments, the amino acid substitution at position 509 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a glycine (G) for a serine (S). In certain embodiments, the amino acid substitution at position 570 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a serine (S) for an asparagine (N). In certain embodiments, the piggyBac™ transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487. In certain embodiments, including those embodiments wherein the piggyBac<sup>TM</sup> transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487, the piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at positions 372, 375 and 450 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, the piggyBac<sup>TM</sup> transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487, a substitution of an alanine (A) for an arginine (R) at position 372 of SEQ ID NO: 14487, and a substitution of an alanine (A) for a lysine (K) at position 375 of SEQ ID NO: 14487. In certain embodiments, the piggyBac<sup>TM</sup> transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487, a substitution of an alanine (A) for an arginine (R) at position 372 of SEQ ID NO: 14487, a substitution of an alanine (A) for a lysine (K) at position 375 of SEO ID NO: 14487 and a substitution of an asparagine (N) for an aspartic acid (D) at position 450 of SEQ ID NO: 14487.

- (85) The sleeping beauty transposon is transposed into the target genome by the Sleeping Beauty transposase that recognizes ITRs, and moves the contents between the ITRs into TA chromosomal sites. In various embodiments, SB transposon-mediated gene transfer, or gene transfer using any of a number of similar transposons, may be used in the compositions and methods of the disclosure.
- (86) In certain embodiments, and, in particular, those embodiments wherein the transposon is a Sleeping Beauty transposon, the transposase is a Sleeping Beauty transposase or a hyperactive Sleeping Beauty transposase (SB100X).
- (87) In certain embodiments of the methods of the disclosure, the Sleeping Beauty transposase enzyme comprises an amino acid sequence at least 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (88) TABLE-US-00020 (SEQ ID NO: 14485) 1 MGKSKEISQD LRKKIVDLHK SGSSLGAISK RLKVPRSSVQ TIVRXYKHHG TTQPSYRSGR 61 RRVLSPRDER TLVRKVQINP RTTAKDLVKM LEETGTKVSI STVKRVLYRH NLKGRSARKK 121 PLLQNRHKKA RLRFATAHGD KDRTFWRNVL WSDETKIELF GHNDHRYVWR KKGEACKPKN 181 TIPTVKHGGG SIMLWGCFAA GGTGALHKID GIMRKENYVD ILKQHLKTSV RKLKLGRKWV 241 FQMDNDPKHT SKVVAKWLKD NKVKVLEWPS QSPDLNPIEN LWAELKKRVR ARRPTNLTQL 301 HQLCQEEWAK IHPTYCGKLV EGYPKRLTQV KQFKGNATKY.
- (89) In certain embodiments of the methods of the disclosure, the hyperactive Sleeping Beauty (SB100X) transposase enzyme comprises an amino acid sequence at least 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (90) TABLE-US-00021 (SEQ ID NO: 14486) 1 KGKSKEISQD LRKRIVDLHK SGSSLGAISK RLAVPRSSVQ TIVRKYKHHG TTQPSYRSGR 61 RRVLSPRDER TLVRKVQINP RTTAKDLVKM LEETGTKVSI STVKRVLYRH NLKGHSARKK 121 PLLQNRHKKA RLRFATAHGD KDRTFWRNVL WSDETKIELF GHNDHRYVWR KKGEACKPKN 181 TIPTVKHGGG SIMLWGCFAA GGTGALHKID GIMDAVQYVD ILKQHLKTSV RKLKLGRKWV 241 FQHDNDPKHT SKVVAKWLKD NKVKVLEWPS QSPDLNPIEN LWAELKKRVR ARRPTNLTQL 301 HQLCQEEWAK IHPNYCGKLV EGYPKRLTQV KQFKGNATKY.
- (91) The Helraiser transposon is transposed by the Helitron transposase. Helitron transposases mobilize the Helraiser transposon, an ancient element from the bat genome that was active about 30 to 36 million years ago. An exemplary Helraiser transposon of the disclosure includes Helibat1, which comprises a

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nucleic acid sequence comprising:
(92) TABLE-US-00022 (SEQ ID NO: 14652)
                                   1 TCCTATATAA TAAAAGAGAA
ACATGCAAAT TGACCATCCC TCCGCTACGC
                                  TCAAGCCACG
                                               61 CCCACCAGCC
           GACTATGCAA ATTAACCCAA CAAAGATGGC AGTTAAATTT
AATCAGAAGT
           GTGTCAAGCG CCCCAGGAGG CAACGGCGGC CGCGGGCTCC
GCATACGCAG
CAGGACCTTG
           181 GCTGGCCCCG GGAGGCGAGG CCGGCCGCGC CTAGCCACAC
CCGCGGGCTC
           CCGGGACCTT 241 CGCCAGCAGA GAGCAGAGCG GGAGAGCGGG
CGGAGAGCGG GAGGTTTGGA GGACTTGGCA
                                   301 GAGCAGGAGG CCGCTGGACA
TAGAGCAGAG CGAGAGAGAG GGTGGCTTGG AGGGCGTGGC 361 TCCCTCTGTC
ACCCCAGCTT
           CCTCATCACA GCTGTGaAAA CTGACAGCAG GGAGGAGGAA
GTCCCACCCC
           CACAGAATCA GCCAGAATCA GCCGTTGGTC AGACAGCTCT
           481 ACAGCCAGGA CTCTCATTCA CCTGCATCTC AGACCGTGAC
CAGCGGCCTG
           TGGGACTATG 541 TCTAAAGAAC AACTGTTGAT ACAACGTAGC
AGTAGAGAGG
TCTGCAGCCG AAAGATGCCG GCGTTATCGA 601 CAGAAAATGT CTGCAGAGCA
           GATCTTGAAA GAAGGCGGCG CCTGCAACAG
ACGTGCGTCT
                                               661 AATGTATCTG
           ACTGGAAAAA CGTCGCTCTG AAGCCGAAAA ACAGCGGCGT
AAGAGCAGCT
CATCGACAGA AAATGTCTAA AGACCAACGT GCCTTTGAAG TTGAAAGAAG
GCGGTGGCGA 781 CGACAGAATA TGTCTAGAGA ACAGTCATCA ACAAGTACTA
CCAATACCGG TAGGAACTGC 641 CTTCTCAGCA AAAATGGAGT ACATGAGGAT
GCAATTCTCG AACATAGTTG TGGTGGAATG 901 ACTGTTCGAT GTGAATTTTG
CCTATCACTA AATTTCTCTG ATGAAAAACC ATCCGATGGG
                                             961 AAATTTACTC
GATGTTGTAG CAAAGGGAAA GTCTGTCCAA ATGATATACA TTTTCCAGAT 1021
TACCCGGCAT ATTTAAAAAG ATTAATGACA AACGAAGATT CTGACAGTAA
AAATTTCATG 1081 GAAAATATTC GTTCCATAAA TAGTTCTTTT GCTTTTGCTT
CCATGGGTGC AAATATTGCA 1141 TCGCCATCAG GATATGGGCC ATACTGTTTT
AGAATACACG GACAAGTTTA TCACCGTACT 1201 GGAACTTTAC ATCCTTCGGA
TGGTGTTTCT CGGAAGTTTG CTCAACTCTA TATTTTGGAT 1261 ACAGCCGAAG
CTACAAGTAA AAGATTAGCA ATGCCAGAAA ACCAGGGCTG CTCAGAAAGA 1321
CTCATGATCA ACATCAACAA CCTCATGCAT GAAATAAATG AATTAAGAAA
ATCGTACAAG 1381 ATGCTACATG AGGTAGAAAA GGAAGCCCAA TCTGAAGCAG
CAGCAAAAGG TATTGCTCCC 1441 ACAGAAGTAA CAATGGCGAT TAAATACGAT
CGTAACAGTG ACCCAGGTAG ATATAATTCT 1501 CCCCGTGTAA CCGAGGTTGC
TGTCATATTC AGAAACGAAG ATGGAGAACC TCCTTTTGAA 1561 AGGGACTTGC
TCATTCATTG TAAACCAGAT CCCAATAATC CAAATGCCAC TAAAATGAAA 1621
CAAATCAGTA TCCTGTTTCC TACATTAGAT GCAATGACAT ATCCTATTCT
TTTTCCACAT 1681 GGTGAAAAAG GCTGGGGAAC AGATATTGCA TTAAGACTCA
GAGACAACAG TGTAATCGAC 1741 AATAATACTA GACAAAATGT AAGGACACGA
GTCACACAAA TGCAGTATTA TGGATTTCAT 1601 CTCTCTGTGC GGGACACGTT
GAATCCTATT TTAAATGCAG GAAAATTAAC TCAACAGTTT 1861 ATTGTGGATT
CATATTCAAA AATCGAGGCC AATCGGATAA ATTTCATCAA AGCAAACCAA 1921
TCTAAGTTGA GAGTTGAAAA ATATAGTGGT TTGATGGATT ATCTCAAATC
TAGATCTGAA 1981 AATGACAATG TGCCGATTGG TAAAATGATA ATACTTCCAT
CATCTTTTGA GGGTAGTCCC 2041 AGAAATATGC AGCAGCGATA
                                                TCAGGATGCT
ATGGCAATTG TAACGAAGTA TGGCAAGCCC 2101 GATTTATTCA TAACCATGAC
ATGCAACCCC AAATGGGCAG ATATTACAAA CAATTTACAA 2161 CGCTGGCAAA
AAGTTGAAAA CAGACCTGAC TTGGTAGCCA GAGTTTTTAA TATTAAGCTG 2221
AATGCTCTTT TAAATGATAT ATGTAAATTC CATTTATTTG GGAAAGTAAT
AGCTAAAATT 2281 CATGTCATTG AATTTCAGAA ACGCGGACTG CCTCACGCTC
         GATATTAGAT 2341 AGTGAGTCCA AATTACGTTC AGAAGATGAC
ACATATTATT
          TAGTTAAGGC AGAAATTCCA 2401 GATGAAGACC AGTGTCCTCG
ATTGACCGTA
ACTTTTTCAA ATTGTAAAAT CAAATATGGT ACATGGACCA 2461 TGTGGAATAC
           TAGTCCATGT ATGGAAAATG GAAAATGTTC AAAGGGATAT 2521
AAAATCGAAA
          TTCAAAATGC GACCA1TGGA AATATTGATG GATATCCGAA
CCAAAAGAAT
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ATACAAACGA 2581 AGATCTGGTA GCACCATGTC TATTGGAAAT AAAGTTGTCG ATAACACTTG GATTGTCCCT 2641 TATAACCCGT ATTTGTGCCT TAAATATAAC TGTCATATAA ATGTTGAAGT CTGTGCATCA 2701 ATTAAAAGTG TCAAATATTT ATTTAAATAC ATCTATAAAG GGCACGATTG TGCAAATATT 2761 CAAATTTCTG AAAAAAATAT TATCAATCAT GACGAAGTAC AGGACTTCAT TGACTCCAGG 2821 TATGTGAGCG CTCCTGAGGC TGTTTGGAGA CTTTTTGCAA TGCGAATGCA TGACCAATCT 2881 CATGCAATCA CAAGATTAGC TATTCATTTG CCAAATGATC AGAATTTGTA TTTTCATACC 2941 GATGATTTTG CTGAAGTTTT AGATAGGGCT AAAAGGCATA ACTCGACTTT GATGGCTTGG 3001 TTCTTATTGA ATAGAGAAGA TTCTGATGCA CGTAATTATT ATTATTGGGA GATTCCACAG 3061 CATTATGTCT TTAATAATTC TTTGTGGACA AAACGCCGAA AGGGTGGGAA TAAAGTATTA 3121 GGTAGACTGT TCACTGTGAG CTTTAGAGAA CCAGAACGAT ATTAGCTTAG ACTTTTGCTT 3181 CTGCATGTAA AAGGTGCGAT AAGTTTTGAG GATCTGCGAA CTGTAGGAGG TGTAACTTAT 3241 GATACATTTC ATGAAGCTGC TAAACACCGA GGATTATTAC TTGATGACAC TATCTGGAAA 3301 GATACGATTG ACGATGCAAT CATCCTTAAT ATGCCCAAAC AACTACGGCA ACTTTTTGCA 3361 TATATATGTG TGTTTGGATG TCCTTCTGCT GCAGACAAAT TATGGGATGA GAATAAATCT 3421 CATTTTATTG TTGATTTCTG TTGGAAATTA CACCGAAGAG AAGGTGCCTG TGTGAACTGT 3481 GAAATGCATG CCCTTAACGA AATTCAGGAG GTATTCACAT TGCATGGAAT GAAATGTTCA 3541 CATTTCAAAC TTCCGGACTA TCCTTTATTA ATGAATGCAA ATACATGTGA TCAATTGTAC 3601 GAGCAACAAC AGGCAGAGGT TTTGATAAAT TCTCTGAATG ATGAACAGTT GGCAGCCTTT 3661 CAGACTATAA CTTCAGCCAT CGAAGATCAA ACTGTACACC CCAAATGCTT TTTCTTGGAT 3721 GGTCCAGGTG GTAGTGGAAA AACATATCTG TATAAAGTTT TAACACATTA TATTAGAGGT 3781 CGTGGTGGTA CTGTTTTACC CACAGCATCT ACAGGAATTG CTGCAAATTT ACTTCTTGGT 3841 GGAAGAACCT TTGATTCCCA ATATAAATTA CCAATTCCAT TAAATGAAAC TTCAATTTCT 3901 AGACTCGATA TAAAGAGTGA AGTTGCTAAA ACCATTAAAA AGGCCCAACT TCTCATTATT 3961 GATGAATGCA CCATGGCATC CAGTCATGCT ATAAACGCCA TAGATAGATT ACTAAGAGAA 4021 ATTATGAATT TGAATGTTGC ATTTGGTGGG AAAGTTCTCC TTCTCGGAGG GGATTTTCGA 4081 CAATGTCTCA GTATTGTACC ACATGCTATG CGATCGGCCA TAGTACAAAC GAGTTTAAAG 4141 TACTGTAATG TTTGGGGGATG TTTCAGAAAG TTGTCTCTTA AAACAAATAT GAGATCAGAG 4201 GATTCTGCTT ATAGTGAATG GTTAGTAAAA CTTGGAGATG GCAAACTTGA TAGCAGTTTT 4261 CATTTAGGAA TGGATATTAT TGAAATCCCC CATGAAATGA TTTGTAACCC ATCTATTATT 4321 GAAGCTACCT TTGGAAATAG TATATCTATA GATAATATTA AAAATATATC TAAACGTGCA 4381 ATTCTTTGTC CAAAAAATGA GCATGTTCAA AAATTAAATG AAGAAATTTT GGATATACTT 4441 GATGGAGATT TTCACACATA TTTGAGTGAT GATTCCATTG ATTCAACAGA TGATGCTGAA 4501 AAGGAAAATT TTCCCATCGA ATTTCTTAAT AGTATTACTC CTTCGGGAAT GCCGTGTCAT 4561 AAATTAAAAT TGAAAGTGGG TGCAATCATC ATGCTATTGA GAAATCTTAA TAGTAAATGG 4621 GGTCTTTGTA ATGGTACTAG ATTTATTATC AAAAGATTAC GACCTAACAT TATCGAAGCT 4681 GAAGTATTAA CAGGATCTGC AGAGGGAGAG GTTGTTCTGA TTCCAAGAAT TGATTTGTCC 4741 CCATCTGACA CTGGCCTCCC ATTTAAATTA ATTCGAAGAC AGTTTCCCGT GATGCCAGCA 4801 TTTGCGATGA CTATTAATAA ATCACAAGGA CAAACTCTAG ACAGAGTAGG AATATTCCTA 4861 CCTGAACCCG TTTTCGCACA TGGTCAGTTA TATGTTGCTT TCTCTCGAGT TCGAAGAGCA 4921 TGTGACGTTA AAGTTAAAGT TGTAAATACT TCATCACAAG GGAAATTAGT CAAGCACTCT 4981 GAAAGTGTTT TTACTCTTAA TGTGGTATAC AGGGAGATAT TAGAATAAGT TTAATCACTT 5041 TATCAGTCAT TGTTTGCATC AATGTTGTTT TTATATCATG TTTTTGTTGT TTTTATATCA 5101 TGTCTTTGTT GTTGTTATAT CATGTTGTTA TTGTTTATTT ATTAATAAAT TTATGTATTA 5161 TTTTCATATA CATTTTACTC ATTTCCTTTC ATCTCTCACA CTTCTATTAT AGAGAAAGGG 5221

CAAATAGCAA TATTAAAATA TTTCCTCTAA TTAATTCCCT TTCAATGTGC ACGAATTTCG 5281 TGCACCGGGC CACTAG.

- (93) Unlike other transposases the Helitron transposase does not contain an RNase-H like catalytic domain, but instead comprises a RepHel motif made up of a replication initiator domain (Rep) and a DNA helicase domain. The Rep domain is a nuclease domain of the HUH superfamily of nucleases. (94) An exemplary Helitron transposase of the disclosure comprises an amino acid sequence comprising: (95) TABLE-US-00023 (SEQ ID NO: 14501) 1 MSKEQLLXQR SSAAERCRRY RQKMSAEQRA SDLERRRRLQ QKVSEEQLLE KRRSEAEKQR 61 RHRQKMSKDQ RAFEVERRRW RRQNMSREQS STSTTNTGRN CLLSKNGVHE DAILEHSCGG 121 MTVRCEFCLS LNFSDEKPSD GKFTRCCSKG KVCPNDIHFP DYPAYLKRLM TNEDSDSKNF 181 MENIRSINSS FAFASMGANI ASPSGYGPYC FRIHGQVYHR TGTLHPSDGV SRKFAQLYIL 241 DTAEATSKRL AMPENQGCSE RLMININNLM HEINELTKSY KMLHEVEKEA QSEAAAKGIA 301 PTEVTMAIKY DRNSDPGRYN SPRVTEVAVI FRNEDGEPPF ERDLLIHCKP DPNNPNATKM 361 KQISILFPTL DAMTYPILFP HGEKGWGTDI ALRLRDNSVI DKNTRQMVRT RVTQMQYYGF 421 HLSVRDTFNP ILNAGKLTQQ FIVDSYSKME ANRINFIKAN QSKLRVEKYS GLMDYLKSRS 481 ENDNVPIGKM IILPSSFEGS PRNMQQRYQD AMAIVTKYSK PDLFITMTCN PKWADITNNL 541 QRWQKVENRP DLVARVFNIK LNALLNDICK FHLFGKVIAK IHVIEFQKRG LPHAHILLIL 601 DSESKLR8ED DIDRIYKAEI PDEDQCPRLF QIVKSMMVHG PCGIQNPNSP CMENGKCSKG 661 YPKEFQNATI GNIDGYPKYK RRSGSTMSIG NKVVDNTWIV PYNPYLCLKY NCHINVEVCA 721 SIKSVKYLFK YIYKGHDCAN IQISEKNIIN HDEVQDFIDS RYVSAPEAVW RLFAMRMHDQ 781 SHAITRLAIH LPMDQMLYFH TDDFAEVLDR AKRHNSTLMA WFLLNREDSD ARNYYYWEIP 841 QHYVFNNSLW TKRRKGGMKV LGRLFTVSFR EPERYYLRLL LLHVKGAISF EDLRTVGGVT 901 YDTFHEAAKH RGLLLDDTIW KDTIDDAIIL NMPKQLRQLF AYICVFGCPS AADKLWDENK 561 SHFIEDFCWK LHRREGACVN CEMHALNEIQ EVFTLHGMKC SHFKLPDYPL LMNANTCDQL 1021 YEQQQAEVLI NSLMDEQLAA FQTITSAIED QTVHPKCFFL DGPGGSGKTY LYKVLTHYIR 1081 GRGGTVLPTA STGIAANLLL GGRTFHSQYK LPIPLNETSI SRLDIKSEVA KTIKKAQLLI 1141 IDECTMASSH AINAIDRLLR EXMNLNVAFG GKVLLLGGDF RQCLSIVPHA MRSAIVQTSL 1201 KYCNVWGCFR KLSLKTNMRS EDSAYSEWLV KLGDGKLDSS FHLGMDIIEI PHEMICNGSI 1261 IEATFGNSIS IDNIKNISKR AILCPKNEHV QKLNEEILDI LDGDFHTYLS DDSIDSTDDA 1321 EKENFPIEFL NSITPSGMPC HKLKLKVGAI IMLLRNLNSK WGLCNGTRET **IKRLRPNIIE** 1381 AEVLTGSAEG EVVLIPRIDL SPSDTGLPFK LIRROFPVMP AFAMTIMKSO GQTLDRVGIF 1441 LPEPVFAHGQ LYVAFSRVRR ACDVKVKVVN TSSQGKLVKH
- (96) In Helitron transpositions, a hairpin close to the 3' end of the transposon functions as a terminator. However, this hairpin can be bypassed by the transposase, resulting in the transduction of flanking sequences. In addition, Helraiser transposition generates covalently closed circular intermediates. Furthermore, Helitron transpositions can lack target site duplications. In the Helraiser sequence, the transposase is flanked by left and right terminal sequences termed LTS and RTS. These sequences terminate with a conserved 5'-TC/CTAG-3' motif. A 19 bp palindromic sequence with the potential to form the hairpin termination structure is located 11 nucleotides upstream of the RTS and consists of the sequence

SESVFTLNVV YREILE.

- (97) TABLE-US-00024 (SEQ ID NO: 14500) GTGCACGAATTTCGTGCACCGGGCCACTAG. (98) Tol2 transposons may be isolated or derived from the genome of the medaka fish, and may be similar to transposons of the hAT family. Exemplary Tol2 transposons of the disclosure are encoded by a sequence comprising about 4.7 kilobases and contain a gene encoding the Tol2 transposase, which contains four exons. An exemplary Tol2 transposase of the disclosure comprises an amino acid sequence comprising the following:
- (99) TABLE-US-00025 (SEQ ID NO: 14502) 1 MEEVCDSSAA ASSTVQNQPQ DQEHPWPYLR EFFSLSGVNK DSFKMKCVLC LDLNKEISAF 61 KSSPSNLRKH

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IERMHPNYLK NYSKLTAQKR KIGTSTHASS SKQLKVDSVF PVKHVSPVTV 121
NKAILRYIIQ GLHPFSTVDL PSFKELISTL QPGISVITRP TLRSKIAEAA LIMKQKVTAA
181 MSEVEWIATT TDCWTARRKS FIGVTAHWIN PGSLERHSAA LACKRLMGSH
TFEVLASAMN 241 DIHSEYEIRD KVVCTTTDSG SNFMKAFRVF GVENNDIETE
ARRCESDDTD SEGCGEGSDG 301 VEFQDASRVL DQDDGFEFQL PKHQKCACHL
LNLVSSVDAQ KALSNEHYKK LYRSVFGKCQ 361 ALWNKSSRSA LAAEAVESES
RLQLLRPNQT RWNSTFMAVD RILQICKEAG EGALRNICTS 421 LEVPMFNPAE
MLFLTEWANT MRPVAKVLDI LQAETNTQLG WLLPSVHQLS LKLQRLHHSL 481
RYCDPLVDAL QQGIQTRFKH MFEDPEIIAA AILLPKFRTS WTNDETIIKR
GMDYIRVHLE 541 PLDHKKELAN SSSDDEDFFA SLKPTTHEAS KELDGYLACV
SDTRESLLTF PAICSLSIKT 601 NTPLTASAAC ERLFSTAGLL FSPKRARLDT
NNFENQLLLK LNLREYNFE
(100) An exemplary Tol2 transposon of the disclosure, including inverted repeats, subterminal sequences
and the Tol2 transposase, is encoded by a nucleic acid sequence comprising the following:
(101) TABLE-US-00026 (SEQ ID NO: 17041)
                                      1 CAGAGGTGTA AAGTACTTGA
GTAATTTTAC TTGATTACTG TACTTAAGTA TTATTTTTGG
                                                61 GGATTTTTAC
TTTACTTGAG TACAATTAAA AATCAATACT TTTACTTTTA CTTAATTACA
                                                          121
TTTTTTTAGA AAAAAAAGTA CTTTTTACTC CTTACAATTT TATTTACAGT
CAAAAAGTAC 181 TTATTTTTG GAGATCACTT CATTCTATTT TCCCTTGCTA
TTACCAAACC AATTGAATTG
                        241 CGCTGATGCC CAGTTTAATT TAAATGTTAT
TTATTCTGCC TATGAAAATC GTTTTCACAT 301 TATATGAAAT TGGTCAGACA
TGTTCATTGG TCCTTTGGAA GTGACGTCAT GTCACATCTA 361 TTACCACAAT
GCACAGCACC TTGACCTGGA AATTAGGGAA ATTATAACAG TCAATCAGTG
                                                            421
GAAGAAAATG GAGGAAGTAT GTGATTCATC AGCAGCTGCG AGCAGCACAG
TCCAAAATCA 481 GCCACAGGAT CAAGAGCACC CGTGGCCGTA TCTTCGCGAA
TTCTTTTCTT TAAGTGGTGT 541 AAATAAAGAT TCATTCAAGA TGAAATGTGT
CCTCTGTCTC CCGCTTAATA AAGAAATATC 601 GGCCTTCAAA AGTTCGCCAT
CAAACCTAAG GAAGCATATT GAGGTAAGTA CATTAAGTAT
                                                661 TTTGTTTTAC
TGATAGTTTT TTTTTTTTT TTTTTTTTT TTTTTGGGTG TGCATGTTTT 721
GACGTTGATG GCGCGCCTTT TATATGTGTA GTAGGCCTAT TTTCACTAAT
GCATGCGATT 781 GACAATATAA GGCTCACGTA ATAAAATGCT AAAATGCATT
TGTAATTGGT AACGTTAGGT
                        841 CCACGGGAAA TTTGGCGCCT ATTGCAGCTT
TGAATAATCA TTATCATTCC GTGCTCTCAT 901 TGTGTTTCAA TTCATGCAAA
ACACAAGAAA ACCAAGCGAG AAATTTTTTT CCAAACATGT
                                                 961 TGTATTGTCA
AAACGGTAAC ACTTTACAAT GAGGTTGATT AGTTCATGTA TTAACTAACA 1021
TTAAATAACC ATGAGCAATA CATTTGTTAC TGTATCTGTT AATCTTTGTT
AACGTTAGTT 1081 AATAGAAATA CAGATGTTCA TTGTTTGTTC ATGTTAGTTC
ACAGTGCATT AACTAATGTT 1141 AACAAGATAT AAAGTATTAG TAAATGTTGA
AATTAACATG TATACGTGCA GTTCATTATT 1201 AGTTCATGTT AACTAATGTA
GTTAACTAAC GAACCTTATT GTAAAAGTGT TACCATCAAA 1261 ACTAATGTAA
TGAAATCAAT TCACCCTGTC ATGTCAGCCT TACAGTCCTG TGTTTTTGTC 1321
AATATAATCA GAAATAAAAT TAATGTTTGA TTGTCACTAA ATGCTACTGT
ATTTCTAAAA 1381 TCAACAAGTA TTTAACATTA TAAAGTGTGC AATTGGCTGC
AAATGTCAGT TTTATTAAAG 1141 GGTTAGTTCA CCCAAAAATG AAAATAATGT
CATTAATGAC TCGCCCTCAT GTCGTTCCAA 1501 GCCCGTAAGA CCTCCGTTCA
TCTTCAGAAC ACAGTTTAAG ATATTTTAGA TTTAGTCCGA 1561 GAGCTTTCTG
TGCCTCCATT GAGAATGTAT GTACGGTATA CTGTCCATGT CCAGAAAGGT 1621
AATAAAAACA TCAAAGTAGT CCATGTGACA TCAGTGGGTT AGTTAGAATT
TTTTGAAGCA 1681 TCGAATACAT TTTGGTCCAA AAATAACAAA ACCTACGACT
TTATTCGGCA TTGTATTCTC 1741 TTCCGGGTCT GTTGTCAATC CGCGTTCACG
ACTTCGCAGT GACGCTACAA TGCTGAATAA 1801 AGTCGTAGGT TTTGTTATTT
TTGGACCAAA ATGTATTTTC GATGCTTCAA ATAATTCTAC 1861 CTAACCCACT
GATGTCAGAT GGACTACTTT GATGTTTTTA TTACCTTTCT GGACATGGAC 1921
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AGTATACCGT ACATACATTT TCAGTGGAGG GACAGAAAGC TCTCGGACTA
AATCTAAAAT 1981 ATCTTAAACT GTGTTCCGAA GATGAACGGA GGTGTTACGG
GCTTGGAACG ACATGAGGGT 2041 GAGTCATTAA TGACATCTTT TCATTTTTGG
GTGAACTAAC CCTTTAATGC TGTAATCAGA 2101 GACTGTATGT GTAATTGTTA
CATTTATTCC ATACAATATA AATATTTATT TGTTGTTTTT 2161 ACAGAGAATG
CACCCAAATT ACCTCAAAAA CTACTCTAAA TTGACAGCAC AGAAGAGAAA 2221
GATCGGGACC TCCACCCATG CTTCCAGCAG TAAGCAACTG AAAGTTGACT
CAGTTTTCCC 2281 AGTCAAACAT GTGTCTCCAG TCACTGTGAA CAAAGCTATA
TTAAGGTACA TCATTCAAGG 2341 ACTTCATCCT TTCAGCACTG TTGATCTGCC
ATCATTTAAA GAGCTGATTA GTACACTGCA 2401 GCCTGGCATT TCTGTCATTA
CAAGGCCTAC TTTACGCTCC AAGATAGCTG AAGCTGCTCT 2461 GATCATGAAA
CAGAAAGTGA CTGCTGCCAT GAGTGAAGTT GAATGGATTG CAACCACAAC 2521
GGATTGTTGG ACTGCACGTA GAAAGTCATT CATTGGTGTA ACTGCTCACT
GGATCAACCC 2581 TGGAAGTCTT GAAAGACATT CCGCTGCACT TGCCTGCAAA
AGATTAATGG GCTCTCATAC 2641 TTTTGAGGTA CTGGCCAGTG CCATGAATGA
TATCCACTCA GAGTATGAAA TACGTGACAA 2701 GGTTGTTTGC ACAACCACAG
ACAGTGGTTC CAACTTTATG AAGGCTTTCA GAGTTTTTGG 2761 TGTGGAAAAC
AATGATATCG AGACTGAGGC AAGAAGGTGT GAAAGTGATG ACACTGATTC 2821
TGAAGGCTGT GGTGAGGGAA GTGATGGTGT GGAATTCCAA GATGCCTCAC
GAGTCCTGGA 2881 CCAAGACGAT GGCTTCGAAT TCCAGCTACC AAAACATCAA
AAGTGTGCCT GTCACTTACT 2941 TAACCTAGTC TCAAGCGTTG ATGCCCAAAA
AGCTCTCTCA AATGAACACT ACAAGAAACT 3001 CTACAGATCT GTCTTTGGCA
AATGCCAAGC TTTATGGAAT AAAAGCAGCC GATCGGCTCT 3061 AGCAGCTGAA
GCTGTTGAAT CAGAAAGCCG GCTTCAGCTT TTAAGGCCAA ACCAAACGCG 3121
GTGGAATTCA ACTTTTATGG CTGTTGACAG AATTCTTCAA ATTTGCAAAG
AAGCAGGAGA 3181 AGGCGCACTT CGGAATATAT CCACCTCTCT TGAGGTTCCA
ATGTAAGTGT TTTTCCCCTC 3241 TATCGATGTA AACAAATGTG GGTTGTTTTT
GTTTAATACT CTTTGATTAT GCTGATTTCT 3301 CCTGTAGGTT TAATCCAGCA
GAAATGCTCT TCTTGACACA CTCCGCCAAC ACAATCCGTC 3361 CAGTTGCAAA
AGTACTCGAC ATCTTGCAAG CGGAAACGAA TACACAGCTG GGGTGGCTGC 3421
TGCCTAGTGT CCATCAGTTA AGCTTGAAAC TTCAGCGACT CCACCATTCT
CTCAGGTACT 3481 GTGACCCACT TGTGGATGCC CTACAACAAG GAATCCAAAC
ACGATTCAAG CATATGTTTG 3541 AAGATCCTGA GATCATAGCA GCTGCCATCC
TTCTCCCTAA ATTTCGGACC TCTTGGACAA 3601 ATGATGAAAC CATCATAAAA
CGAGGTAAAT GAATGCAAGC AACATACACT TGACGAATTG 3661 TAATCTGGGC
AACCTTTGAG CCATACCAAA ATTATTCTTT TATTTATTTA TTTTTGCACT 3721
TTTTAGGAAT GTTATATCCC ATCTTTGGCT GTGATCTCAA TATGAATATT
GNFGTAAAGT 3781 ATTCTTGCAG CAGGTTGTAG TTATCCCTCA GTGTTTCTTG
AAACCAAACT CATATGTATG 3841 ATATGTGGTT TGGAAATGCA GTTAGATTTT
ATGCTAAAAT AAGGGATTTG CATGATTTTA 3901 GATGTAGATG ACTGCACGTA
AATGTAGTTA ATGACAAAAT CCATAAAATT TGTTCCCAGT 3961 CAGAAGCCCC
TCAACCAAAC TTTTCTTTGT GTCTGCTCAC TGTGCTTGTA GGCATGGACT 4021
ACATCAGAGT GCATCTGGAG CCTTTGGACC ACAAGAAGGA ATTGGCCAAC
AGTTCATCTG 4081 ATGATGAAGA TTTTTTCGCT TCTTTGAAAC CGACAACACA
TGAAGCCAGC AAAGAGTTGG 4141 ATGGATATCT GGCCTGTGTT TCAGACACCA
GGGAGTCTCT GCTCACGTTT CCTGCTATTT 4201 GCAGCCTCTC TATCAAGACT
AATACACCTC TTCCCGCATC GGCTGCCTGT GAGAGGCTTT 4261 TCAGCACTGC
AGGATTGCTT TTCAGCCCCA AAAGAGCTAG GCTTGACACT AACAATTTTG 4321
AGAATCAGCT TCTACTGAAG TTAAATCTGA GGTTTTACAA CTTTGAGTAG
CGTGTACTGG 4381 CATTAGATTG TCTGTCTTAT AGTTTGATAA TTAAATACAA
ACAGTTCTAA AGCAGGATAA 4441 AACCTTGTAT GCATTTCATT TAATGTTTTT
TGAGATTAAA AGCTTAAACA AGAATCTCTA 4501 GTTTTCTTTC TTGCTTTTAC
TTTTACTTCC TTAATACTCA AGTACAATTT TAATGGAGTA 4561 CTTTTTTACT
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- TTTACTCAAG TAAGATTCTA GCCAGATACT TTTACTTTTA ATTGAGTAAA 4621 ATTTTCCCTA AGTACTTGTA CTTTCACTTG AGTAAAATTT TTGAGTACTT TTTACACCTC 4681 TG.
- (102) Exemplary transposon/transposase systems of the disclosure include, but are not limited to, piggyBac and piggyBac-like transposons and transposases.
- (103) PiggyBac and piggyBac-like transposases recognizes transposon-specific inverted terminal repeat sequences (ITRs) on the ends of the transposon, and moves the contents between the ITRs into TTAA or TTAT chromosomal sites. The piggyBac or piggyBac-like transposon system has no payload limit for the genes of interest that can be included between the ITRs.
- (104) In certain embodiments, and, in particular, those embodiments wherein the transposon is a piggyBac transposon, the transposase is a piggyBac<sup>TM</sup>, Super piggyBac<sup>TM</sup> (SPB) transposase. In certain embodiments, and, in particular, those embodiments wherein the transposase is a piggyBac<sup>TM</sup>, Super piggyBac<sup>TM</sup> (SPB), the sequence encoding the transposase is an mRNA sequence.
- (105) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme.
- (106) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or a piggyBac-like transposase enzyme. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%. 95%, 99% or any percentage in between identical to: (107) TABLE-US-00027 (SEQ ID NO: 14487) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEI SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEQNVI EOPGSSLASN RILTLPORTI RGKNKHCWST SKSTRRSRVS ALNIVRSORG 121 PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTGATFRD TNEDEIYAFF 181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGFRGRCPF RMYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTQTN GVPLGEYYVK ELSKPVHGSC RNITCDNWFT SIPLALNLLQ 361 EPYKLTIVGT VRSNKREIPE VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481 SFIIYSHNVS SKGEKVQSRK KFMRNLYMSL TSSFMRKRLE APTLKPYLRD NISNILPNEV 541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC OSCF.
- (108) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme that comprises or consists of an amino acid sequence having an amino acid substitution at one or more of positions 30, 165, 282, or 538 of the sequence:
- (109) TABLE-US-00028 (SEQ ID NO: 14487) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEI SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEONVI EQPGSSLASN RILTLPQRTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121 PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTGATFRD TNEDEIYAFF 181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGERGRCPF RMYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTQTN GVPLGEYYVK ELSKPVHGSC RNITCDNWFT SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST GKPQMVNYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481 SFIIYSHNVS SKGEKVQSRK KFMRNLYMSL TSSFMRKRLE APTLKRYLRD NISNILPNEV 541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC QSCF.
- (110) In certain embodiments, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme that comprises or consists of an amino acid sequence having an amino acid substitution at two or more of positions 30, 165, 282, or 538 of the sequence of SEQ ID NO: 14487. In certain embodiments, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme that comprises or consists of an amino acid sequence having an amino acid substitution at three or more of positions 30, 165, 282, or 538

of the sequence of SEQ ID NO: 14487. In certain embodiments, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme that comprises or consists of an amino acid sequence having an amino acid substitution at each of the following positions 30, 165, 282, and 538 of the sequence of SEQ ID NO: 14487. In certain embodiments, the amino acid substitution at position 30 of the sequence of SEQ ID NO: 14487 is a substitution of a valine (V) for an isoleucine (I). In certain embodiments, the amino acid substitution at position 165 of the sequence of SEQ ID NO: 14487 is a substitution of a serine (S) for a glycine (G). In certain embodiments, the amino acid substitution at position 282 of the sequence of SEQ ID NO: 14487 is a substitution of a valine (V) for a methionine (M). In certain embodiments, the amino acid substitution at position 538 of the sequence of SEQ ID NO: 14487 is a substitution of a lysine (K) for an asparagine (N).

- (111) In certain embodiments of the methods of the disclosure, the transposase enzyme is a Super piggyBac<sup>™</sup> (SPB) or piggyBac-like transposase enzyme. In certain embodiments, the Super piggyBac<sup>™</sup> (SPB) or piggyBac-like transposase enzyme of the disclosure may comprise or consist of the amino acid sequence of the sequence of SEQ ID NO: 14487 wherein the amino acid substitution at position 30 is a substitution of a valine (V) for an isoleucine (1), the amino acid substitution at position 165 is a substitution of a serine (S) for a glycine (G), the amino acid substitution at position 282 is a substitution of a valine (V) for a methionine (M), and the amino acid substitution at position 538 is a substitution of a lysine (K) for an asparagine (N). In certain embodiments, the Super piggyBac<sup>™</sup> (SPB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (112) TABLE-US-00029 (SEQ ID NO: 14484) 1 MGSSLDDEHI LSALLQSDDE LVGEDSDSEV SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG 61 SEILDEONVI EQPGSSLASN RILTLPQRTI RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121 PTRMCRNIYD PLLCFKLFFT DEIISEIVKW TNAEISLKRR ESMTSATFRD TNEDEIYAFF 181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS VMSRDRFDFL IRCLRMDDKS IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ LLGFRGRCPF RVYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTQTN GVPLGEYYVK SIPLAKNLLQ 361 EPYKLTIVGT VRSNKREIPE ELSKPVHGSC RNITCDNWFT VLKNSRSRPV GTSMFCFDGP LTLVSYKPKP AKMVYLLSSC 421 DEDASINEST GKPQMVMYYN QTKGGVDTLD QMCSVMTCSR KTNRWPMALL YGMINIACIN 481 SKGEKVOSRK KFMRNLYMSL TSSFMRKRLE APTLKRYLRD SFIIYSHNVS NISNILPKEV 541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA NASCKKCKKV ICREHNIDMC QSCF.
- (113) In certain embodiments of the methods of the disclosure, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>TM</sup>, Super piggyBac<sup>TM</sup> or piggyBac-like transposase enzyme may further comprise an amino acid substitution at one or more of positions 3, 46, 82, 103, 119, 125, 177, 180, 185, 187, 200, 207, 209, 226, 235, 240, 241, 243, 258, 296, 298, 311, 315, 319, 327, 328, 340, 421, 436, 456, 470, 486, 503, 552, 570 and 591 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup>, Super piggyBac<sup>™</sup> or piggyBac-like transposase enzyme may further comprise an amino acid substitution at one or more of positions 46, 119, 125, 177, 180, 185, 187, 200, 207, 209, 226, 235, 240, 241, 243, 296, 298, 311, 315, 319, 327, 328, 340, 421, 436, 456, 470, 485, 503, 552 and 570. In certain embodiments, the amino acid substitution at position 3 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an asparagine (N) for a serine (S). In certain embodiments, the amino acid substitution at position 46 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a serine (S) for an alanine (A). In certain embodiments, the amino acid substitution at position 46 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a threonine (T) for an alanine (A). In certain embodiments, the amino acid substitution at position 82 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a tryptophan (W) for an isoleucine (I). In certain embodiments, the amino acid substitution at position 103 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P) for a serine (S). In certain embodiments, the amino acid substitution at position 119 of SEQ ID NO: 14487 or SEQ ID

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NO: 14484 is a substitution of a proline (P) for an arginine (R). In certain embodiments, the amino acid
substitution at position 125 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an alanine
(A) a cysteine (C). In certain embodiments, the amino acid substitution at position 125 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a cysteine (C). In certain embodiments,
the amino acid substitution at position 177 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution
of a lysine (K) for a tyrosine (Y). In certain embodiments, the amino acid substitution at position 177 of
SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a histidine (H) for a tyrosine (Y). In certain
embodiments, the amino acid substitution at position 180 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a leucine (L) for a phenylalanine (F). In certain embodiments, the amino acid substitution
at position 180 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a
phenylalanine (F). In certain embodiments, the amino acid substitution at position 180 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a valine (V) for a phenylalanine (F). In certain
embodiments, the amino acid substitution at position 185 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a leucine (L) for a methionine (M). In certain embodiments, the amino acid substitution at
position 187 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a glycine (G) for an alanine
(A). In certain embodiments, the amino acid substitution at position 200 of SEQ ID NO: 14487 or SEQ ID
NO: 14484 is a substitution of a tryptophan (W) for a phenylalanine (F) In certain embodiments, the
amino acid substitution at position 207 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
proline (P) for a valine (V). In certain embodiments, the amino acid substitution at position 209 of SEQ ID
NO: 14487 or SEQ ID NO: 14484 is a substitution of a phenylalanine (F) for a valine (V). In certain
embodiments, the amino acid substitution at position 226 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a phenylalanine (F) for a methionine (M). In certain embodiments, the amino acid
substitution at position 235 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an arginine
(R) for a leucine (L). In certain embodiments, the amino acid substitution at position 240 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a valine (V). In certain embodiments, the
amino acid substitution at position 241 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
leucine (L) for a phenylalanine (F). In certain embodiments, the amino acid substitution at position 243 of
SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a proline (P). In certain
embodiments, the amino acid substitution at position 258 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a serine (S) for an asparagine (N). In certain embodiments, the amino acid substitution at
position 2% of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a tryptophan (W) for a
leucine (L). In certain embodiments, the amino acid substitution at position 296 of SEQ ID NO: 14487 or
SEQ ID NO: 14484 is a substitution of a tyrosine (Y) for a leucine (L). In certain embodiments, the amino
acid substitution at position 296 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a
phenylalanine (F) for a leucine (L). In certain embodiments, the amino acid substitution at position 298 of
SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a methionine (M). In
certain embodiments, the amino acid substitution at position 298 of SEQ ID NO: 14487 or SEQ ID NO:
14484 is a substitution of an alanine (A) for a methionine (M). In certain embodiments, the amino acid
substitution at position 298 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a valine (V)
for a methionine (M). In certain embodiments, the amino acid substitution at position 311 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a proline (P). In certain
embodiments, the amino acid substitution at position 311 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
a substitution of a valine for a proline (P). In certain embodiments, the amino acid substitution at position
315 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for an arginine (R). In
certain embodiments, the amino acid substitution at position 319 of SEQ ID NO: 14487 or SEQ ID NO:
14484 is a substitution of a glycine (G) for a threonine (T). In certain embodiments, the amino acid
substitution at position 327 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an arginine
(R) for a tyrosine (Y). In certain embodiments, the amino acid substitution at position 328 of SEQ ID NO:
14487 or SEQ ID NO: 14484 is a substitution of a valine (V) for a tyrosine (Y). In certain embodiments,
the amino acid substitution at position 340 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution
of a glycine (G) for a cysteine (C). In certain embodiments, the amino acid substitution at position 340 of
SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a cysteine (C). In certain
embodiments, the amino acid substitution at position 421 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is
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a substitution of a histidine (H) for the aspartic acid (D). In certain embodiments, the amino acid substitution at position 436 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a valine (V). In certain embodiments, the amino acid substitution at position 456 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a tyrosine (Y) for a methionine (M). In certain embodiments, the amino acid substitution at position 470 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a phenylalanine (F) for a leucine (L). In certain embodiments, the amino acid substitution at position 485 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a serine (S). In certain embodiments, the amino acid substitution at position 503 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a leucine (L) for a methionine (M). In certain embodiments, the amino acid substitution at position 503 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an isoleucine (I) for a methionine (M). In certain embodiments, the amino acid substitution at position 552 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a lysine (K) for a valine (V). In certain embodiments, the amino acid substitution at position 570 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a threonine (T) for an alanine (A). In certain embodiments, the amino acid substitution at position 591 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P) for a glutamine (Q). In certain embodiments, the amino acid substitution at position 591 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an arginine (R) for a glutamine (Q). (114) In certain embodiments of the methods of the disclosure, including those embodiments wherein the

transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> or piggyBac-like transposase enzyme or may comprise or the Super piggyBac<sup>TM</sup> transposase enzyme may further comprise an amino acid substitution at one or more of positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments of the methods of the disclosure, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>TM</sup> or piggyBac-like transposase enzyme may comprise or the Super piggyBac™ transposase enzyme may further comprise an amino acid substitution at two, three, four, five, six or more of positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, including those embodiments wherein the transposase comprises the above-described mutations at positions 30, 165, 282 and/or 538, the piggyBac<sup>™</sup> or piggyBac-like transposase enzyme may comprise or the Super piggyBac<sup>™</sup> transposase enzyme may further comprise an amino acid substitution at positions 103, 194, 372, 375, 450, 509 and 570 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, the amino acid substitution at position 103 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a proline (P) for a serine (S). In certain embodiments, the amino acid substitution at position 194 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a valine (V) for a methionine (M). In certain embodiments, the amino acid substitution at position 372 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an alanine (A) for an arginine (R). In certain embodiments, the amino acid substitution at position 375 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an alanine (A) for a lysine (K) In certain embodiments, the amino acid substitution at position 450 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of an asparagine (N) for an aspartic acid (D). In certain embodiments, the amino acid substitution at position 509 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a glycine (G) for a serine (S). In certain embodiments, the amino acid substitution at position 570 of SEQ ID NO: 14487 or SEQ ID NO: 14484 is a substitution of a serine (S) for an asparagine (N). In certain embodiments, the piggyBac<sup>TM</sup> or piggyBac-like transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487. In certain embodiments, including those embodiments wherein the piggyBac<sup>TM</sup> or piggyBac-like transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487, the piggyBac<sup>™</sup> or piggyBaclike transposase enzyme may further comprise an amino acid substitution at positions 372, 375 and 450 of the sequence of SEQ ID NO: 14487 or SEQ ID NO: 14484. In certain embodiments, the piggyBac™ or piggyBac-like transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487, a substitution of an alanine (A) for an arginine (R) at position 372 of SEQ ID NO: 14487, and a substitution of an alanine (A) for a lysine (K) at position 375 of SEQ ID NO: 14487. In certain embodiments, the piggyBac™ or piggyBac-like transposase enzyme may comprise a substitution of a valine (V) for a methionine (M) at position 194 of SEQ ID NO: 14487, a substitution of

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an alanine (A) for an arginine (R) at position 372 of SEQ ID NO: 14487, a substitution of an alanine (A) for a lysine (K) at position 375 of SEQ ID NO: 14487 and a substitution of an asparagine (N) for an aspartic acid (D) at position 450 of SEQ ID NO: 14487.
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- (115) In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from an insect. In certain embodiments, the insect is *Trichoplusia ni* (GenBank Accession No. AAA87375; SEQ ID NO: 17083). Argyrogramma agnata (GenBank Accession No. GU477713; SEQ ID NO: 17084, SEQ ID NO: 17085), *Anopheles gambiae* (GenBank Accession No. XP\_312615 (SEQ ID NO: 17086); GenBank Accession No. XP\_320414 (SEQ ID NO: 17087); GenBank Accession No. XP\_310729 (SEQ ID NO: 17088)), Aphis gossypii (GenBank Accession No. GU329918; SEQ ID NO: 17089, SEQ ID NO: 17090), Acyrthosiphon pisum (GenBank Accession No. XP\_001948139; SEQ ID NO: 17091), Agrotis ipsilon (GenBank Accession No. GU477714; SEQ ID NO: 17092, SEQ ID NO: 17093). Bombyx mori (GenBank Accession No. BAD11135; SEQ ID NO: 17094). *Chilo suppressalis* (GenBank Accession No. JX294476; SEQ ID NO: 17095, SEQ ID NO: 17096). *Drosophila melanogaster* (GenBank Accession No. AAL39784; SEQ ID NO: 17097), Helicoverpa armigera (GenBank Accession No. ABS18391; SEQ ID NO: 17098). Heliothis virescens (GenBank Accession No. ABD76335; SEQ ID NO: 17099) Macdunnoughia crassisigna (GenBank Accession No. EU287451; SEQ ID NO: 17100, SEQ ID NO: 17101). Pectinophora gossypiella (GenBank Accession No. GU270322; SEQ ID NO: 17102, SEQ ID NO: 17103), Tribolium castaneum (GenBank Accession No. XP\_001814566; SEQ ID NO: 17104), Ctenoplusia agnata (also called Argyrogramma agnata), Messour bouvieri, Megachile rotundata, Bombus impatiens, Mamestra brassicae, Mayetiola destructor or Apis mellifera.
- (116) In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from an insect. In certain embodiments, the insect is *Trichoplusia ni* (AAA87375).
- (117) In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from an insect. In certain embodiments, the insect is *Bombyx mori* (BAD11135).
- (118) In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from a crustacean. In certain embodiments, the crustacean is *Daphnia pulicaria* (AAM76342, SEQ ID NO: 17105).
- (119) In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from a vertebrate. In certain embodiments, the vertebrate is *Xenopus tropicalis* (GenBank Accession No. BAF82026; SEQ ID NO: 17106), *Homo sapiens* (GenBank Accession No. NP\_689808; SEQ ID NO: 17107), *Mus musculus* (GenBank Accession No. NP\_741958; SEQ ID NO: 17108). *Macaca fascicularis* (GenBank Accession No. AB179012; SEQ ID NO: 17108, SEQ ID NO: 17109), *Rattus norvegicus* (GenBank Accession No. XP 220453; SEQ ID NO: 17110) or *Myotis lucifuqus*.
- (120) In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from a urochordate. In certain embodiments, the urochordate is *Ciona intestinalis* (GenBank Accession No. XP\_002123602; SEQ ID NO: 17111).
- (121) In certain embodiments, the piggyBac or piggyBac-like transposase inserts a transposon at the sequence 5'-TTAT-3' within a chromosomal site (a TTAT target sequence).
- (122) In certain embodiments, the piggyBac or piggyBac-like transposase inserts a transposon at the sequence 5'-TTAA-3' within a chromosomal site (a TTAA target sequence).
- (123) In certain embodiments, the target sequence of the piggy Bac or piggyBac-like transposon comprises or consists of 5'-CTAA-3', 5'-TTAG-3', 5'-ATAA-3', 5'-TCAA-3', 5'-AGTT-3', 5'-ATTA-3', 5'-GTTA-3', 5'-TTGA-3', 5'-TTTA-3', 5'-AGGT-3', 5'-AGGG-3', 5'-CTAG-3', 5'-TGAA-3', 5'-AGGT-3', 5'-ATCA-3', 5'-CTCC-3', 5'-TAAA-3', 5'-TCTC-3', 5'TGAA-3', 5'-AAAT-3', 5'-AATC-3', 5'-ACAA-3', 5'-ACAT-3', 5'-ACTC-3', 5'-AGTG-3', 5'-AGAG-3', 5'-CACA-3', 5'-CATA-3', 5'-CCAG-3', 5'-TCCA-3', 5'-TCCA
- (124) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Bombyx mori*. The piggyBac or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:

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1 MDIERQEERI RAMLEEELSD
(125) TABLE-US-00030 (SEQ ID NO: 14504)
YSDESSSEDE TDHCSEHEVN YDTEERIDS
                                          VDVPSNSRQE
                                                         61 EANAIIANES
              SLVRQRASAS RQVSGPFYTS
                                         KDGTKWYKNC QRPNVRLRSE 121
DSDPDDDLPL
NIVTEQAQVK NIARDASTEY ECWNIFVTSD
                                          MLQEILTHTN SSIRHRQTKT
AAENSSAETS 181 FYMQETTLCE LKALIALLYL
                                            AGLIKSNRQS LKDLWRTDGT
GVDIFRTTMS LQRFQFLQNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ
CCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN
FDVVNLEVYA GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG
YELMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421
VSYAPKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD
ELSANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRANKN VTIKRTEFIR
SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NVPGRYVRCQ
DCPYKKDRKT KHSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.
(126) The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid
sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%,
80%, 85%, 90° %, 95%, 99% or any percentage in between identical to:
(127) TABLE-US-00031 (SEQ ID NO: 14505)
                                            1 MDIERQEERI RAMLEEELSD
YSDESSSEDE TDHCSEHEVN YDTEEERIDS VDVPSNSRQE
                                                         61 EANAIIANES
DSDPDDDLPL SLVRQRASAS RQVSGPFYTS KDGTKWYKNC QRPNVRLRSE 121
NIVTEOAOVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRHROTKT
AAENSSAETS 181 FYMQETTLCE LKALIALLYL AGLIKSNRQS LKDLWRTDGT
GVDIFRTTMS LQRFQFLQNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ
CCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN
FYVVNLEVYA GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG
              YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421
YELMLHLLNE
VSYAPKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD
ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR
SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NVPGRYVRCQ
DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.
(128) In certain embodiments, the piggyBac or piggyBac-like transposase is fused to a nuclear localization
signal. In certain embodiments, the amino acid sequence of the piggyBac or piggyBac-like transposase
fused to a nuclear localization signal is encoded by a polynucleotide sequence comprising.
(129) TABLE-US-00032 (SEQ ID NO: 14629)
                                              1 atggcaccca aaaagaaacg taaagtgatg
         gacaggaaga aagaatcagg
                                 61 gcgatgctcg aagaagaact gagcgactac tccgacgaat
gacattgaaa
                    121 gaccactgta gcgagcatga ggttaactac gacaccgagg aggagagaat
cgtcatcaga
         ggatgaaacc
          181 gatgtgccct ccaactcacg ccaagaagag gccaatgcaa ttatcgcaaa cgaatcggac
                                                                           241
cgactctgtg
agcgatccag acgatgatct gccactgtcc ctcgtgcgcc agcgggccag cgcttcgaga
                                                             301 caagtgtcag
                                                  361 cgacctaacg tcagactccg
gtccattcta cacttcgaag
                   gacggcacta agtggtacaa gaattgccag
ctccgagaat atcgtgaccg aacaggctca ggtcaagaat
                                        421 atcgcccgcg acgcctcgac tgagtacgag
                             481 ctgcaagaaa ttctgacgca caccaacagc tcgattaggc
tgttggaata tcttcgtgac ttcggacatg
atcgccagac caagactgca
                     541 gcggagaact catcggccga aacctccttc tatatgcaag
                                                                agactactct
          601 aaggegetga ttgcactget gtacttggee ggeetcatea aateaaatag geagageete
                                                                          661
gtgcgaactg
                    tggaactgga gtggatatct ttcggacgac tatgagcttg
aaagatctct ggagaacgga
                                                            721 cagcggttcc
                                                  781 aggaaacaga ctgacaacat
agtttctgca
         aaacaatatc agattcgacg acaagtccac ccgggacgaa
         cggtcaatat tcgatcagtt tgtgcagtgc
                                       841 tgccaaaacg cttatagccc atcggaattc
ggctgcgttc
                             901 cgggggcgct gcctgttccg agtgtacatc ccgaacaagc
                   tctctccttc
ctgaccatcg
         acgaaatgct
         cggaatcaaa
                    961 atcctggccc tggtggacgc caagaatttc tacgtcgtga atctcgaagt
cggctaaata
gtacgcagga 1021 aagcaaccgt cgggaccgta cgctgtttcg aaccgcccgt ttgaagtcgt cgagcggctt 1081
         tggccagatc ccaccgcaat gttaccttcg acaattggtt caccggctac 1141 gagctgatgc ttcaccttat
attcagccgg
gaacgagtac
         cggctcacta gcgtggggac tgtcaggaag 1201 aacaagcggc agatcccaga atccttcatc
         gccagcctaa ctcgtccgtg 1261 ttcggatttc aaaaggatat cacgcttgtc tcgtacgccc
cgcaccgacc
          caaggtcgtg 1321 gtcgtgatga gcaccatgca tcacgacaac agcatcgacg
                                                                agtcaaccgg
agaaaagcaa 1381 aagcccgaga tgatcacctt ctacaattca actaaggccg gcgtcgacgt cgtggatgaa 1441
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actataacgt gtcccggaac tctaagcggt ggcctatgac tctcttctac 1501 ggagtgctga
ctgtgcgcga
          aatcaacgcg tgcatcatct accgcaccaa caagaacgtg 1561 accatcaagc gcaccgagtt
atatggccgc
         ctgggtttga gcatgatcta cgagcacctc 1621 cattcacgga acaagaagaa gaatatccct
catcagatcg
acttacctga ggcagcgtat cgagaagcag 1681 ttgggagaac caagcccgcg ccacgtgaac gtgccggggc
gctacgtgcg gtgccaagat 1741 tgcccgtaca aaaaggaccg caaaaccaaa
                                                       agatcgtgta acgcgtgcgc
caaacctatc 1801 tgcatggagc atgccaaatt tctgtgtgaa aattgtgctg aactcgattc ctccctg.
(130) In certain embodiments the piggyBac or piggyBac-like transposase is hyperactive. A hyperactive
piggyBac or piggyBac-like transposase is a transposase that is more active than the naturally occurring
variant from which it is derived. In certain embodiments, the hyperactive piggyBac or piggyBac-like
transposase enzyme is isolated or derived from Bombyx mori. In certain embodiments, the piggyBac or
piggy Bac-like transposase is a hyperactive variant of SEQ ID NO: 14505. In certain embodiments, the
hyperactive piggyBac or piggyBac-like transposase comprises a sequence that is at least 90% identical to:
(131) TABLE-US-00033 (SEQ ID NO: 14576) 1 MDIERQEERI RAMLEEELSD
YSDESSSEDE TDHCSEHEVN YDTEEERIDS VDVPSNSRQE
                                                        61 EANAIIANES
DSDPDDDLPL SLVRQRASAS RQMSGPHYTS KDGTKWYKNC QRPNVRLRSE 121
NIVTEQAQVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRWRQTKT
AAENSSASTS 181 FYMQETTLCE LKALIGLLYI AGLIKSNRQS LKDLWRTDGT
GVDIFRTTMS LQRFQFLQNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ
SCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN
FYVKNLEVYA GKQPSGPYAV
                             SNRPFEVVER LIOPVARSHR 361 NVTFDNWFTG
YELMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421
VSYARKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD
ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR
SLGLSMIYEH LHSRNKKKNI 541 PTYLKRQIEK QLGEPSPRHV NVPGRYVRCQ
DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSHL.
(132) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises SEQ ID
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- (132) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14576. In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:
- (133) TABLE-US-00034 (SEQ ID NO: 14630) 1 MDIERQEERI RAMLEEELSD YSDESSSEDE TDHCSEHEVN YDTEEERIDS **VDVPSNSRQE 61 EANAIIANES** DSDPDDDLPL SLVRQRASAS RQVSGPFYTS KDGTKWYKNC QRPNVRLRSE 121 NIVTEQAQVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRWRQTKT AAENSSAFTS 181 FYMQETTLCE LKALIGLLYI AGLIKSNRQS LKDLWRTDGT GVDIFRTTMS LQRFQFLLNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ SCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN FYVHNLEVYA GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG YEVMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VEGFQKDITL 421 VSYAPKKNKV VVVMSTMHHD NSIDESTGEK **QKPEMITFYN STKAGVDVVD** ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NVPGRYVRCQ DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAHLDS.
- (134) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:
- (135) TABLE-US-00035 (SEQ ID NO: 14631) 1 MDIERQEERI RAMLEEELSD **61 EANAIIANES** YSDESSSEDE TDHCSEHEVN YDTEERIDS **VDVPSNSRQE** SLVRQRASAS RQVSGPFYTS KDGTKWYKNC QRPNVRLRSE 121 DSDPDDDLPL MLQEILTHTN SSIRWRQTKT NIVTEQAQVK NIARDASTEY ECWNIFVTSD AAENSSASTS 181 FYMQETTLCE LKALIGLLYI AGLIKSNRQS LKDLWRTDGT GVDIFRTTMS LQRFQFLLNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN SCONAYSPSE FYVKNLEVYA GKQPSGPYAV **SNRPFEVVER** LIQPVARSHR 361 NVTFDNWFTG YELMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421 VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD VSYAPKKNKV

ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIAM QLGEPSPRHV NVPGRYVRCQ DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.

(136) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:

(137) TABLE-US-00036 (SEQ ID NO: 14632) 1 MDIERQEERI RAMLEEELSD YSDESSSEDE TDHCSEHEVN **YDTEERIDS VDVPSNSRQE 61 EANAIIANES** DSDPDDDLPL SLVRQRASAS RQVSGPFYTS KDGTKWYKNC QRPNVRLRSE 121 NIVTEQAQVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRWRQTKT AAENSSAETS 181 FYMQETTLCE LKALIGLLYI AGLIKSNRQS LKDLWRTDGT ERKQTDNMAA FRSIFDQFVQ GVDIFRTTMS LQRFQFLLNN 241 IRFDDKSTRD SCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN FYVKNLEVYA GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG YRLTSVGTVR KNKTQIPENF IRTDROPNSS VFGFQKDITL 421 YELMLHLLNE VSYAPKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD ELQANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NVPGRYVRCQ DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.

(138) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:

(139) TABLE-US-00037 (SEQ ID NO: 14633) 1 MDIERQEERI RAMLEEELSD YSDESSSEDE TDHCSEHEVN YDTEERIDS VDVPSNSRQE **61 EANAIIANES** DSDPDDDLPL SLVRQRASAS RQVSGPFYTS KDGTKWYKNC QRPNVRLRSE 121 NIVTEQAQVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRWRQTKT AAENSSAETS 181 FYMQETTLCE LKALIGLLYI AGLIKSNRQS LKDLWRTDGT GVDIFRTTMS LQRFQFLQNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ SCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN FYVKNLEVYA GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG YELMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421 VSYAPKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NVPGRYVRCQ DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.

(140) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:

(141) TABLE-US-00038 (SEQ ID NO: 14634) 1 MDIERQEERI RAMLEEELSD YSDESSSEDE **TDHCSEHEVN YDTEERIDS VDVPSNSRQE 61 EANAIIANES** DSDPDDDLPL SLVRQRASAS RQVSGPFYTS KDGTKWYKNC QRPNVRLRSE 121 NIVTEQAQVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRHRQTKT AAENSSAETS 181 FYMQETTLCE LKALIALLYL AGLIKSNRQS LKDLWRTDGT GVDIFRTTMS LQRFQFLQNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ CCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG DYVVNLEVYA YELMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421 VSYAPKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKPTEFIR SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSSRHV NVKGRYVRCQ DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.

(142) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase is more active than the transposase of SEQ ID NO: 14505. In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase is at least 90%, at least 95%, at least 97%, at least 98%, or at least 99% or any percentage in between identical to SEQ ID NO: 14505.

(143) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises an amino

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acid substitution at a position selected from 92, 93, 96, 97, 165, 178, 189, 196, 200, 201, 211, 215, 235,
238, 246, 253, 258, 261, 263, 271, 303, 321, 324, 330, 373, 389, 399, 402, 403, 404, 448, 473, 484, 507,
523, 527, 528, 543, 549, 550, 557, 601, 605, 607, 609, 610 or a combination thereof (relative to SEQ ID
NO: 14505). In certain embodiments, the hyperactive piggy Bac or piggyBac-like transposase comprises
an amino acid substitution of Q92A, V93L, V93M, P96G, F97H, F97C, H165E, H165W, E178S, E178H,
C189P, A196G, L200I, A201Q, L211A, W215Y, G219S, Q235Y, Q235G, Q238L, K246I, K253V, M258V,
F261L, S263K, C271S, N303R, F321W, F321D, V324K, V324H, A330V, L373C, L373V, V389L, S399N,
R402K, T403L, D404Q, D404S, D404M, N441R, G448W, E449A, V469T, C473Q, R484K T507C,
G523A, I527M, Y528K Y543I, E549A, K550M, P557S, E601V, E605H, E605W, D607H, S609H, L610I
or any combination thereof. In certain embodiments, the hyperactive piggyBac or piggyBac-like
transposase comprises an amino acid substitution of Q92A, V93L, V93M, P96G, F97H, F97C, H165E,
H165W, E178S, E178H, C189P, A196G, L200I, A201Q, L211A, W215Y, G219S, Q235Y, Q235G,
Q238L, K246I, K253V, M258V, F261L, S263K, C271S, N303R, F321W, F321D, V324K, V324H, A330V,
L373C, L373V, V389L, S399N, R402K, T403L, D404Q, D404S, D404M, N441R, G448W, E449A,
V469T, C473Q, R484K T507C, G523A, I527M, Y528K Y543I, E549A, K550M, P557S, E601V, E605H,
E605W, D607H, S609H and L610I.
(144) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises one or
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- more substitutions of an amino acid that is not wild type, wherein the one or more substitutions a for wild type amino acid comprises a substitution of E4X, A12X, M13X, L14X, E15X, D20X, E24X, S25X, S26X, S27X, D32X, H33X, E36X, E44X, E45X, E46X, I48X, D49X, R58X, A62X, N63X, A64X, I65X, I66X, N68X, E69X, D71X, S72X, D76X, P79X, R84X, Q85X, A87X, S88X, Q92X, V93X, S94X, G95X, P96X, F97X, Y98X, T99X, I145X, S149X, D150X, L152X, E154X, T157X, N160X, S161 X, S162X, H165X, R166X, T168X, K169X, T170X, A171X, E173X, S175X, S176X, E178X, T179X, M183X, Q184X, T186X, T187X, L188X, C189X, L194X, I195X, A196X, L198X, L200X, A201X, L203X, I204X, K205X, A206X, N207X, Q209X, S210X, L211X, K212X, D213X, L214X, W215X, R216X, T217X, G219X, V222X, D223X, I224X, T227X, M229X, Q235X, L237X, Q238X, N239X, N240X, P302X, N303X, P305X, A306X, K307X, Y308X, I310X, K311X, 1312X, L313X, A314X, L315X, V316X, D317X, A318X, K319X, N320X, F321X, Y322X, V323X, V324X, L326X, E327X, V328X, A330X, Q333X, P334X, S335X, G336X, P337X, A339X, V340X, S341X, N342X, R343X, P344X, F345X, E346X, V347X, E349X, I352X, Q353X, V355X, A356X, R357X, N361X, D365X, W367X, T369X, G370X, L373X, M374X, L375X, H376X, N379X, E380X, R382X, V386X, V389X, N392X, R394X, O395X, S399X, F400X, I401X, R402X T403X, D404X, R405X, O406X, P407X, N408X, S409X, S410X, V411X, F412X, F414X, Q415X, I418X, T419X, L420X, N428X V432X, M434X, D440X, N441X, S442X, I443X, D444X, E445X, G448X, E449X, Q451X, K452X, M455X, 1456X, T457X, F458X, S461X, A464X, V466X, Q468X, V469X, E471X, L472X, C473X, A474X, K483X, W485X, T488X, L489X, Y491X, G492X, V493X, M496X, I499X, C502X, I503X, T507X, K509X, N510X, V511X, T512X, I513X, R515X, E517X, S521X, G523X, L524X, S525X, I527X, Y528X, E529X, H532X, S533X, N535X, K536X, K537X, N539X, I540X, T542X, Y543X, Q546X, E549X, K550X, Q551X, G553X, E554X, P555X, S556X, P557X, R558X, H559X, V560X, N561X, V562X, P563X, G564X, R565X, Y566X, V567X, Q570X, D571X, P573X, Y574X, K576X, K581X, S583X, A586X, A588X, E594X, F598X, L599X, E601X, N602X, C603X, A604X, E605X, L606X, D607X, S608X, S609X or L610X (relative to SEQ ID NO: 14505). A list of hyperactive amino acid substitutions can be found in U.S. Pat. No. 10,041,077, the contents of which are incorporated herein by reference in their entirety. (145) In certain embodiments, the piggyBac or piggyBac-like transposase is integration deficient. In certain embodiments, an integration deficient piggyBac or piggyBac-like transposase is a transposase that can excise its corresponding transposon, but that integrates the excised transposon at a lower frequency than a corresponding wild type transposase. In certain embodiments, the piggyBac or piggyBac-like transposase is an integration deficient variant of SEQ ID NO: 14505.
- (146) In certain embodiments, the excision competent, integration deficient piggyBac or piggyBac-like transposase comprises one or more substitutions of an amino acid that is not wild type, wherein the one or more substitutions a for wild type amino acid comprises a substitution of R9X, A12X, M13X, D20X, Y21K, D23X, E24X, S25X, S26X, S27X, E28X, E30X, D32X, H33X, E36X, H37X, A39X, Y41X, D42X, T43X, E44X, E45X, E46X, R47X, D49X, S50X, S55X, A62X, N63X, A64X, I66X, A67X, N68X,

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E69X, D70X, D71X, S72X, D73X, P74X, D75X, D76X, D77X, I78X, S81 X, V83X, R84X, Q85X, A7X,
S88X, A89X, S90X, R91X, Q92X, V93X, S94X, G95X, P96X, F97X, Y98X, T99X, W012X, G103X,
Y107X, K108X, L117X, I122X, Q128X, I312X, D135X, S137X, E139X, Y140X, I145X, S149X, D150X,
Q153X, E154X, T157X, S61X, S162X, R164X, H165X, R166X, Q167X, T168X, K169X, T170X,
A171X, A172X, E173X, R174X, S175X, S176X, A177X, E178X, T179X, S180X, Y182X, Q184X,
E185X, T187X, L188X, C189X, L194X, I195X, A196X, L198X, L200X, A201X, L203X, I204X, K205X,
N207X, Q209X, L21X, D213X, L214X, W215X, R216X, T217X, G219X, T220X, V222X, D223X,
I224X, T227X, T228X, F234X, Q235X, L237X, Q238X, N239X, N240X, N303X, K304X, I310X,
I312X, L313X, A314X, L315X, V316X, D317X, A318X, K319X, N320X, F321X, Y322X, V323X,
V324X, N325X, L326X, E327X, V328X, A330X, G331X, K332X, Q333X, S335X, P337X, P344X,
F345X, E349X, H359X, N361X, V362X, D365X, F368X, Y371X, E372X, L373X, H376X, E380X,
R382X, R382X, V386X, G387X, T388X, V389X, K391X, N392X, R394X, Q395X, E398X, S399X,
F400X, I401X, R402X T403X, D404X, R405X, Q406X, P407X, N408X, S409X, S410X, Q415X,
K416X, A424X, K426X, N428X, V430X, V432X, V433X, M434X, D436X, D440X, N441X, S442X,
I443X, D444X, E445X, S446X, 0,447X, G448X, E449X, K450X, Q451X, E454X, M455X, I456X,
T457X, F458X, S461X, A464X, V466X, Q468X, V469X, C473X, A474X, N475X, N477X, K483X,
R484X, P486X, T488X, L489X, G492X, V493X, M496X, I499X, I503X, Y505X, T507X, N510X,
V511X, T512X, I513X, K514X, T516X, E517X, S521X, G523X, L524X, S525X, I527X, Y528X, L531X,
H532X, S533X, N535X, I540X, T542X, Y543X, R545X, Q546X, E549X, L552X, G553X, E554X,
P555X, S556X, P557X, R558X, H559X, V560X, N561X, V562X, P563X, G564X, V567X, Q570X,
D571X, P573X, Y574X, K575X, K576X, N585X, A586X, M593X, K596X, E60X, N602X, A604X,
E605X, L606X, D607X, S608X, S609X or L610X (relative to SEQ ID NO: 14505). A list of integration
deficient amino acid substitutions can be found in U.S. Pat. No. 10,041,077, the contents of which are
incorporated by reference in their entirety.
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(147) In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises a sequence of:

(148) TABLE-US-00039 (SEQ ID NO: 14606) 1 MDIERQEERI RAMLEEELSD **TDHCSEHEVN VDVPSNSRQE** YSDESSSEDE **YDTEERIDS 61 EANAIIANES** DSDPDDDLPL **SLVRQRASAS RQVSSPFYTS** KDGTKWYKNC **ORPNVRLRSE 121** NIVTEQAQVK NIARDASTEY **ECWNIFVTSD** MLQEILTHTN **SSIRHRQTKT** AAENSSAETS 181 FYMQETTLCE LKALIALLYL **AGLIKSNRQS** LKDLWRKDGT **GVDIFRTTMS** LQRFQFLLNN 241 IRFDDISTRD ERKQTDNMAA FRSIFDQFVQ CCQNAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN GKQPSGPYAV **SNRPFEVVER** LIQPVARSHR 361 NVTFDNWFTG FYVVNLEVYA YELMLHLLNE YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421 VSYAPKKNKV VVVMSTMHHD NSIDESTGEK **QKPEMITFYN STKAGVDVVD** ELCANYNVSR 481 NSKKWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR LHSRNKKKNI 541 PTYLQQRIEK QLGEPVPRHV SLGLSMMYEH NVPGRYVRCO KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL. DCPYKKDRKT

In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises a sequence of:

(149) TABLE-US-00040 (SEQ NO: 14607) 1 MDIERQEERI RAMLEEELSD **TDHCSEHEVN VDVPSNSRQE** YSDESSSEDE **YDTEERIDS 61 EANAIIANES** DSDPDDDLPL **SDVRQRASAS RQVSGPFYTS** KDGTKWYKNC **QRPNVRLRSE 121** NIVTEQAQVK **NIARDASTEY ECWNIFVTSD** MLQEILTHTN **SSIRHRQTKT** AAENSSAETS 181 FYMQETTLCE LKALIGLLYL AGLIKSNRQS LKDLWRTDGT **GVDIFRTTMS** LORFYFLONN 241 IRFDDKSTLD **ERKOTDNMAA** FRSIFDOFVO **SCQNAYSPSE** FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN FYVVNLEVYA **GKQPSGPYAV SNPRFEVVER** LIQPVARSHR 361 NVTFDNWFTG YELMLHLLNE YRLTSVGTVR **KNKROIPESF IRTDROPNSS** VFGFQKDITL 421 NSIDESTGEK **QKPEMITFYN STKAGVDVVD** VSYAPKKNKV VVVMSTMHHD ELCANYNVSR 481 NSKRWPMTLF YGVLNMAAIN ACIIYPTNKN VTIKRTEFIR SLGLSMIYEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NYPGRYVRCQ

DCPYKKDRKT KRSCNACAKP ICMEHAKFLC 601 VNCAELDSSL.

In certain embodiments, the piggyBac or piggyBac-like transposase that is integration deficient comprises a sequence of:

(150) TABLE-US-00041 (SEQ ID NO: 14608) 1 MDIERQEERI RAMLEEELSD YSDESSSEDE TDHCSEHEVN YDTEERIDS **VDVPSNSRQE 61 EANAIIANES** DSDPDDDLPL SLVPQRASAS RQVSGPFYTS KDGTKWYKNC QPPNVLRRSE 121 NIVTEQAQVK NIARDASTEY ECWNIFVTSD MLQEILTHTN SSIRHRQTKT AAENSSAETS 181 FYMQETTLCE LKALIALLYL AGLIKSNRQS LKDLWRKDGT GVDIFRTTMS LQRFQFLLNN 241 IRFDDKSTRD ERKQTDNMAA FRSIFDQFVQ CCONAYSPSE FLTIDEMLLS FRGRCLFRVY 301 IPNKPAKYGI KILALVDAKN DYVVNLEVYA GKQPSGPYAV SNRPFEVVER LIQPVARSHR 361 NVTFDNWFTG YRLTSVGTVR KNKRQIPESF IRTDRQPNSS VFGFQKDITL 421 YECMLHLLNE VSYAPKKNKV VVVMSTMHHD NSIDESTGEK QKPEMITFYN STKAGVDVVD ELCANYNVSR 421 NSKKWPMTLF YGVLNMAAIN ACIIYRTNKN VTIKRTEFIR SLGLSMIKEH LHSRNKKKNI 541 PTYLRQRIEK QLGEPSPRHV NVPGRYVRCQ DCPYRKDRKT KRSCNACAKP ICMEHAKFLC 601 ENCAELDSSL.

In certain embodiments, the integration deficient transposase comprises a sequence that is at least 90% identical to SEQ ID NO: 14608.

- (151) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Bombyx mori*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (152) TABLE-US-00042 (SEQ ID NO: 14506) 1 ttatcccggc gagcatgagg cagggtatct cataccatgg taaaatttta aagttgtgta 61 ttttataaaa ttttcgtctg acaacactag cgcgctcagt agctggaggc aggagcgtgc 121 gggaggggat agtggcgtga tcgcagtgtg gcacgggaca ccggcgagat attcgtgtgc 181 aaacctgttt cgggtatgtt ataccctgcc tcattgttga cgtattttt ttatgtaatt 241 tttccgatta ttaatttcaa ctgttttatt ggtattttta tgttatccat tgttcttttt 301 ttatgattta ctgtatcggt tgtctttcgt tcctttagtt gagtttttt ttattatttt 361 cagtttttga tcaaa.

In certain embodiments, the piggyBac $^{\text{TM}}$  (PB) or piggyBac-like transposon comprises a sequence of: (155) TABLE-US-00045 (SEQ ID NO: 14509) 1 taaataataa taatttcata attaaaact tctttcattg aatgccatta aataaaccat 61 tattttacaa aataagatca acataattga gtaaataata ataagaacaa tattatagta 121 caacaaaata tgggtatgtc ataccctgcc acattcttga tgtaactttt tttcacctca 181 tgctcgccgg gttat.

(156) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a left sequence corresponding to SEQ ID NO: 14507. In certain embodiments, one piggyBac or piggyBac-like transposon end is at least 85%, at least 90%, at least 95%, at least 98%, at least 990% identical or any percentage in between identical to SEQ ID NO: 14506 and the other piggyBac or piggyBac-like transposon end is at least 85%, at least 90%, at least 95%, at least 98%, at least 99% or any percentage in between identical to SEQ ID NO: 14507. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14506 and SEQ ID NO: 14507 or SEQ ID NO: 14509. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14508 and SEQ ID NO: 14507 or SEQ ID NO: 14508 and SEQ ID NO: 14507 or SEQ ID NO: 14509. In certain embodiments, the left and right transposon ends share a 16 bp repeat sequence at their ends of CCCGGCGAGCATGAGG (SEQ ID NO:

14510) immediately adjacent to the 5′-TTAT-3 target insertion site, which is inverted in the orientation in the two ends. In certain embodiments, left transposon end begins with a sequence comprising 5′-TTATCCCGGCGAGCATGAGG-3 (SEQ ID NO: 14511), and the right transposon ends with a sequence comprising the reverse complement of this sequence: 5′-CCTCATGCTCGCCGGGTTAT-3′ (SEQ ID NO: 14512).

(157) In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end comprising at least 14, 16, 18, 20, 30 or 40 contiguous nucleotides of SEQ ID NO: 14506 or SEQ ID NO: 14508. In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end comprising at least 14, 16, 18, 20, 30 or 40 contiguous nucleotides of SEQ ID NO: 14507 or SEQ ID NO: 14509. In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end with at least 90% identity to SEQ ID NO: 14506 or SEQ ID NO: 14508. In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end with at least 90% identity to SEQ ID NO: 14507 or SEQ ID NO: 14509. (158) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (159) TABLE-US-00046 (SEQ ID NO: 14515) 1 ttaacccggc gagcatgagg cagggtatct acaacactag cgcgctcagt agctggaggc 61 ttttataaaa ttttcgtctg cataccctgg taaaatttta aagttgtgta aggagcgtgc 121 gggaggggat agtggcgtga tcgcagtgtg gcacgggaca ccggcgagat attcgtgtgc 181 aaacctgttt cgggtatgtt ataccctgcc tcattgttga cgtatttttt ttatgtaatt 241 tttccgatta ctgttttatt ggtattttta tgttatccat tgttcttttt 301 ttatgattta ctgtatcggt tgtctttcgt tcctttagtt gagtttttt ttattatttt 361 cagtttttga tcaaa.

(160) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (161) TABLE-US-00047 (SEQ ID NO: 14516) 1 tcatattttt agtttaaaaa aataattata tgttttataa 61 ttcagtatta aataatattt ttgttaaatt gttgattttt 121 tgaaaagaat ctcattatct ggttgattta tattccaaag gtaaacctct aaatgtttgc tgctaaaatt actgtgttta agaaaaagat taataaataa 181 taataatttc ataattaaaa acttctttca ttgaatgcca ttaaataatt cattatttta 241 caaaataaga tcaacataac tgagtaaata ataataagaa caatattata gtacaacaaa 301 atatgggtat gtcataccct tttttttttt ttttttttt ttctttcggg tagagggccg 361 aacctcctac gaggtccccg cgcaaaaggg gcgcgcgggg tatgtgagac tcaacgatct 421 gcatggtgtt gtgagcagac cgcgggccca aggattttag agcccaccca ctaaacgact 481 cctctgcact cttacacccg ccctccgagg tcagaacccg gatgaggtag 541 gggggctacc gcggtcaaca ctacaaccag acgtccgatc acggcgcggc tcaccccaag gacgcccagc 601 cgacggagcc ttcgaggcga atcgaaggct ctgaaacgtc ggccgtctcg gtacggcagc 661 ccgtcgggcc gcccagacgg tgccgctggt gtcccggaat accccgctgg accagaacca 721 gcctgccggg tcgggacgcg atacaccgtc gaccggtcgc tccaatcact ccacggcagc 721 gcgctagagt gctggta.

(162) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of CCCGGCGAGCATGAGG (SEQ ID NO: 14510). In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of SEQ ID NO: 14510. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of TTATCCCGGCGAGCATGAGG (SEQ ID NO: 14511). In certain embodiments, the piggyBac or piggyBac-like transposon comprises at least 16 contiguous nucleotides from SEQ 1D NO: 14511. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of CCTCATGCTCGCCGGGTTAT (SEQ ID NO: 14512). In certain embodiments, the piggyBac or piggyBac-like transposon comprises at least 16 contiguous nucleotides from SEQ ID NO: 14512. In certain embodiments, the piggyBac-like transposon comprising at least 16 contiguous nucleotides from SEQ ID NO: 14511 and one end comprising at least 16 contiguous nucleotides from SEQ ID NO: 14512. In certain embodiments, the piggyBac-like transposon comprises SEQ ID NO: 14511 and SEQ ID NO: 14512. In certain embodiments, the piggyBac-like transposon comprises a sequence of TTAACCCGGCGAGCATGAGG (SEQ ID NO 14513). In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of CCTCATGCTCGCCGGGTTAA (SEQ ID NO: 14514).

(163) In certain embodiments, the piggyBac or piggyBac-like transposon may have ends comprising SEQ ID NO: 14506 and SEQ ID NO: 14507, or a variant of either or both of these having at least 90% sequence identity to SEQ ID NO: 14506 or SEQ ID NO: 14507, and the piggyBac or piggyBac-like transposase has the sequence of SEQ ID NO: 14504 or SEQ ID NO: 14505, or a sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identity to SEQ ID NO: 14504 or SEQ ID NO: 14505. In certain

embodiments, the piggyBac or piggyBac-like transposon comprises a heterologous polynucleotide inserted between a pair of inverted repeats, where the transposon is capable of transposition by a piggyBac or piggyBac-like transposase having at least 5%, 10%, 15%, 20%, 25%, 30%, 35%. 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identity to SEQ ID NO: 14504 or SEQ ID NO: 14505. In certain embodiments, the transposon comprises two transposon ends, each of which comprises SEQ ID NO: 14510 in inverted orientations in the two transposon ends. In certain embodiments, each inverted terminal repeat (ITR) is at least 90% identical to SEQ ID NO: 14510. (164) In certain embodiments, the piggyBac or piggyBac-like transposon is capable of insertion by a piggyBac or piggyBac-like transposase at the sequence 5'-TTAT-3 within a target nucleic acid. In certain embodiments, one end of the piggyBac or piggyBac-like transposon comprises at least 16 contiguous nucleotides from SEQ ID NO: 14506 and the other transposon end comprises at least 16 contiguous nucleotides from SEQ ID NO: 14507 In certain embodiments, one end of the piggyBac or piggyBac-like transposon comprises at least 17, at least 18, at least 19, at least 20, at least 22, at least 25, at least 30 contiguous nucleotides from SEQ ID NO: 14506 and the other transposon end comprises at least 17, at least 18, at least 19, at least 20, at least 22, at least 25, at least 30 contiguous nucleotides from SEQ ID NO: 14507.

(165) In certain embodiments, the piggyBac or piggyBac-like transposon comprises transposon ends (each end comprising an ITR) corresponding to SEQ ID NO: 14506 and SEQ ID NO: 14507, and has a target sequence corresponding to 5'-TTAT3'. In certain embodiments, the piggyBac or piggyBac-like transposon also comprises a sequence encoding a transposase (e.g. SEQ ID NO: 14505). In certain embodiments, the piggyBac or piggyBac-like transposon comprises one transposon end corresponding to SEQ ID NO: 14506 and a second transposon end corresponding to SEQ ID NO: 14516. SEQ ID NO: 14516 is very similar to SEQ ID NO: 14507, but has a large insertion shortly before the ITR. Although the ITR sequences for the two transposon ends are identical (they are both identical to SEQ ID NO: 14510), they have different target sequences: the second transposon has a target sequence corresponding to 5'-TTAA-3', providing evidence that no change in ITR sequence is necessary to modify the target sequence specificity. The piggyBac or piggyBac-like transposase (SEQ ID NO: 14504), which is associated with the 5'-TTAA-3' target site differs from the 5'-TTAT-3'-associated transposase (SEQ ID NO: 14505) by only 4 amino acid changes (D322Y, S473C, A507T, H582R). In certain embodiments, the piggyBac or piggyBac-like transposase (SEQ ID NO: 14504), which is associated with the 5'-TTAA-3' target site is less active than the 5'-TTAT-3'-associated piggyBac or piggyBac-like transposase (SEQ ID NO: 14505) on the transposon with 5'-TTAT-3' ends. In certain embodiments, piggyBac or piggyBac-like transposons with 5'-TTAA-3' target sites can be converted to piggyBac or piggyBac-like transposases with 5'-TTAT-3 target sites by replacing 5'-TTAA-3' target sites with 5'-TTAT-3'. Such transposons can be used either with a piggyBac or piggyBac-like transposase such as SEQ ID NO: 14504 which recognizes the 5'-TTAT-3' target sequence, or with a variant of a transposase originally associated with the 5'-TTAA-3' transposon. In certain embodiments, the high similarity between the 5'-TTAA-3' and 5'-TTAT-3' piggyBac or piggyBac-like transposases demonstrates that very few changes to the amino acid sequence of a piggyBac or piggyBaclike transposase alter target sequence specificity. In certain embodiments, modification of any piggyBac or piggyBac-like transposon-transposase gene transfer system, in which 5'-TTAA-3' target sequences are replaced with 5'-TTAT-3'-target sequences, the ITRs remain the same, and the transposase is the original piggyBac or piggyBac-like transposase or a variant thereof resulting from using a low-level mutagenesis to introduce mutations into the transposase. In certain embodiments, piggyBac or piggyBac-like transposon transposase transfer systems can be formed by the modification of a 5'-TTAT-3'-active piggyBac or piggyBac-like transposon-transposase gene transfer systems in which 5'-TTAT-3' target sequences are replaced with 5'-TTAA-3'-target sequences, the ITRs remain the same, and the piggyBac or piggy Bac-like transposase is the original transposase or a variant thereof.

(166) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Bombyx mori*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:

(167) TABLE-US-00048 (SEQ ID NO: 14577) 1 cccggcgagc atgaggcagg gtatctcata ccctggtaaa attttaaagt tgtgtatttt 61 ataaaatttt cgtctgacaa cactagcgcg ctcagtagct ggaggcagga gcgtgcggga 121 ggggatagtg gcgtgatcgc agtgtggcac gggacaccgg cgagatattc gtgtgcaaac 181

ctgtttcggg tatgttatac cctgcctcat tgttgacgta t.

In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
(168) TABLE-US-00049 (SEQ ID NO: 14578) 1 tttaagaaaa agattaataa ataataataa
tttcataatt aaaaacttct ttcattgaat 61 gccattaaat aaaccattat tttacaaaat aagatcaaca taattgagta
aataataata 121 agaacaatat tatagtacaa caaaatatgg gtatgtcata ccctgccaca ttcttgatgt 181 aacttttttt

cacctcatgc tcgccggg.

In certain embodiments, the transposon comprises at least 16 contiguous bases from SEQ ID NO: 14577 and at least 16 contiguous bases from SEQ ID NO: 14578, and inverted terminal repeats that are at least 87% identical to CCCGGCGAGCATGAGG (SEQ ID NO: 14510). In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:

(169) TABLE-US-00050 (SEQ ID NO: 14595) 1 cccggcgagc atgaggcagg gtatctcata ccctggtaaa attttaaagt tgtgtatttt 61 ataaaatttt cgtctgacaa cactagcgcg ctcagtagct ggaggcagga gcgtgcggga 121 ggggatagtg gcgtgatcgc agtgtggcac gggacaccgg cgagatattc gtgtgcaaac 181 ctgtttccgg tatgttatac cctgcctcat tgttgacgta ttttttttat gtaatttttc 241 cgattattaa tttcaactgt tttattggta ttttttgtt atccattgtt cttttttat 301 gatttactgt atcggttgtc tttcgttcct ttagttgagt tttttttat tattttcagt 361 ttttgatcaa a.

In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (170) TABLE-US-00051 (SEQ ID NO: 14596) 1 tcatattttt agtttaaaaa aataattata tgttttataa tgaaaagaat ctcattatct 61 ttcagtatta ggttgattta tattccaaag aataatattt ttgttaaatt gttgattttt 121 gtaaacctct aaatgtttgt tgctaaaatt actgtgttta agaaaaagat taataaataa 181 taataatttc ataattaaaa acttctttca ttgaatgcca ttaaataaac cattatttta 241 caaaataaga tcaacataat tgagtaaata ataataagaa caatattata gtacaacaaa 301 atatgggtat gtcataccct gccacattct tgatgtaact tttttcacc tcatgctcgc 361 cggg.

- (171) In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14595 and SEQ ID NO: 14596, and is transposed by the piggyBac or piggyBac-like transposase of SEQ ID NO: 14505. In certain embodiments, the ITRs of SEQ ID NO: 14595 and SEQ ID: 14596 are not flanked by a 5'-TTAA-3' sequence. In certain embodiments, the ITRs of SEQ ID NO: 14595 and SEQ ID: 14596 are flanked by a 5'-TTAT-3' sequence.
- (172) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (173) TABLE-US-00052 (SEQ ID NO: 14597) 1 cccggcgagc atgaggcagg gtatctcata ccctggtaaa attttaaagt tgtgtatttt 61 ataaaatttt cgtctgacaa cactagcgcg ctcagtagct ggaggcagga gcgtgcggga 121 ggggatagtg gcgtgatcgc agtgtggcac gggacaccgg cgagatattc gtgtgcaaac 181 ctgtttcggg tatgttatac cctgcctcat tgttgacgta ttttttttat gtaatttttc 241 cgattattaa tttcaactgc tttattggta tttttttgtt atccattgtt ctttttttat 301 g.

In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (174) TABLE-US-00053 (SEQ ID NO: 14598) 1 cagggtatct catacctgg taaaatttta aagttgtgta ttttataaaa ttttcgtctg 61 acaacactag cgcgctcagt agctggaggc aggagggtgc gggaggggat agtggcgtga 121 tcgcagtgtg gcacgggaca ccggcgagat attcgtgtgc aaacctgttt cgggtatgtt 181 ataccctgcc tcattgttga cgtatttttt ttatgtaatt tttccgatta ttaatttcaa 241 ctgttttatt ggtattttta tgttatccat tgttcttttt ttatg.

In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (175) TABLE-US-00054 (SEQ ID NO: 14599) 1 cagggtatct cataccetgg taaaatttta aagttgtgta ttttataaaa ttttcgtctg 61 acaacactag cgcgctcagt agctggaggc aggagggtc gggaggggat agtggcgtga 121 tcgcagtgtg gcacgggaca ccggcgagat attcgtgtgc aaacctgttt cgggtatgtt 181 ataccetgcc tcattgttga cgtat.

In certain embodiments, the left end of the piggyBac or piggyBac-like transposon comprises a sequence of SEQ ID NO: 14577, SEQ ID NO: 14595, or SEQ ID NOs: 14597-14599. In certain embodiments, the left end of the piggyBac or piggyBac-like transposon is preceded by a left target sequence.

In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (176) TABLE-US-00055 (SEQ ID NO: 14600) 1 tcatattttt agtttaaaaa aataattata tgttttataa tgaaaagaat ctcattatct 61 ttcagtatta ggttgattta tattccaaag aataatattt ttgttaaatt gttgattttt 121 gtaaacctct aaatgtttgt tgctaaaatt actgtgttta agaaaaagat taataaataa 181 taataatttc ataattaaaa acttctttca ttgaatgcca ttaaataaac cattatttta 241 caaaataaga tcaacataat tgagtaaata ataataagaa

caatattata gtacaacaaa 301 atatgggtat gtcataccct gccacattct tgatgtaact ttttttcacc tcatgctcgc 351 cggg.

In certain embodiments the piggyBac or piggyBac-like transposon comprises a sequence of: (177) TABLE-US-00056 (SEQ ID NO: 14601) 1 tttaagaaaa agattaataa ataataataa tttcataatt aaaaacttct ttcattgaat 61 gccattaaat aaaccattat tttacaaaat aagatcaaca taattgagta aataataata 121 agaacaatat tatagtacaa caaaatatgg gtatgtcata ccctgccaca ttcttgatgt 181 aactttttt ca. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (178) TABLE-US-00057 (SEQ ID NO: 14602) 1 cccggcgagc atgaggcagg gtatctcata ccctggtaaa attttaaagt tgtgtatttt 61 ataaaatttt cgtctgacaa cactagcgcg ctcagtagct ggaggcagga gcgtgcggga 121 ggggatagtg gcgtgatcgc agtgtggcac gggacaccgg cgagatattc gtgtgcaaac 181 ctgtttcqgq tatgttatac cctqcctcat tgttgacqta ttttttttat gtaattttc 241 cgattattaa tttcaactgt tttattggta ttttatgtt atccattgtt ctttttttat 301 gatttactgt atcggttgtc tttcgttcct ttagttgagt ttttttttat tattttcagt 361 ttttgatcaa a.

(179) In certain embodiments, the right end of the piggyBac or piggyBac-like transposon comprises a sequence of SEQ ID NO: 14578, SEQ ID NO: 14596, or SEQ ID NOs: 14600-14601. In certain embodiments, the right end of the piggyBac or piggyBac-like transposon is followed by a right target sequence. In certain embodiments, the transposon is transposed by the transposase of SEQ ID NO: 14505 In certain embodiments, the left and right ends of the piggyBac or piggyBac-like transposon share a 16 bp repeat sequence of SEQ ID NO: 14510 in inverted orientation and immediately adjacent to the target sequence. In certain embodiments, the left transposon end begins with SEO ID NO: 14510, and the right transposon end ends with the reverse complement of SEQ ID NO: 14510, 5'-CCTCATGCTCGCCGGG-3' (SEQ ID NO: 14603). In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR with at least 93%, at least 87%, or at least 81% or any percentage in between identity to SEQ ID NO: 14510 or SEQ ID NO: 14603. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a target sequence followed by a left transposon end comprising a sequence selected from SEQ ID NOs: 88, 105 or 107 and a right transposon end comprising SEQ ID NO: 14578 or 106 followed by a target sequence, in certain embodiments, the piggyBac or piggyBac like transposon comprises one end that comprises a sequence that is at least 90%, at least 95% or at least 99% or any percentage in between identical to SEQ ID NO: 14577 and one end that comprises a sequence that is at least 90%, at least 95% or at least 99% or any percentage in between identical to SEQ ID NO: 14578. In certain embodiments, one transposon end comprises at least 14, at least 16, at least 18 or at least 20 contiguous bases from SEQ ID NO: 14577 and one transposon end comprises at least 14, at least 16, at least 18 or at least 20 contiguous bases from SEQ ID NO: 14578.

(180) In certain embodiments, the piggyBac or piggyBac-like transposon comprises two transposon ends wherein each transposon ends comprises a sequence that is at least 81% identical, at least 87% identical or at least 93% identical or any percentage in between identical to SEQ ID NO: 14510 in inverted orientation in the two transposon ends. One end may further comprise at least 14, at least 18 or at least 20 contiguous bases from SEQ ID NO: 14599, and the other end may further comprise at least 14, at least 16, at least 18 or at least 20 contiguous bases from SEQ ID NO: 14601. The piggyBac or piggyBac-like transposon may be transposed by the transposase of SEQ ID NO: 14505, and the transposase may optionally be fused to a nuclear localization signal.

(181) In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14595 and SEQ ID NO: 14596 and the piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14504 or SEQ ID NO: 14505. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14597 and SEQ ID NO: 14596 and the piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14504 or SEQ ID NO: 14505. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14504 or SEQ ID NO: 14505. In certain embodiments, the piggyBac-like transposase comprises SEQ ID NO: 14504 or SEQ ID NO: 14505. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14602 and SEQ ID NO: 14600 and the piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14504 or SEQ ID NO: 14505.

(182) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a left end comprising 1, 2, 3, 4, 5, 6, or 7 sequences selected from ATGAGGCAGGGTAT (SEQ ID NO: 14614), ATACCCTGCCTCAT (SEQ ID NO: 14615), GGCAGGGTAT (SEQ ID NO: 14616), ATACCCTGCC

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(SEQ ID NO: 14617), TAAAATITTA (SEQ ID NO: 14618), ATITUATAAAAT (SEQ ID NO: 14619).
TCATACCCTG (SEQ ID NO: 14620) and TAAATAATAA (SEQ ID NO: 14621). In certain
embodiments, the piggyBac or piggyBac-like transposon comprises a right end comprising 1, 2 or 3
sequences selected from SEQ ID NO: 14617. SEQ ID NO: 14620 and SEQ ID NO: 14621.
(183) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or
piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase
enzyme is isolated or derived from Xenopus tropicalis. The piggyBac or piggyBac-like transposase
enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%,
35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in
between identical to:
(184) TABLE-US-00058 (SEQ ID NO: 14317)
                                            1 MAKRFYSAEE
                                                           AAAHCMASSS
                          TEESWCSSST VSALEEPMEV
EEFSGSDSEY VPPASESDSS
                                                       61 DEDVDDLEDQ
EAGDRADAAA GGEPAWGPPC
                             NFPPEIPPFT TVPGVKVDTS
                                                        NEEPINFFQL 121
FMTEAILQDM VLYTNVYAEQ
                            YLTQNPLPRY APAHAWHPTD
                                                         IAEMKRFVGL
TLAMGLIKAN 181 SLESYWDTTT VLSIPVFSAT
                                                          RFLHFNNNAT
                                           MSRNRYQLLL
AVPPDQPGHD RLHKLRPLID 241 SLSERFAAVY
                                           TPCQNICIDE
                                                        SLLLFKGRLQ
FRQYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL
                                                        DPPGCPPDLT
VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLDT 361 PACGTINRNR
KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKNNVFMLT
                                                         SIHDESVIRE 421
                            MGGVDRTDQL
                                           OHYYNATRKT
ORVGRPPKNK PLCSKEYSKY
                                                          RAWYKKVGIY
LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV
                                                          EEQTVPEMPP
SDNVARLIGK
             HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD
                                                          TRYYCPKCPR
NPGLCFKPCF EIYETQLHY.
(185) In some embodiments, the piggyBac or piggyBac-like transposase is a hyperactive variant of SEQ
ID NO: 14517. In certain embodiments, the piggyBac or piggyBac-like transposase is an integration
defective variant of SEQ ID NO: 14517. The piggyBac or piggyBac-like transposase enzyme may
comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%,
50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
(186) TABLE-US-00059 (SEQ ID NO: 14518)
                                            1 MAKRFYSAEE AAAHCMAPSS
EEFSGSDSEY VRPASESDSS
                           TEESWCSSST
                                        VSALEEPMEV
                                                        61 DEDVDDLEDQ
EAGDRADAAA GGEPAWGPPC
                            NFPPEIPPFT TVPGVKVDTS
                                                        NFEPINFFQL 121
FMTEAILQDM VLYTNVYAEQ YLTQNPLPRY ARAHAWHPTD
                                                          IAEMKRFVGL
TLAMGLIKAN 181 SLESYWNTTT VLSIPVFSAT
                                           MSRNRYQLLL
                                                          RFLHFNNNAT
AVPPDQPDHD RLHKLRPLID 241 SLSERFAAVY
                                           TPCQNICIDE SLLLFKGRLR
FROYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL
                                                        DPPGCPPDLT
VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLDT 361 PACGTINRTR
KGLPRALLDK KLNRGETYAL RKNELLAIKF
                                          FDKKNVFMLT
                                                         SIHDESVIRE 421
QRVGRPPKNK PLCSKEYSKY MGGVDPTDQL
                                           QHYYNATRKT
                                                          SAWYKKVGIY
LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV
                                                          EEQTVPEMLP
SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD
                                                          TRYYCPKCPR
NPGLCFKPCF EIYHTOLHY.
(187) In certain embodiments, the piggyBac or piggyBac-like transposase is isolated or derived from
Xenopus tropicalis. In certain embodiments, the piggyBac or piggyBac-like transposase is a hyperactive
piggyBac or piggyBac-like transposase. In certain embodiments, the hyperactive piggyBac or piggyBac-
like transposase comprises a sequence at least 90% identical to:
(188) TABLE-US-00060 (SEQ ID NO: 14572)
                                            1 MAKRFYSAEE AAAHCSASSS
EEFSGSDSEY VPPASESDSS
                          TEESWCSSST
                                       VSALEEPMEV
                                                       61 DEDVDDLEDQ
EAGDRADAAA GGEPAWGPPC
                             NFPPEIPPFT TVPGVKVDTS
                                                        NFEPINFFQL 121
FMTEAILQDM VLYTNVYAEQ YLTQNPLTRG
                                          ARAHAWHPTD IAEMKRFVGL
TLAMGLIKAN 181 SIESYWDTTT VLSIPVFGAT
                                           MSRNRYOLLL
                                                          RFLHFNNNAT
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TPCQNICIDE

**SLMLFKGRLQ** 

**DPPGCPPDLT** 

AVPPDQPGHD RLHKLRPLID 241 SLSERFANVY

RYGIKFYKLC ESSTGYTSYF 301 LIYEGKDSKL

SPLLGQGFHL YVDNFYSSIP LFTALYCLNT 361 PACGTINRNR

FROYIPSKRA

VSGKIVWELI

- KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDPTDQL QHYYNATRKT RHWYKKVGIY LIQMALRNSY 481 IVYKAAYPGP KLSYYKYQLQ ILPALLFGGV EEQTVPEMPD SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCRKPCF EIYHTQLHY.
- (189) In certain embodiments, piggyBac or piggyBac-like transposase is a hyperactive piggyBac or piggyBac-like transposase. A hyperactive piggyBac or piggyBac-like transposase is a transposase that is more active than the naturally occurring variant from which it is derived. In certain embodiments, a hyperactive piggyBac or piggyBac-like transposase is more active than the transposase of SEQ ID NO: 14517. In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:
- (190) TABLE-US-00061 (SEQ ID NO: 14572) 1 MAKRFYSAEE AAAHCSASSS EEFSGSDSEY VPPASESDSS TEESWCSSST VSALEEPMEV 61 DEDVDDLEDQ EAGDRADAAA GGEPAWGPPC NFPPEIPPFT **TVPGVKVDTS** NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQNPLTRG ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SIESYWDTTT VLSIPVFGAT MSRNRYQLLL RFLHFNNNAT AVPPDQPGHD RLHKLRPLID 241 SLSERFANVY TPCQNICIDE SLMLFKGRLQ FRQYIPSKRA RYGIKFYKLC ESSTGYTSYF 301 LIYEGKDSKL DPPGCPPDLT VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLNT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDPTDQL QHYYNATRKT RHWYKKVGIY LIQMALRNSY 481 IVYKAAYPGP KLSYYKYQLQ ILPALLFGGV **EEQTVPEMPD** SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCRKPCF EIYHTQLHY.
- (191) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:
- (192) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:
- (193) TABLE-US-00062 (SEQ ID NO: 14624) 1 MAKRFYSAEE AAAHCMASSS TEESWCSSST VSALEEPMEV EEFSGSDSEY VPPASESDSS 61 DEDVDDLEDQ EAGDRADAAA GGEPAWGPPC NFPPEIPPFT TVPGVKVDTS NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQNPLTRY ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SLESYWDTTT VLSIPVESAT MSRNRYQLLL RFLHENNNAT AVPPDQPGHD RLHKLRPLID 241 SLSERFAAVY TPCQNICIDE SLLLFKGRLQ FRQYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL DPPGCPPDLT VSGKIVWELI SPLLSQGFHL YVDNFYSSIP LFTALYCLNT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDRTDQL QHYYNATRKT RHWYKKVGIY LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV **EEQTVPEMPP** SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCRKPCF EIYHTQLHY.
- (194) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:
- 1 MAKRFYSAEE AAAHCMASSS (195) TABLE-US-00063 (SEQ ID NO: 14625) EEFSGSDSEY VPPASESDSS TEESVCSSST VSALEEPMEV 51 DEDVDDLEDQ EAGDRADAAA GGEPAWGPPC NFPPEIPPFT TVPGVKVDTS NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQNPLPRY ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SLESYWDTTT VLKIPVFSAT MSRNRYQLLL RFLHFNNNAT AVPPDQPGHD RLHKLRPLID 241 SLSERFAAVY TPCQNICIDE SLLIFKGRLQ FROYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL DPPGCPPDLT VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLNT 351 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDRTDQL QHYYNATRKT RHWYKKVGIY

LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV EEQTVPEMPP SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCFKPCF EIYHTQLHY.

(196) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:

(197) TABLE-US-00064 (SEQ ID NO: 14627) 1 MAKRFYSAEE AAAHCMASSS **TEESWCSSST** EOTSGSDSEY VPPASESDSS VSALEEPMEV 61 DEDVDDLEDQ EAGDRADAAA **GGEPAWGPPC** NFPPEIPPFT **TVPCVKVDTS** NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQNPLTRY ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SIESYWDTTT VLSIPVFGAT **MSRNRYQLLL RFLHFNNNAT** AVPPDQPGHD RLHKLRPLID 241 SLSERFANVY **TPCQNICIDE SLLLFKGRLQ** FRQYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL **DPPGCPPDLT** VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLNT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF **FDKKNVFMLT** SIHDESVIRE 421 QRVGRKPKNK PLCSKEYSKY MGGVDRTDQL **QHYYNATRKT** RHWYKKVGIY KLSYYKYQLQ ILPALLFGGV LIQMALRNSY 481 IVYKAAVPGP **EEQTVPEMPP** HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD SDNVARLIGK **TRYYCPKCPR** NPGLCRKPCF EIYHTQLHY.

(198) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:

(199) TABLE-US-00065 (SEQ ID NO: 14628) 1 MAKRFYSAEE AAAHCSASSS EEFSGSDSEY VPPASESDSS TEESWCSSST VSALEEPMEV 61 DEDVDDLEDQ EAGDRADAAA GGEPAWGPPC NFPPEIPPFT TVPGVKVDTS NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQNPLTRG ARAHAWHPTD **IAEMKRFVGL** TLAMGLIKAN 181 SLESYWDTTT VLSIPVFGAT **MSRNRYQLLL RFLHFNNNAT** AVPPDQPGHD RLHKLRPLID 241 SLSERFANVY **TPCQNICIDE** SLMLFKGRLQ FRQYIPSKRA RYGIKFYKLC ESSTGYTSYF 301 LIYEGKDSKL DPPGCPPDLT VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLNT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDRTDQL **QHYYNATRKT** RHWYKKVGIY LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV **EEQTVPEMPP** HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD SDNVARLIGK TRYYCPKCPR NPGLCRKPCF EIYHTQLHY.

(200) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises a sequence of:

(201) (SEQ ID NO: 17042).

(202) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises an amino acid substitution at a position selected from amino acid 6, 7, 16, 19, 20, 21, 22, 23, 24, 26, 28, 31, 34, 67, 73, 76, 77, 88, 91, 141, 145, 146, 148, 150, 157, 162, 179, 182, 189, 192, 193, 196, 198, 200, 210, 212, 218, 248, 263, 270, 294, 297, 308, 310, 333, 336, 354, 357, 358, 359, 377, 423, 426, 428, 438, 447, 450, 462, 469, 472, 498, 502, 517, 520, 523, 533, 534, 576, 577, 582, 583 or 587 (relative to SEQ ID NO: 14517). In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises an amino acid substitution of Y6C, S7G, M16S, S19G, S20Q, S20G, S20D, E21D, E22Q, F23T, F23P, S24Y, S26V, S28Q, V31K, A34E, L67A, G73H, A76V, D77N, P88A, N91D, Y141Q, Y141A, N145E, N145V, P146T, P146V, P146K, P148T, P148H, Y150G, Y150S, Y50C, H157Y, A162C, A179K, L182I, L182V, T189G, L192H, S193N, S193K, V1%, S198G, T200W, L210H, F212N, N218E, A248N, L263M, Q270L, S294T, T297M, S308R, L310R, L333M, Q336M, A354H, C357V, L358F, D359N, L3771, V423H, P426K, K428R, S438A, T447G, T447A, L450V, A462H, A462Q, I469V, I472L, Q498M, L502V, E5171, P520D, P520G, N523S, I533E, D534A, F576R, F576E, K577I, I582R, Y583F, L587Y or L587W, or any combination thereof including at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or all of these mutations (relative to SEQ ID NO: 14517).

(203) In certain embodiments, the hyperactive piggyBac or piggyBac-like transposase comprises one or more substitutions of an amino acid that is not wild type, wherein the one or more substitutions a for wild

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type amino acid comprises a substitution of A2X, K3X, R4X, F5X, Y6X, S7X, A11X, A3X, C15X, M16X,
A17X, S18X, S19X, S20X, E21X, E22X, F23X, S24X, G25X, 26X, D27X, S28X, E29X, E42X, E43X,
S44X, C46X, S47X, S48X, S49X, T50X, V51X, S52X, A53X, L54X, E55X, E56X, P57X, M58X, E59X,
E62X, D63X, V64X, D65X, D66X, L67X, E68X, D69X, Q70X, E71X, A72X, G73X, D74X, R75X,
A76X, D77X, A78X, A79X, A80X, G81X, G82X, E83X, P84X, A85X, W86X, G87X, P88X, P89X,
C90X, N91X, F92X, P93X, E95X, I96X, P97X, P98X, F99X, T100X, T101X, P103X, G104X, V105X,
K106X, V107X, D108X, T109X, N111 X, P114X, I115X, N116X, F117X, F118X, Q119X, M122X,
T123X, E124X, A125X, I126X, L127X, Q128X, D129X, M130X, L132X, Y133X, V126X, Y127X,
A138X, E139X, Q140X, Y141X, L142X, Q144X, N145X, P146X, L147X, P148X, Y150X, A151X,
A155X, H157X, P158X, I161X, A162X, V168X, T171X, L72X, A173X, M174X, I177X, A179X, L182X,
D187X, T188X, T189X, T190X, L192X, S193X, I194X, P195X, V196X, S198X, A199X, T200X, S202X,
L208X, L209X, L210X, R21 X, F212X, F215X, N217X, N218X, A219X, T220X, A221X, V222X,
P224X, D225X, Q226X, P227X, H229X, R231X, H233X, L235X, P237X, I239X, D240X, L242X,
S243X, E244X, R244X, F246X, A247X, A248X, V249X, Y250X, T251X, P252X, C253X, Q254X,
I256X, C257X, I258X, D259X, E260X, S261X, L262X, L263X, L264X, F265X, K266X, G267X,
R268X, L269X, Q270X, F271X, R272X, Q273X, Y274X, I275X, P276X, S277X, K278X, R279X,
A280X, R281X, Y282X, G283X, I284X, K285X, F286X, Y287X, K288X, L289X, C290X, E291X,
S292X, S293X S294X, G295X, Y296X, T297X, S298X, Y299X, F300X, E304X, L310X, P313X,
G314X, P316X, P317X, D318X, L319X, T320X, V321X, K324X, E328X, I330X, S331X, P332X,
L333X, L334X, G335X, O336X, F338X, L340X, D343X, N344X, F345X, Y346X, S347X, L351X,
F352X, A354X, L355X, Y356X, C357X, L358X, D359X, T360X, R422X, Y423X, G424X, P426X,
K428X, N429X, K430X, P431X, L432X, S434X, K435X, E436X, S438X, K439X, Y440X, G443X,
R446X, T447X, L450X, Q451X, N455X, T460X, R461X, A462X, K465X, V467X, G468X, I469X,
Y470X, L471X, I472X, M474X, A475X, L476X, R477X, S479X, Y480X, V482X Y483X, K484X,
A485X, A486X, V487X, P488X, P490X, K491X, S493X, Y494X, Y495X, K496X, Y497T, Q498X,
L499X, Q500X, I501X, L502X, P503X, A504X, L505X, L506X, F507X, G508X, G509X, V510X,
E511X, E512X, Q513X, T514X, V515X, E517X, M518X, P519X, P520X, S521X, D522X, N523X,
V524X, A525X, L527X, I528X, K530X, H531X, F532X, 1533X, D534X, T535X, L536X, T539X,
P540X, Q546X, K550X, R553X, K554X, R555X, G556X, I557X, R558X, R559X, D560X, T561X,
Y564X, P566X, K567X, P569X, R570X, N571X, L574X, C575X, F576X, K577X, P578X, F580X,
E581X, I582X, Y583X, T585X, Q586X, L587X, H588X or Y589X (relative to SEQ ID NO: 14517). A list
of hyperactive amino acid substitutions can be found in U.S. Pat. No. 10,041,077, the contents of which
are incorporated by reference in their entirety.
(204) In certain embodiments, the piggyBac or piggyBac-like transposase is integration deficient. In
certain embodiments, an integration deficient piggyBac or piggyBac-like transposase is a transposase that
can excise its corresponding transposon, but that integrates the excised transposon at a lower frequency
than a corresponding naturally occurring transposase. In certain embodiments, the piggyBac or piggyBac-
like transposase is an integration deficient variant of SEQ ID NO: 14517. In certain embodiments, the
integration deficient piggyBac or piggyBac-like transposase is deficient relative to SEQ ID NO: 14517.
(205) In certain embodiments, the piggyBac or piggyBac-like transposase is active for excision but
deficient in integration. In certain embodiments, the integration deficient piggyBac or piggyBac-like
transposase comprises a sequence that is at least 90% identical to a sequence of:
(206) TABLE-US-00066 (SEQ
                            ID
                               NO: 14605)
                                                 1 MAKRFYSAEE AAAHCMASSS
                              TEESWCSSST
EEFSGSDSEY
               VPPASESDSS
                                             VSALEEPMEV
                                                             61 DEDVDDLEDO
EAGDRVDAAA
                GGEPAWGPPC
                                NFPPEIPPFT
                                              TVPGVKVDTS
                                                              NFEPINFFQL 121
FMTEAILQDM
                VLYTNVYAEQ
                                YLTQNPLPRY
                                               ARAHAWHPTD
                                                                IAEMKRFVGL
TLAMGLIKAN 181 SLESYWDTTT VLSIPVFSAT
                                                MSRNRYQLLL
                                                                KFLHFNNEAT
AVPPDQPGHD
               RLHKLRPLID 241 SLSERFAAVY
                                                TPCQNICIDE
                                                               SLLLFKGRLQ
FRQYIPSKRA
               RYGIKFYKLC
                              ESSSGYTSYF 301 LIYEGKDSKL
                                                               DPPGCPPDLT
VSGKIVWELI
               SPLLGQGFHL
                              YVDNFYSSIP
                                             LFTALYCLDT 361 PACGTINRNR
KGLPRALLDK
               KLNRGETYAL
                                RKNELLAIKF
                                               FDKKNVFMLT
                                                               SIHDESVIRE 421
ORVGRPPKNK
                PLCSKEYSKY
                               MGGVDRTDQL
                                                OHYYNATRKT
                                                                 RAWYKKVGIY
LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV
                                                                EEQTVPEMPP
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SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCFKPCF EIYHTQLHYG RR.

(207) In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises a sequence that is at least 90% identical to a sequence of:

(208) TABLE-US-00067 (SEQ ID NO: 14604) 1 MAKRFYSAEE AAAHCMASSS EEFSGSDSEY VPPASESDSS TEESWCSSST VSALEEPMEV 61 DEDVDDLEDQ EAGDRADAAA GGEPAWGPPC NFPPEIPPFT TVPGVKVDTS NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQVPLPRY ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SLESYWDTTT VLNIPVFSAT MSRNRYQLLL RFLEFNNEAT AVPPDQPGHD RLHKLRPLID 241 SLSERFAAVY TPCQNICIDE SLLLFKGRLQ FRQYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL DPPGCPPDLT VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLDT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QPVGRPPKNK PLCSKEYSKY MGGVDRTDQL QHYYNATRKT RAWYKKVGIY LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV **EEQTVPEMPP** SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCFKPCF EIYHTQLHY.

- (209) In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises a sequence that is at least 90% identical to a sequence of:
- (210) TABLE-US-00068 (SEQ ID NO: 14611) 1 MAKRFYSAEE AAAHCMASSS EEFSGSDSEY VPPASESDSS TEESWCSSST VSALEEPMEV 61 DEDVDDLEDQ EAGDRADAAA GGEPAWGPPC NFPPEIPPFT TVPGVKVDTS NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQNVLPRY ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SLESYWDTTT VLSIPVFSAT MSRNRYQLLL **RFLHFNNDAT** AVPPDQPGHD RLHKLRPLID 241 SLTERFAAVY TPCQNICIDE SLLLFKGRLQ FRQYIPSKRA RYGIKFYKLC ESSSGYTSYF 301 LIYEGKDSKL DPPGCPPDLT VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLDT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDRTDQL QHYYNATRKT RAWYKKVGIY LIQMALRNSY 481 IVYKAAYPGP KLSYYKYQLQ ILPALLFGGV **EEQTVPEMPP** SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD TRYYCPKCPR NPGLCFKPCF EIYHTOLHYG RR.
- (211) In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14611. In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises a sequence that is at least 90% identical to a sequence of:
- (212) TABLE-US-00069 (SEQ ID NO: 14612) 1 MAKRFYSAEE ALAHCMASSS TEESWCSSST VSALEEPMEV EEFSGSDSEY VPPASESDSS 61 DEDVDDLEDQ EAGDRADAAP GGEPAWGPPC NFPPEIPPFT TVPGVKVDTS NFEPINFFQL 121 FMTEAILQDM VLYTNVYAEQ YLTQVPLPRY ARAHAWHPTD IAEMKRFVGL TLAMGLIKAN 181 SLESYWDTTT VLSIPVFSAT MSRNRYQLLL RFLHFNNEAT AVPPDQPGHD RLHKLRPLID 241 SLSERFAAVY TPCQNICIDE SLLLFKGRLQ FRQYIPSKRA RYGIYFYKLC ESSSGYTSYF 301 LIYEGKDSKL DPPGCPDDLT VSGKIVWELI SPLLGQGFHL YVDNFYSSIP LFTALYCLDT 361 PACGTINRNR KGLPRALLDK KLNRGETYAL RKNELLAIKF FDKKNVFMLT SIHDESVIRE 421 QRVGRPPKNK PLCSKEYSKY MGGVDRTDQL QHYYNATRKT RAWYKKVGIY LIQMALRNSY 481 IVYKAAVPGP KLSYYKYQLQ ILPALLFGGV **EEQTVPEMPP** SDNVARLIGK HFIDTLPPTP 541 GKQRPQKGCK VCRKRGIRRD **TRYYCPKCPR** NPGLCFKPCF EIYHTQLHYG RR.
- (213) In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14612. In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises a sequence that is at least 90% identical to a sequence of (214) TABLE-US-00070 (SEQ ID NO: 14613) 1 MAKRFYSAEE AAAHCMASSS EEFSGSDSEY VPPASESDSS TEESWCSSST VSALEEPMEV

61 DEDVDDLEDO

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EAGDRADAAA
             GGEPAWGPPC
                          NFPPEIPPFT
                                     TVPGVKVDTS
                                                 NFEPINFFQL 121
FMTEAILQDM
            VLYTNVYAEQ
                         YLTQVPLPRY
                                     ARAHAWHPTD
                                                   IAEMKRFVGL
TLAMGLIKAN 181 SLESYWDTTT
                          VLNIPVFSAT
                                      MSRNRYQLLL
                                                   RFLEFNNNAT
AVPPDQPGHD
            RLHKLRPLID 241 SLSERFAAVY
                                      TPCQNICIDE
                                                  SLLLFKGRLO
FROYIPSKRA
            RYGIKFYKLC
                        ESSSGYTSYF 301 LIYEGKDSKL
                                                  DPPGCPPDLT
            SPLLGQGFHL
                                    LFTALYCLDT 361 PACGTINRNR
VSGKIVWELI
                        YVDNEYSSIP
KGLPRALLDK
            KLNRGETYAL
                         RKNELLAIKF
                                     FDKKNVFMLT
                                                  SIHDESVIRE 421
QRVGRPPKNK
            PLCSKEYSKY
                         MGGVDRTDQL
                                      QHYYNATRKT
                                                   RAWYKKVGIY
                          KLSYYKYQLQ
LIQMALRNSY 481 IVYKAAVPGP
                                       ILPALLFGGV
                                                   EEQTVPEMPP
SDNVARLIGK
            HFIDTLPPTP 541 GKORPOKGCK
                                      VCRKRGIRRD
                                                   TRYYCPKCPR
NPGLCFKPCF
            EIYHTQLHYG RR.
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- (215) In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14613. In certain embodiments, the integration deficient piggyBac or piggyBac-like transposase comprises an amino acid substitution wherein the Asn at position 218 is replaced by a Glu or an Asp (N218D or N218E) (relative to SEQ ID NO: 14517).
- (216) In certain embodiments, the excision competent, integration deficient piggyBac or piggyBac-like transposase comprises one or more substitutions of an amino acid that is not wild type, wherein the one or more substitutions a for wild type amino acid comprises a substitution of A2X, K3X, R4X, F5X, Y6X, S7X, A8X, E9X, E10X, A11X, A12X, A13X, H14X, C15X, M16X, A17X, S18X, S19X, S20X, E21X, E22X, F23X, S24X, G25X, 26X, D27X, S28X, E29X, V31X, P32X, P33X, A34X, S35X, E36X, S37X, D38X, S39X, S40X, T41X, E42X, E43X, S44X, W45X, C46X, S47X, S48X, S49X, T50X, V51X, S52X, A53X, L54X, E55X, E56X, P57X, M58X, E59X, V60X, M122X, T123X, E124X, A125X, L127X, Q128X, D129X, L132X, Y133X, V126X, Y127X, E139X, Q140X, Y141X, L142X, T43X, Q144X, N145X, P146X, L147X, P148X, R149X, Y150X, A151X, H154X, H157X, P158X, T159X, D160X, I161X, A162X, E163X, M164X, K165X, R166X, F167X, V168X, G169X, L170X, T171X, L172X, A173X, M174X, G175X, L176X, I177X, K178X, A179X, N180X, S181X, L182X, S184X, Y185X, D187X, T188X, T89X, T190X, V191X, L192X, S193X, 1194X, P195X, V196X, F197X, S198X, A199X, T200X, M201X, S202X, R203X, N204X, R205X, Y206X, Q207X, L208X, L209X, L210X, R211X, F212X, L213X, H241X, F215X, N216X, N217X, N218X, A219X, T220X, A221X, V222X, P223X, P224X, D225X, Q226X, P227X, G228X, H229X, D230X, R231X, H233X, K234X, L235X, R236X, L238X, I239X, D240X, L242X, S243X, E244X, R244X, F246X, A247X, A248X, V249X, Y250X, T251X, P252X, C253X, Q254X, N255X, I256X, C257X, I258X, D259X, E260X, S261X, L262X, L263X, L264X, F265X, K266X, G267X, R268X, L269X, Q270X, F271X, R272X, Q273X, Y274X, I275X, P276X, S277X, K278X, R279X, A280X, R281X, Y282X, G283X, I284X, K285X, F286X, Y287X, K288X, L289X, C290X, E291X, S292X, S293X, S294X, G295X, Y296X, T297X, S298X, Y299X, F300X, I302X, E304X, G305X, K306X, D307X, S308X, K309X, L310X, D311X, P312X, P313X, G314X, C315X, P316X, P317X, D318X, L319X, T320X, V321X, S322X, G323X, K324X, I325X, V326X, W327X, E328X, L329X, I330X, S331X, P332X, L333X, L334X, G335X, Q336X, F338X, H339X, L340X, V342X, N344X, F345X, Y346X, S347X, S348X, I349X, L351X, T353X, A354X, Y356X, C357X, L358X, D359X, T360X, P361X, A362X, C363X, G364X, I366X, N367X, R368X, D369X, K371X, G372X, L373X, R375X, A376X, L377X, L378X, D379X, K380X, K381X, L382X, N383X, R384X G385X, T387X, Y388X, A389X, L390X, K392X, N393X, E394X, A397X, K399X, F400X, F401X, D402X, N405X, L406X, L409X, R422X, Y423X, G424X, E425X, P426X, K428X, N429X, K430X, P431X, L432X, S434X, K435X, E436X, S438X, K439X, Y440X, G442X, G443X, V444X, R446X, T447X, L450X, Q451X, H452X, N455X, T457X, R458X, T460X, R461X, A462X, Y464X, K465X, V467X, G468X, I469X, L471X, I472X, Q473X, M474X, L476X, R477X, N478X, S479X, Y480X, V482XY483X, K484X, A485X, A486X, V487X, P488X, G489X, P490X, K491X, L492X, S493X, Y494X, Y495X, K496X, Q498X, L499X, Q500X, I501X, L502X, P503X, A504X, L505X, L506X, F507X, G508X, G509X, V510X, E511X, E512X, Q513X, T514X, V515X, E517X, M518X, P519X, P520X, S521X, D522X, N523X, V524X, A525X, L527X, I528X, G529X, K530X, F532X, I533X, D534X, T535X, L536X, P537X, P538X, T539X, P540X, G541X, F542X, Q543X, R544X, P545X, Q546X, K547X, G548X, C549X, K550X, V551 X, C552X, R553X, K554X, R555X, G556X, 1557X, R558X, R559X, D560X, T561X, R562X, Y563X, Y564X, C565X, P566X,

K567X, C568X, P569X, R570X, N571X, P572X, G573X, L574X, C575X, F576X, K577X, P578X, C579X, F580X, E581X, I582X, Y583X, H584X, T585X, Q586X, L587X, H588X or Y589X (relative to SEQ ID NO: 14517). A list of excision competent, integration deficient amino acid substitutions can be found in U.S. Pat. No. 10,041,077, the contents of which are incorporated by reference in their entirety. (217) In certain embodiments, the piggyBac or piggyBac-like transposase is fused to a nuclear localization signal. In certain embodiments, SEQ ID NO: 14517 or SEQ ID NO: 14518 is fused to a nuclear localization signal. In certain embodiments, the amino acid sequence of the piggyBac or piggyBac like transposase fused to a nuclear localization signal is encoded by a polynucleotide sequence comprising: (218) TABLE-US-00071 (SEQ ID NO: 14626) 1 atggcaccca aaaagaaacg taaagtgatg gccaaaagat ttcacagcgc cgaagaagca 61 gcagcacatt gcatggcatc gtcatccgaa gaattctcgg cgaatatgtc 121 ccaccggcct cggaaagcga ttcgagcact gaggagtcgt ggcgttcctc ggagcgattc ctcaactgtc 181 tcggctcttg aggagccgac ggaagtggat gaggatgtgg acgacttgga ggaccaggaa 241 gccggagaca gggccgacgc tgccgcggga ggggagccgg cgcggggacc tccatgcaat 301 tttcctcccg aaatcccacc gttcactact gtgccgggag tgaaggtcga cacgtccaac 361 ttcgaaccga tcaatttctc tcaactcttc atgactgaag cgatcctgca agatatggtg 421 ctctacacta atgtgtacgc cgagcagtac accegetgee tegetaegeg 481 agagcgcatg cgtggcaccc gaccgatatc gcggagatga ctgactcaaa 541 ctcgcaatgg gcctgatcaa ggccaacagc ctcgagtcat accgggatac gggactgacc agcggttcgt cacgactgtg 601 cttagcattc cggtgttctc cgctaccatg tcccgtaacc gccaccaact cctgctgcgg 661 ttcctccact tcaacaacaa tgcgaccgct gtgccacctg accagccagg acacgacaga 721 ctccacaagc tgcggccatc gatcgactcg ctgagcgagc gactcgccgc ggtgtacacc 781 ccttgccaaa acatttgcaa cgacgagtcg cttctgctgt ttaaaggccg gcttcagttc 841 cgccagtaca tcccatcgaa gcgcgctcgc tatggtatca aattctacaa actctgcgag 901 tcgtccagcg gctacacgtc atacttcttg atctacgagg ggaaggactc taagctggac 951 ccaccggggt gtccaccgga tcttactgtc tccggaaaaa tcgtgtggga actcatctca 1021 cctctcctcg gacaaggctc tcatctctac gtcgacaatt tccactcatc gatccctctg 1081 ttcaccgccc tctactgccc ggatactcca gcctgtggga ccattaacag aaaccggaag 1141 ggtctgccga gagcactgcc ggataagaag ttgaacaggg gagagactta cgcgctgaga 1201 aagaacgaac tcctcgccat caaattette gacaagaaaa atgtgtttat geteacetee 1321 etgtgeteta aggaataete caagtacatg accggaccga tcagctgcag 1381 cattactaca acgccactag aaagacccgg gcctggtaca agaaagtcgg catctacctg 1441 atccaaatgg cactgaggaa ttcgtatatt gtctacaagg ctgccgttcc gggcccgaaa 1501 ctgtcatact acaagtacca gcttcaaatc ctgccggcgc tgctgttcgg tggagtggaa 1561 gaacagactg tgcccgagat gccgccatcc gacaacgtgg cccggttgat cggaaagcac 1621 ttcattgata ccctgcctcc gacgcctgga aagcagcggc cacagaaggg atgcaaagtt 1681 tgccgcaagc gcggaatacg gcgcgatacc cgctactatt gcccgaagtg cccccgcaat 1741 cccggactgt gtttcaagcc ctgttttgaa atctaccaca cccagttgca ttac.

- (219) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Xenopus tropicalis*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (220) TABLE-US-00072 (SEQ ID NO: 14519) 1 ttaacctttt tactgccaat gacgcatggg atacgtcgtg gcagtaaaag ggcttaaatg 61 ccaacgacgc gtcccatacg ttgttggcat tttaagtctt ctatctgcag cggcagcatg 121 tgccgccgct gcagagagtt tctagcgatg acagcccctc tgggcaacga gccggggggg 181 ctgt. (221) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (222) TABLE-US-00073 (SEQ ID NO: 14520) 1 tttgcatttt tagacattta gaagcctata tcttgttaca gaattggaat tacacaaaaa 61 ttctaccata ttttgaaagc ttaggttgtt ctgaaaaaaa caatatattg ttttcctggg 121 taaactaaaa gtcccctcga ggaaaggccc ctaaagtgaa acagtgcaaa acgttcaaaa 181 actgtctggc aatacaagtt ccactttgac caaaacggct ggcagtaaaa gggttaa.
- (223) In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14519 and SEQ ID NO: 14520. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (224) TABLE-US-00074 (SEQ ID NO: 14521) 1 ttaacccttt gcctgccaat cacgcatggg atacgtcgtg gcagtaaaag ggcttaaatg 61 ccaacgacgc gtcccatacg ttgttggcat tttaagtctt ctctctgcag cggcagcatg 121 tgccgccgct gcagagagtt tctagcgatg acagcccctc tgggcaacga gccggggggg 181 ctgtc. (225) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (226) TABLE-US-00075 (SEQ ID NO: 14522) 1 tttgcatttt tagacattta gaagcctata

tcttgttaca gaattggaat tacacaaaaa 61 ttctaccata ttttgaaagc ttaggttgtt ctgaaaaaaa caatatattg ttttcctggg 121 taaactaaaa gtcccctcga ggaaaggccc ctaaagtgaa acagtgcaaa acgttcaaaa 181 actgtctggc aatacaagtt ccactttggg acaaatcggc tggcagtgaa agggttaa. (227) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (228) TABLE-US-00076 (SEQ ID NO: 14523) 1 ttaacctttt tactgccaat gacgcatggg atacgtcgtg gcagtaaaag ggcttaaatg 61 ccaacgacgc gtcccatacg ttgttggcat tttaattctt ctctctgcag cggcagcatg 121 tgccgccgct gcagagagtt tctagcgatg acagcccctc tgggcaacga gccggggggg 181 ctgtc. (229) In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14520 and SEQ ID NO: 14519, SEQ ID NO: 14521 or SEQ ID NO: 14523. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14522 and SEQ ID NO: 14519, SEQ ID NO: 14521 or SEQ ID NO: 14523. In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end comprising at least 14, 16, 18, 20, 30 or 40 contiguous nucleotides from SEQ ID NO: 14519, SEQ ID NO: 14521 or SEQ ID NO: 14523. In certain embodiments, the piggy Bac or piggyBaclike transposon comprises one end comprising at least 14, 16, 18, 20, 30 or 40 contiguous nucleotides from SEQ ID NO: 14520 or SEQ ID NO: 14522. In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end with at least 90% identity to SEQ ID NO: 14519, SEQ ID NO: 14521 or SEQ ID NO: 14523. In certain embodiments, the piggyBac or piggyBac-like transposon comprises one end with at least 90% identity to SEQ ID NO: 14520 or SEQ ID NO: 14522. In one embodiment, one transposon end is at least 90% identical to SEQ ID NO: 14519 and the other transposon end is at least 90% identical to SEQ ID NO: 14520.

(230) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of TTAACCTTTTTACTGCCA (SEQ ID NO: 14524). In certain embodiments, the piggyBac or piggyBaclike transposon comprises a sequence of TTAACCCTTTGCCTGCCA (SEQ ID NO: 14526). In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of TTAACCYTTTTACTGCCA (SEQ ID NO: 14527). In certain embodiments, the piggyBac or piggyBaclike transposon comprises a sequence of TGGCAGTAAAAGGGTTAA (SEQ ID NO: 14529). In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of TGGCAGTGAAAGGGTTAA (SEQ ID NO: 14531). In certain embodiments, the piggyBac or piggyBaclike transposon comprises a sequence of TTAACCYTITKMCTGCCA (SEQ ID NO: 14533). In certain embodiments, one end of the piggyBac or piggyBac-like transposon comprises a sequence selected from SEQ ID NO: 14524, SEQ ID NO: 14526 and SEQ ID NO: 14527. In certain embodiments, one end of the piggyBac<sup>™</sup> (PB) or piggyBac-like transposon comprises a sequence selected from SEQ ID NO: 14529 and SEQ ID NO: 14531. In certain embodiments, each inverted terminal repeat of the piggyBac or piggyBac-like transposon comprises a sequence of ITR sequence of CCYTTTKMCTGCCA (SEQ ID NO: 14563). In certain embodiments, each end of the piggyBac<sup>™</sup> (PB) or piggyBac-like transposon comprises SEQ ID NO: 14563 in inverted orientations. In certain embodiments, one ITR of the piggyBac or piggyBac-like transposon comprises a sequence selected from SEQ ID NO: 14524, SEQ ID NO: 14526 and SEQ ID NO: 14527. In certain embodiments, one ITR of the piggyBac or piggyBac-like transposon comprises a sequence selected from SEQ ID NO: 14529 and SEQ ID NO: 14531. In certain embodiments, the piggyBac or piggyBac like transposon comprises SEQ ID NO: 14533 in inverted orientation in the two transposon ends.

(231) In certain embodiments. The piggyBac or piggyBac-like transposon may have ends comprising SEQ ID NO: 14519 and SEQ ID NO: 14520 or a variant of either or both of these having at least 90% sequence identity to SEQ ID NO: 14519 or SEQ ID NO: 14520, and the piggyBac or piggyBac-like transposase has the sequence of SEQ ID NO: 14517 or a variant showing at least %, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between sequence identity to SEQ ID NO: 14517 or SEQ ID NO: 14518. In certain embodiments, one piggyBac or piggyBac-like transposon end comprises at least 14 contiguous nucleotides from SEQ ID NO: 14523, and the other transposon end comprises at least 14 contiguous nucleotides from SEQ ID NO: 14520 or SEQ ID NO: 14522 In certain embodiments, one transposon end comprises at least 15, at least 16, at least 17, at least 18, at least 20, at least 22, at least 25, at least 30 contiguous nucleotides from SEQ ID NO: 14519, SEQ ID NO: 14521 or SEQ ID NO: 14523, and the other transposon end comprises at least 15, at least 15, at least 15, at least 16, at least 17, at least 18, at least 19, at least 19, at least 20, at

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least 22, at least 25 or at least 30 contiguous nucleotides from SEQ ID NO: 14520 or SEQ ID NO: 14522.
(232) In certain embodiments, the piggyBac or piggyBac-like transposase recognizes a transposon end
with a left sequence corresponding to SEQ ID NO: 14519, and a right sequence corresponding to SEQ ID
NO: 14520. It will excise the transposon from one DNA molecule by cutting the DNA at the 5'-TTAA-3'
sequence at the left end of one transposon end to the 5'-TTAA-3' at the right end of the second transposon
end, including any heterologous DNA that is placed between them, and insert the excised sequence into a
second DNA molecule. In certain embodiments, truncated and modified versions of the left and right
transposon ends will also function as part of a transposon that can be transposed by the piggyBac or
piggyBac-like transposase. For example, the left transposon end can be replaced by a sequence
corresponding to SEQ ID NO: 14521 or SEQ ID NO: 14523, the right transposon end can be replaced by a
shorter sequence corresponding to SEQ ID NO: 14522. In certain embodiments, the left and right
transposon ends share an 18 bp almost perfectly repeated sequence at their ends (5'-
TTAACCYTITKMCTGCCA: SEQ ID NO: 14533) that includes the 5'-TTAA-3' insertion site, which
sequence is inverted in the orientation in the two ends. That is in SEQ ID NO: 14519 and SEQ ID NO:
14523 the left transposon end begins with the sequence 5'-TTAACCTTTITACTGCCA-3' (SEQ ID NO:
14524), or in SEQ ID NO: 14521 the left transposon end begins with the sequence 5'-
TTAACCCTTTGCCTGCCA-3' (SEQ ID NO: 14526); the right transposon ends with approximately the
reverse complement of this sequence: in SEQ ID NO: 14520 it ends 5' TGGCAGTAAAAGGGTTAA-3'
(SEQ ID NO: 14529), in SEQ ID NO: 14522 it ends 5'-TGGCAGTGAAAGGGTTAA-3' (SEQ ID NO:
14531.) One embodiment of the invention is a transposon that comprises a heterologous polynucleotide
inserted between two transposon ends each comprising SEQ ID NO: 14533 in inverted orientations in the
two transposon ends. In certain embodiments, one transposon end comprises a sequence selected from
SEQ ID NOS: 14524, SEQ ID NO: 14526 and SEQ ID NO: 14527. In some embodiments, one transposon
end comprises a sequence selected from SEQ ID NO: 14529 and SEQ ID NO: 14531.
(233) In certain embodiments, the piggyBac<sup>™</sup> (PB) or piggyBac-like transposon is isolated or derived
from Xenopus tropicalis. In certain embodiments, the piggyBac or piggyBac-like transposon comprises at
a sequence of:
                                                   1 ccctttgcct gccaatcacg
(234) TABLE-US-00077 (SEQ ID NO: 14573)
                                                                            catgggatac
gtcgtggcag taaaagggct taaatgccaa 61 cgacgcgtcc catacgtt.
(235) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(236) TABLE-US-00078 (SEQ ID NO: 14574)
                                                   1 cctgggtaaa
                                                                 ctaaaagtcc cctcgaggaa
aggecectaa agtgaaacag tgeaaaacgt 61 teaaaaactg tetggeaata caagtteeac tttgggacaa
agtgaaaggg.
(237) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at least 16 contiguous
bases from SEQ ID NO: 14573 or SEQ ID NO: 14574, and inverted terminal repeat of
CCYTTTBMCTGCCA (SEQ ID NO: 14575).
(238) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(239) TABLE-US-00079 (SEQ ID NO: 14579)
                                                      1 ccctttgcct gccaatcacg catgggatac
gtcgtggcag taaaagggct taaatgccaa
                                   61 cgacgcgtcc catacgttgt tggcatttta agtcttctct ctgcagcggc
agcatgtgcc 121 gccgctgcag agagtttcta gcgatgacag cccctctggg caacgagccg ggggggctgt 181 c.
(240) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(241) TABLE-US-00080 (SEQ ID NO: 14580)
                                                      1 cctttttact gccaatgacg catgggatac
gtcgtggcag taaaagggct taaatgccaa
                                     61 cgacgcgtcc catacgttgt tggcatttta attcttctct ctgcagcggc
agcatgtgcc 121 gccgctgcag agagtttcta gcgatgacag cccctctggg caacgagccg ggggggctgt 181 c.
(242) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(243) TABLE-US-00081 (SEQ ID NO: 14581)
                                                      1 cctttttact gccaatgacg catgggatac
gtcgtggcag taaaagggct taaatgccaa
                                                    catacgttgt tggcatttta agtcttctct ctgcagcggc
                                     61 cgacgcgtcc
agcatgtgcc 121 gccgctgcag agagtttcta gcgatgacag cccctctggg caacgagccg ggggggctgt 181 c.
(244) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
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gtcgtggcag taaaagggct taaatgccaa 61 cgacgcgtcc catacgttgt tggcatttta agtcttctct ctgcagcggc

(246) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:

1 cctttttact gccaatgacg catgggatac

(245) TABLE-US-00082 (SEQ ID NO: 14582)

agcatgtgcc 121 gccgctgcag

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(247) TABLE-US-00083 (SEQ ID NO: 14583)
                                                  1 cctttttact gccaatgacg catgggatac
gtcgtggcag taaaagggct taaatgccaa 61 cgacgcgtcc catacgttgt tggcatttta agtctt.
(248) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(249) TABLE-US-00084 (SEQ ID NO: 14584)
                                                  1 ccctttgcct gccaatcacg catgggatac
gtcgtggcag taaaagggct taaatgccaa 61 cgacgcgtcc catacgttgt tggcatttta agtctt .
(250) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(251) TABLE-US-00085 (SEQ ID NO: 14585)
                                                     1 ttatcctttt tactgccaat gacgcatggg
atacgtcgtg gcagtaaaag ggcttaaatg
                                   61 ccaacgacgc gtcccatacg ttgttggcat tttaagtctt ctctctgcag
cggcagcatg 121 tgccgccgct gcagagagtt tctagcgatg acagcccctc tgggcaacga gccggggggg 131 ctgtc.
(252) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of;
(253) TABLE-US-00086 (SEQ ID NO: 14586)
                                                     1 tttgcatttt tagacattta gaagcctata
tcttgttaca gaattggaat tacacaaaaa
                                  61 ttctaccata ttttgaaagc ttaggttgtt ctgaaaaaaa caatatattg
ttttcctggg 121 taaactaaaa gtcccctcga ggaaaggccc ctaaagtgaa acagtgcaaa acgttcaaaa 161
actgtctggc aatacaagtt ccactttggg acaaatcggc tggcagtgaa aggg.
In certain embodiments, the piggyBac or piggyBac-like transposon comprises a left transposon end
sequence selected from SEQ ID NO: 14573 and SEQ ID NOs: 14579-14585. In certain embodiments, the
left transposon end sequence is preceded by a left target sequence. In certain embodiments, the piggyBac
or piggyBac-like transposon comprises at a sequence of:
(254) TABLE-US-00087 (SEQ ID NO: 14587)
                                                     1 tttgcatttt tagacattta gaagcctata
tcttgttaca gaattggaat tacacaaaaa
                                  61 ttctaccata ttttgaaagc ttaggttgtt ctgaaaaaaa caatatattg
ttttcctggg 121 taaactaaaa gtcccctcga ggaaaggccc ctaaagtgaa acagtgcaaa acgttcaaaa 181
actgtctggc aatacaagtt ccactttgac caaaacggct ggcagtaaaa ggg.
(255) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(256) TABLE-US-00088 (SEQ ID NO: 14588)
                                                    1 ttgttctgaa aaaaacaata tattgttttc
ctgggtaaac taaaagtccc ctcgaggaaa
                                    61 ggcccctaaa gtgaaacagt gcaaaacgtt caaaaactgt
           aagttccact 121 ttgaccaaaa cggctggcag taaaaggg.
(257) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(258) TABLE-US-00089 (SEQ ID NO: 14589)
                                                     1 tttgcatttt tagacattta gaagcctata
tcttgttaca gaattggaat tacacaaaaa
                                  61 ttctaccata ttttgaaagc ttaggttgtt ctgaaaaaaa caatatattg
ttttcctggg 121 taaactaaaa gtcgcctcga ggaaaggccc ctaaagtgaa acagtgcaaa acgttcaaaa 181
actgtctggc aatacaagtt ccactttgac caaaacggct ggcagtaaaa gggttat.
(259) In certain embodiments, the piggyBac or piggyBac-like transposon comprises at a sequence of:
(260) TABLE-US-00090 (SEQ ID NO: 14590)
                                                     1 ttgttctgaa aaaaacaata tattgttttc
ctgggtaaac taaaagtccc ctcgaggaaa
                                    61 ggcccctaaa gtgaaacagt gcaaaacgtt caaaaactgt
           aagttccact 121 ttgggacaaa tcggctggca gtgaaaggg.
(261) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a right transposon end
sequence selected from SEQ ID NO: 14574 and SEQ ID NOs: 14587-14590. In certain embodiments, the
right transposon end sequence is followed by a right target sequence. In certain embodiments, the left and
right transposon ends share a 14 repeated sequence inverted in orientation in the two ends (SEQ ID NO:
14575) adjacent to the target sequence. In certain embodiments, the piggyBac or piggyBac-like transposon
comprises a left transposon end comprising a target sequence and a sequence that is selected from SEQ ID
NOs: 14582-14584 and 14573, and a right transposon end comprising a sequence selected from SEQ ID
NOs: 14588-14590 and 14574 followed by a right target sequence.
(262) In certain embodiments, the left transposon end of the piggyBac or piggyBac-like transposon
comprises
(263) TABLE-US-00091 (SEQ ID NO: 14591)
                                                  1 atcacgcatg ggatacgtcg tggcagtaaa
agggettaaa tgccaacgac gegteecata 61 egtt,
and an ITR. In certain embodiments, the left transposon end comprises
(264) TABLE-US-00092 (SEQ ID NO: 14592)
                                                  1 atgacgcatg ggatacgtcg
                                                                            tggcagtaaa
agggettaaa tgecaaegae gegteecata 61 egttgttgge attttaagte tt
and an ITR In certain embodiments, the right transposon end of the piggyBac or piggyBac-like transposon
comprises
(265) TABLE-US-00093 (SEQ ID NO: 14593)
                                                  1 cctgggtaaa ctaaaagtcc cctcgaggaa
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- aggcccctaa agtgaaacag tgcaaaacgt 61 tcaaaaactg tctggcaata caagttccac tttgggacaa atcggc and an ITR. In certain embodiments, the right transposon end comprises (266) TABLE-US-00094 (SEQ ID NO: 14594) 1 ttgttctgaa aaaaacaata tattgttttc
- (266) TABLE-US-00094 (SEQ ID NO: 14594) 1 ttgttctgaa aaaaacaata tattgttttc ctgggtaaac taaaagtccc ctcgaggaaa 61 ggcccctaaa gtgaaacagt gcaaaacgtt caaaaactgt ctggcaatac aagttccact 121 ttgaccaaaa cggc and an ITR.
- (267) In certain embodiments, one transposon end comprises a sequence that is at least 90%, at least 95%, at least 99% or any percentage in between identical to SEQ ID NO: 14573 and the other transposon end comprises a sequence that is at least 90%, at least 95%, at least 99% or any percentage in between identical to SEQ ID NO: 14574. In certain embodiments, one transposon end comprises at least 14, at least 16, at least 18, at least 20 or at least 25 contiguous nucleotides from SEQ ID NO: 14573 and one transposon end comprises at least 14, at least 16, at least 20 or at least 25 contiguous nucleotides from SEQ ID NO: 14574. In certain embodiments, one transposon end comprises at least 14, at least 16, at least 18, at least 20 from SEQ ID NO: 14591, and the other end comprises at least 14, at least 16, at least 18, at least 20 from SEQ ID NO: 14593. In certain embodiments, each transposon end comprises SEQ ID NO: 14575 in inverted orientations.
- (268) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence selected from of SEQ ID NO: 14573, SEQ ID NO: 14579, SEQ ID NO: 14581, SEQ ID NO: 14582, SEQ ID NO: 14583, and SEQ ID NO: 14588, and a sequence selected from SEQ ID NO: 14587, SEQ ID NO: 14588, SEQ ID NO: 14589 and SEQ ID NO: 14586 and the piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14517 or SEQ ID NO: 14518.
- (269) In certain embodiments, the piggyBac or piggyBac-like transposon comprises ITRs of CCCTITGCCTGCCA (SEQ ID NO: 14622) (left ITR) and TGGCAGTGAAAGGG (SEQ ID NO: 14623) (right ITR) adjacent to the target sequences.
- (270) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Helicoverpa armigera*. The piggyBac or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30% 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (271) TABLE-US-00095 (SEQ ID NO: 14525) 1 MASRQRLNHD EIATILENDD DYSPLDSESE KEDCVVEDDV WSDNEDAIVD FVEDTSAQED 61 PDNNIASRES PNLEVTSLTS HRIITLPQRS IRGKNNHVWS TTKGRTTGRT SAINIIRTNR 121 GPTRMCRNIV DPLLCFQLFI TDEIIHEIVK WTNVEIIVKR QNLKDISASY RDTNTMEIWA 181 LVGILTLTAV MKDNHLSTDE LFDATFSGTR YVSVMSRERF EFLIRCIRMD DKTLRPTLRS 241 DDAFLPVRKI WEIFINQCRQ NHVPGSNLTV DEQLLGFRGR CPFRMYIPNK PDKYGIKFPM 301 MCAAATKYMI DAIPYLGKST KTNGLPLGEF YVKDLTKTVH GTNRNITCDN WFTSIPLAKN 361 MLQAPYNLTI VGTIRSNKRE MPEEIKNSRS RPVGSSMFCF DGPLTLVSYK PKPSKMVFLL 421 SSCDENAVIN ESNGKPDMIL FYNQTKGGVD SFDQMCKSMS ANRKTNRWPM AVFYGMLNMA 481 FVNSYIIYCH NKINKQEKPI SRKEFMKKLS IQLTTPWMQE RLQAPTLKRT LRDNITNVLK 541 NVVPASSENI SNEPEPKKRR YCGVCSYKKR RMTKAQCCKC KKAICGEHNI DVCQDCI.
- (272) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Helicoverpa armigera*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (273) TABLE-US-00096 (SEQ ID NO: 14570) 1 ttaaccctag aagcccaatc tacgtaaatt tgacgtatac cgcggcgaaa tatctctgtc 61 tctttcatgt ttaccgtcgg atcgccgcta acttctgaac caactcagta gccattggga 121 cctcgcagga cacagttgcg tcatctcggt aagtgccgcc atcttgttgt actctctatt 161 acaacacacg tcacgtcacg tcgttgcacg tcattttgac gtataattgg gctttgtgta 241 acttttgaat ttgtttcaaa ttttttatgt ttgtgattta tttgagttaa tcgtattgtt 301 tcgttacatt tttcatataa taataatatt ttcaggttga gtacaaa. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (274) TABLE-US-00097 (SEQ ID NO: 14528) 1 agactgtttt tttgtaagag acttctaaaa

- (275) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Pectinophora gossypiella*. The piggyBac or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (276) TABLE-US-00098 (SEQ ID NO: 14530) 1 MDLRKQDEKI RQWLEQDIEE DSKGESDNSS SETEDIVEME VHKNTSSESE VSSESDYEPV 61 CPSKRQRTQI IESEESDNSE SIRPSRRQTS RVIDSDETDE DVMSSTPQNI PRNPNVIQPS 121 SRFLYGKNKH KWSSAAKPSS VRTSRRNIIH FIPGPKERAR EVSEPIDIFS LFISEDMLQQ 181 VVTFTNAEML IRKNKYKTET FTVSPTNLEE IRALLGLLFN AAAMKSNHLP TRMLFNTHRS 241 GTIFKACMSA ERLNFLIKCL RFDDKLTRNV RQRDDRFAPI RDLWQALISN FQKWYTPGSY 301 ITVDEQLVGF RGRCSFRMYI PNKPNKYGIK LVMAADVNSK YIVNAIPYLG KGTDPQNQPL 361 ATFFIKEITS TLHGTNRNIT MDNWFTSVPL ANELLMAPYN LTLVGTLRSN KREIPEKLKN 421 SKSRAIGTSM FCYDGDKTLV SYKAKSNKVV FILSTIHDQP DINQETGKPE MIHFYNSTKG 481 AVDTVDQMCS SISTNRKTQR WPLCVFYNML NLSIINAYVV YVYNNVRNNK KPMSRRDFVI 541 KLGDQLMEPW LRQRLQTVTL RRDIKVMIQD ILGESSDLEA PVPSVSNVRK IYYLCPSKAR 601 RMTKHRCIKC KQAICGPHNI DICSRCIE. (277) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from Pectinophora gossypiella. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (278) TABLE-US-00099 (SEQ ID NO: 14532) 1 ttaaccctag ataactaaac attcgtccgc tcgacgacgc gctatgccgc gaaattgaag 61 tttacctatt attccgcgtc ccccgccccc gccgcttttt ctagcttcct gatttgcaaa 121 atagtgcatc gcgtgacacg ctcgaggtca cacgacaatt aggtcgaaag ttacaggaat 181 ttcgtcgtcc gctcgacgaa agtttagtaa ttacgtaagt ttggcaaagg taagtgaatg 241 aagtattttt ttataattat tttttaattc tttatagtga taacgtaagg tttatttaaa 301 tttattactt ttatagttac ttagccaatt gttataaatt ccttgttatt gctgaaaaat 361 ttgcctgttt tagtcaaaat ttattaactt ttcgatcgtt ttttag.
- (280) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Ctenoplusia agnata*. The piggyBac or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (281) TABLE-US-00101 (SEQ ID NO: 14534) 1 MASRQHLYQD EIAAILENED DYSPHDTDSE MEDCVTQDDV RSDVEDEMVD NIGNGTSPAS 61 RHEDPETPDP SSEASNLEVT LSSHRIIILP QRSIREKNNH IWSTTKGQSS GRTAAINIVR 121 TNRGPTRMCR NIVDPLLCFQ LFIKEEIVEE IVKWTNVEMV QKRVNLKDIS ASYRDTNEME 181 IWAIISMLTL SAVMKDNHLS TDELFNVSYG TRYVSVMSRE RFEFLLRLLR MGDKLLRPNL 241 RQEDAFTPVR KIWEIFINQC RLNYVPGTNL TVDEQLLGFR GRCPFRMYIP NKPDKYGIKF 301 PMVCDAATKY MVDAIPYLGK STKTQGLPLG EFYVKELTQT VHGTNRNVTC DNWFTSVPLA 361 KSLLNSPYNL TLVGTIRSNK REIPEEVKNS RSRQVGSSMF CFDGPLTLVS YKPKPSKMVF 421 LLSSCNEDAV VNQSNGKPDM ILFYNQTKGG VDSFDQMCSS MSTNRKTNRW

- PMAVFYGMLN 481 MAFVNSYIIY CHNMLAKKEK PLSRKDFMKK LSTDLTTPSM QKRLEAPTLK RSLPDNITNV 541 LKIVPQAAID TSFDEPEPKK RRYCGFCSYK KKRMTKTQCF KCKKPVCGEH NIDVCQDCI.
- (282) In certain embodiments, the piggy Bac or piggyBac-like transposon is isolated or derived from *Ctenoplusia agnata*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (283) TABLE-US-00102 (SEQ ID NO: 14535) 1 ttaaccctag aagcccaatc tacgtcattc tgacgtgtat gtcgccgaaa atactctgtc 61 tctttctcct gcacgatcgg attgccgcga acgctcgatt caacccagtt ggcgccgaga 121 tctattggag gactgcggcg ttgattcggt aagtcccgcc attttgtcat agtaacagta 181 ttgcacgtca gcttgacgta tatttgggct ttgtgttatt tttgtaaatt ttcaacgtta 241 gtttattatt gcatcttttt gttacattac tggtttattt gcatgtatta ctcaaatatt 301 atttttattt tagcgtagaa aataca.
- (284) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of (285) TABLE-US-00103 (SEQ ID NO: 14536) 1 agactgtttt ttttgtattt gcattatata ttatattcta aagttgattt aattctaaga 61 aaaacattaa aataagtttc tttttgtaaa atttaattaa ttataagaaa aagtttaagt 121 tgatctcatt ttttataaaa atttgcaatg tttccaaagt tattattgta aaagaataaa 181 taaaagtaaa ctgagtttta attgatgttt tattatatca ttatactaa tattacttaa 241 ataaaacaat aactgaatgt atttctaaaa ggaatcacta gaaaatatag tgatcaaaaa 301 tttacacgtc atttttgcgt atgattgggc tttataggtt ctaaaaatat gattgggcct 361 ctagggttaa.
- (286) In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of CCCTAGAAGCCCAATC (SEQ ID NO: 14564).
- (287) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Agrotis ipsilon*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (288) TABLE-US-00104 (SEQ ID NO: 14537) 1 MESPQRLNQD EIATILENDD DYSPLDSDSE AEDRVVEDDV WSDNEDAMID YVEDTSRQED 61 PDNNIASQES ANLEVTSLTS HRIISLPQRS ICGKNNHVWS TTKGRTTGRT SAINIIRTNR 121 GPTRMCRNIV DPLICFQLFI TDEIIHEIVK WTNVEMIVKR QNLIDISASY RDTNTMEMWA 181 LVGILTLTAV MKDNHLSTDE LFDATFSGTR YVSVMSREPF EFLIRCMRMD DKTLRPTLRS 241 DDAFIPVRKL WEIFINQCRL NYVPGGNLTV DEQLLGFRGR CPFRMYIPNK PDKYGIRFPM 301 MCDAATKYMI DAIPYLGKST KTNGLPLGEF YVKELTKTVH GTNRNVTCDN WFTSIPLAKN 361 MLQAPYNLTI VGTIRSNKRE IPEEIKNSRS RPVGSSMFCF DGPLTLVSYK PKPSRMVFLL 421 SSCDENAVIN ESNGKPDMIL FYNQTKGGVD SFDQMCKSMS ANRKTNRWPM AVFYGMLNMA 481 FVNSYIIYCH NKINKQKKPI NRKEFMKNLS TDLTTPWMQE RLKAPTLKRT LRDNITNVLK 541 NVVPPSPANN SEEPGRKKRS YCGFCSYKKR RMTKTQFYKC KKAICGEHNT DVCQDCV.
- (289) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Agrotis ipsilon*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (290) TABLE-US-00105 (SEQ ID NO: 14538) 1 ttaaccctag aagcccaatc tacgtaaatt tgacgtatac cgcggcgaaa tatatctgtc 61 tctttcacgt ttaccgtcgg attcccgcta acttcggaac caactcagta gccattgaga 121 actcccagga cacagttgcg tcatctcggt aagtgccgcc attttgttgt aatagacagg 181 ttgcacgtca ttttgacgta taattgggct ttgtgtaact tttgaaatta tttataattt 241 ttattgatgt gatttatttg agttaatcgt attgtttcgt tacatttttc atatgatatt 301 aatattttca gattgaatat aaa.
- In certain embodiments, the piggyBac or piggy Bac-like transposon comprises a sequence of: (291) TABLE-US-00106 (SEQ ID NO: 14539) 1 agactgtttt ttttaaaagg cttataaagt attactattg cgtgatttaa ttttataaaa 61 atatttaaaa ccagttgatt tttttaataa ttacctaatt ttaagaaaaa atgttagaag 121 cttgatattt ttagttgattt ttttctaaga tttgattaaa aggccataat tgtattaata 181 aagagtattt ttaacttcaa attatttta tttattaatt aaaacttcaa ttatgataat 241 acatgcaaaa atatagttca tcaacagaaa aatataggaa aactctaata gttttatttt 301 tacacgtcat ttttacgtat gattgggctt tatagctagt caaatatgat tgggcttcta 351 gggttaa.

- (292) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Megachile rotundata*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (293) TABLE-US-00107 (SEQ ID NO: 14540) 1 MNGKDSLGEF YLDDLSDCLD CRSASSTDDE SDSSNIAIRK RCRIPLIYSD SEDEDMNNNV 61 EDNNHFVKES NRYHYQIVEK YKITSKTKKW KDVTVTEMKK FLGLIILMGQ VKKDVLYDYW 121 STDPSIETPF FSKVMSRNRF LQIMQSWHFY NNNDISPNSH RLVKIQPVID YFKEKFNNVY 181 KSDQQLSLDE CLIPWRGRLS IKTYNPAKIT KYGILVRVLS EARTGYVSNF CVYAADGKKI 241 EETVLSVIGP YKNMWHHVYQ DNYYNSVNIA KIFLKNKLRV CGTIRKNRSL PQILQTVKLS 301 RGQHQFLRNG HTLLEVWNNG KRNVNMISTI HSAQMAESRN RSRTSDCPIQ KPISIIDYNK 361 YMKGVDRADQ YLSYYSIFRK TKKWTKRVVM FFINCALFNS FKVYTTLNGQ KITYKNFLHK 421 AALSLIEDCG TEEQGTDLPN SEPTTTRTTS RVDHPGRIEN FGKHKLVNIV TSGQCKKPLR 481 QCRVCASKKK LSRTGFACKY CNVPLHKGDC FERYHSLKKY.
- (294) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Megachile rotundata*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (295) TABLE-US-00108 (SEQ ID NO: 14541) 1 ttaaataatg cccactctag atgaacttaa cactttaccg accggccgtc gattattcga 61 cgtttgctcc ccagcgctta ccgaccggcc atcgattatt cgacgtttgc ttcccagcgc 121 ttaccgaccg gtcatcgact tttgatcttt ccgttagatt tggttaggtc agattgacaa 181 gtagcaagca tttcgcattc tttattcaaa taatcggtgc tttttctaa gctttagcocc 241 ttagaa.
- (297) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Bombus impatiens*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (298) TABLE-US-00110 (SEQ ID NO: 14543) 1 MNEKNGIGEF YLDDLSDCPD SYSRSNSGDE SDGSDTIIRK RGSVLPPRYS DSEDDEINNV 61 EDNANNVENN DDIWSTNDEA IILEPFEGSP GLKIMPSSAE SVTDNVNLFF GDDFFEHLVR 121 ESNRYHYQVM EKYKIPSKAK KWTDITVPEM KKFLGLIVLM GQIKKDVLYD YWSTDPSIET 181 PFFSQVMSRN RFVQIMQSWH FCNNDNIPHD SHRLAKIQPV IDYFRRKFND VYKPCQQLSL 241 DESIIPWPGR LSIKTYNPAK ITKYGILVRV LSEAVTGYVC NEFDYAADGK KLEDTAVIEP 301 YKNIWHQIYQ DNYYNSVKMA RILLKNKVRV CGTIRKNRGL PRSLKTIQLS RGQYEFRRNH 361 QILLEVWNNG RRNVNMISTI HSAQLMESRS KSKRSDVPIQ KPNSIIDYNK YMKGVDRADQ 421 YLAYYSIFRK TKKWTKRVVM FFINCALFNS FRVYTILNGK NITYKNFLHK VAVSWIEDGE 481 TNCTEQDDNL PNSEPTRRAP RLDHPGRLSN YGKHKLINIV TSGRSLKPQR QCRVCAVQKK 541 RSRTCFVCKF CNVPLHKGDC FERYHTLKKY. (299) In certain embodiments, the piggyBac or piggy Bac-like transposon comprises a sequence of:
- (300) TABLE-US-00111 (SEQ ID NO: 14544) 1 ttaatttttt aacattttac cgaccgatag ccgattaatc gggtttttgc cgctgacgct 61 taccgaccga taacctatta atcggctttt tgtcgtcgaa gcttaccaac ctatagccta 121 cctatagtta atcggttgcc atggcgataa acaatctttc tcattatatg agcagtaatt 181 tgttatttag tactaaggta

- ccttgctcag ttgcgtcagt tgcgttgctt tgtaagctcc 241 cacagtttta taccaattcg aaaaacttac cgttcgcg. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of (301) TABLE-US-00112 (SEQ ID NO: 14545) 1 actatttcac atttgaacta aaaaccgttg taatagataa aataaatata atttagtatt 61 aatattatgg aaacaaaaga ttttattcaa tttaattatc ctatagtaac aaaaagcggc 121 caattttatc tgagcatacg aaaagcacag atactcccgc ccgacagtct aaaccgaaac 181 agagccggcg ccagggagaa tctgcgcctg agcagccggt cggacgtgcg tttgctgttg 241 aaccgctagt ggtcagtaaa ccagaaccag tcagtaagcc agtaactgat cagttaacta 301 gattgtatag ttcaaattga acttaatcta gtttttaagc gtatgaatgt tgtctaactt 361 cgttatatat tatattcttt ttaa.
- (302) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Mamestra brassicae*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (303) TABLE-US-00113 (SEQ ID NO: 14546) 1 MFSFVPNKEQ TRTVLIFCFH LKTTAAESHR PLVEAFGEQV PTVKTCERWF QRFKSGDFDV 61 DDKEHGKPPK RYEDAELQAL LDEDDAQTQK QLAEQLEVSQ QAVSNRLREG GKIQKVGRWV 121 PHELNERQRE RRKNTCEILL SRYKRKSFLH RIVTGEEKWI FFVNPKRKKS YVDPGQPATS 181 TARPNRFGKK TRLCVWWDQS GVIYYELLKP GETVNTARYQ QQLINLNRAL QRKRPEQKR 241 QHRVIFLHDN APSHTARAVR DTLETLNWEV LPHAAYSPDL APSDYHLFAS MGHALAEQRF 301 DSYESVEEWL DEWFAAKDDE FYWRGIHKLP ERWDNCVASD GKYFE.
- (304) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Mamestra brassicae*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (305) TABLE-US-00114 (SEQ ID NO: 14547) 1 ttattgggtt gcccaaaaag taattgcgga tttttcatat acctgtcttt taaacgtaca 61 tagggatcga actcagtaaa actttgacct tgtgaaataa caaacttgac tgtccaacca 121 ccatagtttg gcgcgaattg agcgtcataa ttgttttgac tttttgcagt caac.
- In certain embodiments, the piggyBac or piggyBac-1e transposon comprises a sequence of: (306) TABLE-US-00115 (SEQ ID NO: 14548) 1 atgattttt ctttttaaac caattttaat tagttaattg atataaaaat ccgcaattac 61 tttttgggca acccaataa.
- (307) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Mayetiola destructor*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65% 70%, 75% 80% 85% 90%, 95% 99% or any percentage in between identical to:
- (308) TABLE-US-00116 (SEQ ID NO: 14549) 1 MENFENWRKR RHLREVLLGH HRLLVEVYGE HALAKTQCFE WFQRFKSGDF FFAKKTAAES 61 DTEDKERPGQ PKKFEDEELE ALLDEDCCQT QEELAKSLGV **TQQAISKRLK** AAGYIQKQGN 121 WVPHELKPRD VERRFCMSEM LLQRHKKKSF LSRIITGDEK WIHYDNSKRK KSYVKRGGRA 181 KSTPKSNLHG AKVMLCIKWD EPGQTITGDL YRTQLIRLKQ ALAEKRPEYA 241 KRHGAVIFHH QRGVLYYELL DNARPHVALP VKNYLENSGW EVLPHPPYSP DLAPSDYHLF RSMQNDLAGK SDGOYFE. 301 RFTSEQGIPK WLDSFLAAKP AKFFEKGIHE LSERWEKVIA (309) In certain embodiments, the piggy ac or piggyBac-like transposon is isolated or derived from Mayetiola destructor. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (310) TABLE-US-00117 (SEQ ID NO: 14550) 1 taagacttcc aaaatttcca cccgaacttt accttccccg cgcattatgt ctctcttttc 61 accctctgat ccctggtatt gttgtcgagc acgatttata ttgggtgtac aacttaaaaa 121 ccggaattgg acgctagatg tccacactaa cgaatagtgt aaaagcacaa atttcatata 181 tacgtcattt tgaaggtaca tttgacagct atcaaaatca gtcaataaaa ctattctatc 241 tgtgtgcatc atatttttt attaact.

- In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of (311) TABLE-US-00118 (seq ID NO: 14551) 1 tgcattcatt cattttgtta tcgaaataaa gcattaattt ccactaaaaa attccggttt 61 ttaagttgta cacccaatat catccttagt gacaattttc aaatggcttt cccattgagc 121 tgaaaccgtg gctatagtaa gaaaaacgcc caacccgtca tcatatgcct tttttttctc 161 aacatccg.
- (312) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Apis mellifera*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (313) TABLE-US-00119 (SEQ ID NO: 14552) 1 MENQKEHYRH ILLFYFRKGK AVYGDEALKE RQCQNWFDKF RSGDFSLKDE NASQAHKKLC 61 KRSGRPVEVD DDLIKAIIDS DRHSTTREIA EKLHVSHTCI ENHLKQLGYV QKLDTWVPHE 121 LKEKHLTQRI NSCDLLKKRN ENDPFLKRLI TGDEKWVVYN NIKRKRSWSR PREPAQTTSK 181 AGIHRKKVLL SVWWDYKGIV INSVVYIEQL TKLNNAVEEK RPELTNRKGV 241 VFHHDNARPH YFELLPPNRT PYSPDLAPSD YFLFRSLQNS LNGKNFNNDD 301 TSLVTRQKLL ELGWDVLPHP DIKSYLIQFF ANKNQKFYER GIMMLPERWQ KVIDQNGQHI TE. (314) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Apis*
- *mellifera*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (315) TABLE-US-00120 (SEQ ID NO: 14553) 1 ttgggttggc aactaagtaa ttgcggattt cactcataga tggcttcagt tgaattttta tttgatagtt ggcaactcag 121 ctgtcaatca gtaaaaaaag ttttttgatc ggtagaaaaaag ttttttgatc

cattttgtta tttgatagtt ggcaactcag 121 ctgtcaatca gtaaaaaaag ttttttgatc ggttgcgtag ttttcgtttg gcgttcgttg 181 aaaa.

- (316) In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (317) TABLE-US-00121 (SEQ ID NO: 14554) 1 agttatttag ttccatgaaa aaattgtctt tgattttcta aaaaaaatcc gcaattactt 61 agttgccaat ccaa.
- (318) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase enzyme is isolated or derived from *Messor bouvieri*. The piggyBac (PB) or piggyBac-like transposase enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between identical to:
- (319) TABLE-US-00122 (SEQ ID NO: 14555) 1 MSSFVPENVH LRHALLFLFH QKKRAAESHR LLVETYGEHA PTIRTCETWF RQFKCGDFNV 61 QDKERPGRPK TFEDAELQEL LDEDSTQTQK QLAEKLNVSR **VAICERLQAM** GKIQKMGRWV 121 PHELNDRQME NRKIVSEMLL QRYERKSFLH RIVTGDEKWI YFENPKRKKS WLSPGEAGPS 181 TARPNRFGRK TMLCVWWDQI GETVNTDRYR QQMINLNCAL IEKRPQYAQR 241 GVVYYELLKP HDKVILQHDN APSHTAKPVK EMLKSLGWEV LSHPPYSPDL APSDYHLFAS MGHALAEQHF 301 ADFEEVKKWL DEWFSSKEKL FFWNGIHKLS ERWTKCIESN GQYFE.
- (320) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Messor bouvieri*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
- (321) TABLE-US-00123 (SEQ ID NO: 14556) 1 agtcagaaat gacacctcga tcgacgacta atcgacgtct aatcgacgtc gattttatgt 61 caacatgtta ccaggtgtgt cggtaattcc tttccggttt ttccggcaga tgtcactagc 121 cataagtatg aaatgttatg atttgataca tatgtcattt taftctactg acattaacct 131 taaaactaca caagttacgt tccgccaaaa taacagcgtt atagatttat aattttttga 241 aa.

In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of: (322) TABLE-US-00124 (SEQ ID NO: 14557) 1 ataaatttga actatccatt ctaagtaacg

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tgttttcttt aacgaaaaaa ccggaaaaaa 61 attaccgaca ctcctggtat gtaaacatgt
tattttcgac attgaatcgc gtcgattcga 121 agtcgatcga ggtgtcattt ctgact.
(323) In certain embodiments of the methods of the disclosure, the transposase enzyme is a piggyBac or
piggyBac-like transposase enzyme. In certain embodiments, the piggyBac or piggyBac-like transposase
enzyme is isolated or derived from Trichoplusia ni. The piggyBac (PB) or piggyBac-like transposase
enzyme may comprise or consist of an amino acid sequence at least 5%, 10%, 15%, 20%, 25%, 30%,
35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 900%, 95%, 99% or any percentage in
between identical to:
(324) TABLE-US-00125 (SEQ ID NO: 14558)
                                                1 MGSSLDDEHI LSALLQSDDE
                      SDHVSEDDVQ SDTEEAFIDE VHEVQPTSSG
                                                                     61 SEILDEONVI
EQPGSSLASN RILTLPQRTI
                                    RGKNKHCWST SKSTRRSRVS ALNIVRSQRG 121
PTRMCRNIYD PLLCFKLFFT DEIISEIVKW
                                                   TNAEISLKRR ESMTSATFRD
TNEDEIYAFF 181 GILVMTAVRK DNHMSTDDLF DRSLSMVYVS
                                                                         VMSRDRFDFL
IRCLRMDDKS IRPTLRENDV 241 FTPVRKIWDL FIHQCIQNYT PGAHLTIDEQ
LLGFRGRCPF RVYIPNKPSK YGIKILMMCD 301 SGTKYMINGM PYLGRGTQTN
                                                      SIPLAKNLLQ 361 EPYKLTIVGT
GVPLGEYYVK
                       ELSKPVHGSC RNITCDNWFT
VRSNKREIPE VLKNSRSRPV
                                     GTSMECFDGP LTLVSYKPKP AKMVYLLSSC 421
DEDASINEST GKPQMVMYYN QTKGGVDTLD
                                                        QMCSVNTCSR KTNPWPMALL
YGMINIACIN 481 SFIIYSHNVS SKGEKVQSRK KFMRNLYMSL
                                                                        TSSEMPEPIE
APTLKRYLRD NISNILPKEV 541 PGTSDDSTEE PVMKKRTYCT YCPSKIRRKA
NASCKKCKKV ICREHNIDMC QSCF.
(325) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from
Trichoplusia ni. In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence
of:
(326) TABLE-US-00126 (SEQ ID NO: 14559)
                                                 1 ttaaccctag aaagatagtc tgcgtaaaat
        tgacgcatgc attcttgaaa tattgctctc 61 tctttctaaa tagcgcgaat ccgtcgctgt
gcatttagga cacctcagtc gccgcttgga 121 gctcccgtga ggcgtgcttg tcaatgcggt
                                                                            aagtgtcact
gattttgaac tataacgacc 181 gcgtgagtca aaatgacgca tgattatctt
                                                               ttacgtgact tttaagattt
                                                  acgtgataac ttattatata tatattttct 301
aactcatacg 241 ataattatat cgttatttca tgttctactt
tgttatagat atc.
In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
(327) TABLE-US-00127 (SEQ ID NO: 14560)
                                                 1 tttgttactt tatagaagaa attttgagtt
tttgtttttt ttcaataaat aaataaacat 61 aaataaattg tttgttgaat ttattattag
                                                                      tatgtaagtg
taaatataat aaaacttaat 121 atctattcaa attaataaat aaacctcgat
                                                             atacagaccg ataaaacaca
tgcgccaatt 181 tcacgcatga ttatcttcaa cgtacgtcac
                                                    aatatgatta tctttccagg
In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
(328) TABLE-US-00128 (SEQ ID NO: 14561) 1 ccctagaaag atagtctgcg taaaattgac
        gcatgcattc ttgaaatatt gctctctctt 61 tctaaatagc gcgaatccgt cgctgtgcat
ttaggacatc tcagtcgccg cttggagctc 121 ccgtgaggcg tgcttgtcaa tgcggtaagt
                                                                           gtcactgatt
ttgaactata acgaccgcgt 181 gagtcaaaat gacgcatgat tatcttttac
                                                               gtgactttta agatttaact
catacgataa 241 ttatattgtt atttcatgtt ctacttacgt
                                                  gataacttat tatatatata ttttcttgtt 301
atagatatc.
In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
(329) TABLE-US-00129 (SEQ ID NO: 14562)
                                                 1 tttgttactt tatagaagaa attttgagtt
                             61 aaataaattg tttgttgaat ttattattag
tttgtttttt tttaataaat aaataaacat
                                                                      tatgtaagtg
taaatataat aaaacttaat 121
                        atctattcaa attaataaat
                                            aaacctcgat
                                                             atacagaccg ataaaacaca
tgcgtcaatt 181 ttacgcatga ttatctttaa cgtacgtcac
                                                   aatatgatta tctttctagg g.
In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
(330) TABLE-US-00130 (SEQ ID NO: 14609)
                                                 1 tctaaatagc gcgaatccgt cgctgtgcat
        ttaggacatc tcagtcgccg cttggagctc 61 ccgtgaggcg tgcttgtcaa tgcggtaagt
gtcactgatt ttgaactata acgaccgcgt 121 gagtcaaaat gacgcatga tatcttttac
                                                                         gtgactttta
agattLaact catacgataa 181 ttatattgtt atttcatgtt ctacttacgt
                                                            gataacttat tatatatata ttttcttgtt
241 atagatatc.
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In certain embodiments, the piggyBac or piggyBac-like transposon comprises a sequence of:
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- (331) TABLE-US-00131 (SEQ ID NO: 14610) 1 tttgttactt tatagaagaa attttgagtt tttgtttttt tttaataaat aaataaacat 61 aaataaattg tttgttgaat ttattattag tatgtaagtg taaatataat aaaacttaat 121 atccattcaa attaataaat aaacctcgat atacagaccg ataaaacaca tgcgtcaatt 181 ttacgcatga ttatctttaa cgtacgtcac aatatgatta tccttctagg g
- (332) In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14561 and SEQ ID NO: 14562, and the piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14558. In certain embodiments, the piggyBac or piggyBac-like transposon comprises SEQ ID NO: 14609 and SEQ ID NO: 14610, and the piggyBac or piggyBac-like transposase comprises SEQ ID NO: 14558.
- (333) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Aphis gossypii*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of CCTTCCAGCGGCGCGC (SEQ ID NO: 14565).
- (334) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Chilo suppressalis*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of CCCAGATTAGCCT (SEQ ID NO: 14566).
- (335) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Heliothis virescens*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of CCCTTAATTACTCGCG (SEQ ID NO: 14567).
- (336) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Pectinophora gossypiella*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of CCCTAGATAACTAAAC (SEQ ID NO: 14568).
- (337) In certain embodiments, the piggyBac or piggyBac-like transposon is isolated or derived from *Anopheles stephensi*. In certain embodiments, the piggyBac or piggyBac-like transposon comprises an ITR sequence of CCCTAGAAAGATA (SEQ ID NO: 14569).
- (338) Immune and Immune Precursor Cells
- (339) In certain embodiments, immune cells of the disclosure comprise lymphoid progenitor cells, natural killer (NK) cells, T lymphocytes (T-cell), stem memory T cells (T.sub.SCM cells), central memory T cells (T.sub.CM), stem cell-like T cells, B lymphocytes (B-cells), myeloid progenitor cells, neutrophils, basophils, eosinophils, monocytes, macrophages, platelets, erythrocytes, red blood cells (RBCs), megakaryocytes or osteoclasts.
- (340) In certain embodiments, immune precursor cells comprise any cells which can differentiate into one or more types of immune cells. In certain embodiments, immune precursor cells comprise multipotent stem cells that can self renew and develop into immune cells. In certain embodiments, immune precursor cells comprise hematopoietic stem cells (HSCs) or descendants thereof. In certain embodiments, immune precursor cells comprise precursor cells that can develop into immune cells. In certain embodiments, the immune precursor cells comprise hematopoietic progenitor cells (HPCs).
- (341) Hematopoietic Stem Cells (HSCs)
- (342) Hematopoietic stem cells (HSCs) are multipotent, self-renewing cells. All differentiated blood cells from the lymphoid and myeloid lineages arise from HSCs. HSCs can be found in adult bone marrow, peripheral blood, mobilized peripheral blood, peritoneal dialysis effluent and umbilical cord blood. (343) HSCs of the disclosure may be isolated or derived from a primary or cultured stem cell. HSCs of the disclosure may be isolated or derived from an embryonic stem cell, a multipotent stem cell, a pluripotent stem cell, an adult stem cell, or an induced pluripotent stem cell (iPSC).
- (344) Immune precursor cells of the disclosure may comprise an HSC or an HSC descendent cell. Exemplary HSC descendent cells of the disclosure include, but are not limited to, multipotent stem cells, lymphoid progenitor cells, natural killer (NK) cells, T lymphocyte cells (T-cells), B lymphocyte cells (B-cells), myeloid progenitor cells, neutrophils, basophils, eosinophils, monocytes, and macrophages. (345) HSCs produced by the methods of the disclosure may retain features of "primitive" stem cells that, while isolated or derived from an adult stem cell and while committed to a single lineage, share characteristics of embryonic stem cells. For example, the "primitive" HSCs produced by the methods of the disclosure retain their "stemness" following division and do not differentiate. Consequently, as an adoptive cell therapy, the "primitive" HSCs produced by the methods of the disclosure not only replenish their numbers, but expand in vivo. "Primitive" HSCs produced by the methods of the disclosure may be

therapeutically-effective when administered as a single dose. In some embodiments, primitive HSCs of the disclosure are CD34+. In some embodiments, primitive HSCs of the disclosure are CD34+ and CD38–. In some embodiments, primitive HSCs of the disclosure are CD34+, CD38– and CD90+. In some embodiments, primitive HSCs of the disclosure are CD34+. CD38–, CD90+ and CD45RA–. In some embodiments, primitive HSCs of the disclosure are CD34+, CD38–, CD90+, CD45RA–, and CD49f+. In some embodiments, the most primitive HSCs of the disclosure are CD34+, CD38–, CD90+, CD45RA–, and CD49f+.

(346) In some embodiments of the disclosure, primitive HSCs, HSCs, and/or HSC descendent cells may be modified according to the methods of the disclosure to express an exogenous sequence (e.g. a chimeric antigen receptor or therapeutic protein). In some embodiments of the disclosure, modified primitive HSCs, modified HSCs, and/or modified HSC descendent cells may be forward differentiated to produce a modified immune cell including, but not limited to, a modified T cell, a modified natural killer cell and/or a modified B-cell of the disclosure.

(347) T Cells

- (348) Modified T cells of the disclosure may be derived from modified hematopoietic stem and progenitor cells (HSPCs) or modified HSCs.
- (349) Unlike traditional biologics and chemotherapeutics, modified-T cells of the disclosure possess the capacity to rapidly reproduce upon antigen recognition, thereby potentially obviating the need for repeat treatments. To achieve this, in some embodiments, modified-T cells of the disclosure not only drive an initial response, but also persist in the patient as a stable population of viable memory T cells to prevent potential relapses. Alternatively, in some embodiments, when it is not desired, modified-T cells of the disclosure do not persist in the patient.
- (350) Intensive efforts have been focused on the development of antigen receptor molecules that do not cause T cell exhaustion through antigen-independent (tonic) signaling, as well as of a modified-T cell product containing early memory T cells, especially stem cell memory (T.sub.SCM) or stem cell-like T cells. Stem cell-like modified-T cells of the disclosure exhibit the greatest capacity for self-renewal and multipotent capacity to derive central memory (T.sub.CM) T cells or T.sub.CM like cells, effector memory (T.sub.EM) and effector T cells (T.sub.E), thereby producing better tumor eradication and long-term modified-T cell engraftment. A linear pathway of differentiation may be responsible for generating these cells: Naïve T cells (T.sub.N)>T.sub.SCM>T.sub.CM>T.sub.EM>T.sub.E>T.sub.TE, whereby T.sub.N is the parent precursor cell that directly gives rise to T.sub.SCM, which then, in turn, directly gives rise to T.sub.CM, etc. Compositions of T cells of the disclosure may comprise one or more of each parental T cell subset with T.sub.SCM cells being the most abundant (e.g.

T.sub.SCM>T.sub.CM>T.sub.EM>T.sub.E>T.sub.TE).

- (351) In some embodiments of the methods of the disclosure, the immune cell precursor is differentiated into or is capable of differentiating into an early memory T cell, a stem cell like T-cell, a Naïve T cells (T.sub.N), a T.sub.SCM, a T.sub.CM, a T.sub.EM, a T.sub.E, or a T.sub.TE. In some embodiments, the immune cell precursor is a primitive HSC, an HSC, or a HSC descendent cell of the disclosure.
- (352) In some embodiments of the methods of the disclosure, the immune cell is an early memory T cell, a stem cell like T-cell, a Naïve T cells (T.sub.N), a T.sub.SCM, a T.sub.CM, a T.sub.EM, a T.sub.E, or a T.sub.TE.
- (353) In some embodiments of the methods of the disclosure, the immune cell is an early memory T cell.
- (354) In some embodiments of the methods of the disclosure, the immune cell is a stem cell like T-cell.
- (355) In some embodiments of the methods of the disclosure, the immune cell is a T.sub.SCM.
- (356) In some embodiments of the methods of the disclosure, the immune cell is a T.sub.CM.
- (357) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of an early memory T cell. In certain embodiments, the plurality of modified early memory T cells comprises at least one modified T.sub.SCM. In certain embodiments, the plurality of modified early memory T cells comprises at least one modified T.sub.CM.

(358) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of a stem cell-like T cell. In certain embodiments, the plurality of modified stem cell-like T cells comprises at least one modified T.sub.SCM. In certain embodiments, the plurality of modified stem cell-like T cells comprises at least one modified T.sub.CM.

- (359) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of a stem memory T cell (T.sub.SCM). In certain embodiments, the cell-surface markers comprise CD62L and CD45RA. In certain embodiments, the cell-surface markers comprise one or more of CD62L, CD45RA, CD28, CCR7, CD127, CD45RO, CD95, CD95 and IL-2R $\beta$ . In certain embodiments, the cell-surface markers comprise one or more of CD45RA, CD95, IL-2R $\beta$ , CCR7, and CD62L.
- (360) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of a central memory T cell (T.sub.CM). In certain embodiments, the cell-surface markers comprise one or more of CD45RO, CD95, IL-2Rβ, CCR7, and CD62L.
- (361) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of a naïve T cell (T.sub.N). In certain embodiments, the cell-surface markers comprise one or more of CD45RA, CCR7 and CD62L. (362) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of an effector T-cell (modified T.sub.EFF). In certain embodiments, the cell-surface markers comprise one or more of CD45RA, CD95, and IL-2Rβ.
- (363) In some embodiments of the methods of the disclosure, the methods modify and/or the methods produce a plurality of modified T cells, wherein at least 2%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, 50%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 99% or any percentage in between of the plurality of modified T cells expresses one or more cell-surface marker(s) of a stem cell-like T cell, a stem memory T cell (T.sub.SCM) or a central memory T cell (T.sub.CM).
- (364) In some embodiments of the methods of the disclosure, a buffer comprises the immune cell or precursor thereof. The buffer maintains or enhances a level of cell viability and/or a stem-like phenotype of the immune cell or precursor thereof, including T-cells. In certain embodiments, the buffer maintains or enhances a level of cell viability and/or a stem-like phenotype of the primary human T cells prior to the nucleofection. In certain embodiments, the buffer maintains or enhances a level of cell viability and/or a stem-like phenotype of the primary human T cells during the nucleofection. In certain embodiments, the buffer maintains or enhances a level of cell viability and/or a stem-like phenotype of the primary human T cells following the nucleofection. In certain embodiments, the buffer comprises one or more of KCl, MgCl.sub.2, ClNa, Glucose and Ca(NO.sub.3).sub.2 in any absolute or relative abundance or concentration, and, optionally, the buffer further comprises a supplement selected from the group consisting of HEPES, Tris/HCl, and a phosphate buffer. In certain embodiments, the buffer comprises 5 mM KCl, 15 mM MgCl2, 90 mM ClNa, 10 mM Glucose and 0.4 mM Ca(NO.sub.3).sub.2. In certain embodiments, the buffer comprises 5 mM KCl, 15 mM MgCl.sub.2, 90 mM ClNa, 10 mM Glucose and 0.4 mM Ca(NO.sub.3).sub.2 and a supplement comprising 20 mM HEPES and 75 mM Tris/HCl. In certain embodiments, the buffer comprises 5 mM KCl, 15 mM MgCl.sub.2, 90 mM ClNa, 10 mM Glucose and 0.4 mM Ca(NO.sub.3).sub.2 and a supplement comprising 40 mM

Na.sub.2HPO.sub.4/NaH.sub.2PO.sub.4 at pH 7.2. In certain embodiments, the composition comprising primary human T cells comprises 100  $\mu$ l of the buffer and between 5×10.sup.6 and 25×10.sup.6 cells. In certain embodiments, the composition comprises a scalable ratio of 250×10.sup.6 primary human T cells per milliliter of buffer or other media during the introduction step.

(365) In some embodiments of the methods of the disclosure, the methods comprise contacting an immune cell of the disclosure, including a T cell of the disclosure, and a T-cell expansion composition. In some embodiments of the methods of the disclosure, the step of introducing a transposon and/or transposase of the disclosure into an immune cell of the disclosure may further comprise contacting the immune cell and a T-cell expansion composition. In some embodiments, including those in which the introducing step of the methods comprises an electroporation or a nucleofection step, the electroporation or a nucleofection step may be performed with the immune cell contacting T-cell expansion composition of the disclosure. (366) In some embodiments of the methods of the disclosure, the T-cell expansion composition comprises, consists essentially of or consists of phosphorus; one or more of an octanoic acid, a palmitic acid, a linoleic acid, and an oleic acid; a sterol; and an alkane.

(367) In certain embodiments of the methods of producing a modified T cell of the disclosure, the expansion supplement comprises one or more cytokine(s). The one or more cytokine(s) may comprise any cytokine, including but not limited to, lymphokines. Exemplary lymphokines include, but are not limited to, interleukin-2 (IL-2), interleukin-3 (IL-3), interleukin-4 (IL-4), interleukin-5 (IL-5), interleukin-6 (IL-6), interleukin-7 (IL-7), interleukin-15 (IL-15), interleukin-21 (IL-21), granulocyte-macrophage colony-stimulating factor (GM-CSF) and interferon-gamma (INFγ). The one or more cytokine(s) may comprise IL-2.

(368) In some embodiments of the methods of the disclosure, the T-cell expansion composition comprises human serum albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, and an expansion supplement. In certain embodiments of this method, the T-cell expansion composition further comprises one or more of octanoic acid, nicotinamide, 2,4,7,9-tetramethyl-5-decyn-4,7-diol (TMDD), diisopropyl adipate (DIPA), n-butyl-benzenesulfonamide, 1,2-benzenedicarboxylic acid, bis(2methylpropyl) ester, palmitic acid, linoleic acid, oleic acid, stearic acid hydrazide, oleamide, a sterol and an alkane. In certain embodiments of this method, the T-cell expansion composition further comprises one or more of octanoic acid, palmitic acid, linoleic acid, oleic acid and a sterol. In certain embodiments of this method, the T-cell expansion composition further comprises one or more of octanoic acid at a concentration of between 0.9 mg/kg to 90 mg/kg, inclusive of the endpoints; palmitic acid at a concentration of between 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; linoleic acid at a concentration of between 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; oleic acid at a concentration of 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; and a sterol at a concentration of about 0.1 mg/kg to 10 mg/kg, inclusive of the endpoints. In certain embodiments of this method, the T-cell expansion composition further comprises one or more of octanoic acid at a concentration of about 9 mg/kg, palmitic acid at a concentration of about 2 mg/kg, linoleic acid at a concentration of about 2 mg/kg, oleic acid at a concentration of about 2 mg/kg and a sterol at a concentration of about 1 mg/kg. In certain embodiments of this method, the T-cell expansion composition further comprises one or more of octanoic acid at a concentration of between 6.4 µmol/kg and 640 µmol/kg, inclusive of the endpoints; palmitic acid at a concentration of between 0.7 µmol/kg and 70 µmol/kg, inclusive of the endpoints; linoleic acid at a concentration of between 0.75 µmol/kg and 75 µmol/kg, inclusive of the endpoints; oleic acid at a concentration of between 0.75 µmol/kg and 75 µmol/kg, inclusive of the endpoints; and a sterol at a concentration of between 0.25 µmol/kg and 25 µmol/kg, inclusive of the endpoints. In certain embodiments of this method, the T-cell expansion composition further comprises one or more of octanoic acid at a concentration of about 64 µmol/kg, palmitic acid at a concentration of about 7 µmol/kg, linoleic acid at a concentration of about 7.5 µmol/kg, oleic acid at a concentration of about 7.5 µmol/kg and a sterol at a concentration of about 2.5 µmol/kg.

(369) In certain embodiments, the T-cell expansion composition comprises one or more of human serum albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, and an expansion supplement to produce a plurality of expanded modified T-cells, wherein at least 2% of the plurality of modified T-cells expresses one or more cell-surface marker(s) of an early memory T cell, a stem cell-like T cell, a stem memory T cell (T.sub.SCM) and/or a central memory T cell (T.sub.CM). In certain embodiments, the

T-cell expansion composition comprises or further comprises one or more of octanoic acid, nicotinamide, 2,4,7,9-tetramethyl-5-decyn-4,7-diol (TMDD), diisopropyl adipate (DIPA), n-butyl-benzenesulfonamide, 1,2-benzenedicarboxylic acid, bis(2-methylpropyl) ester, palmitic acid, linoleic acid, oleic acid, stearic acid hydrazide, oleamide, a sterol and an alkane. In certain embodiments, the T-cell expansion composition comprises one or more of octanoic acid, palmitic acid, linoleic acid, oleic acid and a sterol (e.g. cholesterol). In certain embodiments, the T-cell expansion composition comprises one or more of octanoic acid at a concentration of between 0.9 mg/kg to 90 mg/kg, inclusive of the endpoints; palmitic acid at a concentration of between 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; linoleic acid at a concentration of between 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; oleic acid at a concentration of 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; and a sterol at a concentration of about 0.1 mg/kg to 10 mg/kg, inclusive of the endpoints (wherein mg/kg=parts per million). In certain embodiments, the Tcell expansion composition comprises one or more of octanoic acid at a concentration of about 9 mg/kg, palmitic acid at a concentration of about 2 mg/kg, linoleic acid at a concentration of about 2 mg/kg, oleic acid at a concentration of about 2 mg/kg, and a sterol at a concentration of about 1 mg/kg (wherein mg/kg=parts per million). In certain embodiments, the T-cell expansion composition comprises one or more of octanoic acid at a concentration of 9.19 mg/kg, palmitic acid at a concentration of 1.86 mg/kg, linoleic acid at a concentration of about 2.12 mg/kg, oleic acid at a concentration of about 2.13 mg/kg, and a sterol at a concentration of about 1.01 mg/kg (wherein mg/kg=parts per million). In certain embodiments, the T-cell expansion composition comprises octanoic acid at a concentration of 9.19 mg/kg, palmitic acid at a concentration of 1.86 mg/kg, linoleic acid at a concentration of 2.12 mg/kg, oleic acid at a concentration of about 2.13 mg/kg, and a sterol at a concentration of 1.01 mg/kg (wherein mg/kg=parts per million). In certain embodiments, the T-cell expansion composition comprises one or more of octanoic acid at a concentration of between 6.4 µmol/kg and 640 µmol/kg, inclusive of the endpoints; palmitic acid at a concentration of between 0.7 µmol/kg and 70 µmol/kg, inclusive of the endpoints; linoleic acid at a concentration of between 0.75 µmol/kg and 75 µmol/kg, inclusive of the endpoints; oleic acid at a concentration of between 0.75 µmol/kg and 75 µmol/kg, inclusive of the endpoints; and a sterol at a concentration of between 0.25 µmol/kg and 25 µmol/kg, inclusive of the endpoints. In certain embodiments, the T-cell expansion composition comprises one or more of octanoic acid at a concentration of about 64 µmol/kg, palmitic acid at a concentration of about 7 µmol/kg, linoleic acid at a concentration of about 7.5 µmol/kg, oleic acid at a concentration of about 7.5 µmol/kg and a sterol at a concentration of about 2.5 µmol/kg. In certain embodiments, the T-cell expansion composition comprises one or more of octanoic acid at a concentration of about 63.75 µmol/kg, palmitic acid at a concentration of about 7.27 μmol/kg, linoleic acid at a concentration of about 7.57 μmol/kg, oleic acid at a concentration of about 7.56 µmol/kg and a sterol at a concentration of about 2.61 μmol/kg. In certain embodiments, the T-cell expansion composition comprises octanoic acid at a concentration of about 63.75 µmol/kg, palmitic acid at a concentration of about 7.27 μmol/kg, linoleic acid at a concentration of about 7.57 μmol/cg, oleic acid at a concentration of 7.56 µmol/kg and a sterol at a concentration of 2.61 µmol/kg. (370) As used herein, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of human serum albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, and an expansion supplement at 37° C. Alternatively, or in addition, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of phosphorus, an octanoic fatty acid, a palmitic fatty acid, a linoleic fatty acid and an oleic acid. In certain embodiments, the media comprises an amount of phosphorus that is 10-fold higher than may be found in, for example, Iscove's Modified Dulbecco's Medium (IMDM); available at ThermoFisher Scientific as Catalog number 12440053).

(371) As used herein, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of human serum albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol. Iscove's MDM, and an expansion supplement at 37° C. Alternatively, or in addition, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of the following elements: boron, sodium, magnesium, phosphorus, potassium, and calcium. In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion

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composition" may be used interchangeably with a media comprising one or more of the following
elements present in the corresponding average concentrations: boron at 3.7 mg/L, sodium at 3000 mg/L,
magnesium at 18 mg/L, phosphorus at 29 mg/L, potassium at 15 mg/L and calcium at 4 mg/L.
(372) As used herein, the terms "supplemented T-cell expansion composition" or "T-cell expansion
composition" may be used interchangeably with a media comprising one or more of human serum
albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, and an expansion supplement
at 37° C. Alternatively, or in addition, the terms "supplemented T-cell expansion composition" or "T-cell
expansion composition" may be used interchangeably with a media comprising one or more of the
following components: octanoic acid (CAS No. 124-07-2), nicotinamide (CAS No. 98-92-0), 2,4,7,9-
tetramethyl-5-decyn-4,7-diol (TMDD) (CAS No. 126-86-3), diisopropyl adipate (DIPA) (CAS No. 6938-
94-9), n-butyl-benzenesulfonamide (CAS No. 3622-84-2), 1,2-benzenedicarboxylic acid, bis(2-
methylpropyl) ester (CAS No. 84-69-5), palmitic acid (CAS No. 57-10-3), linoleic acid (CAS No. 60-33-
3), oleic acid (CAS No. 112-80-1), stearic acid hydrazide (CAS No. 4130-54-5), oleamide (CAS No.
3322-62-1), sterol (e.g., cholesterol) (CAS No. 57-88-5), and alkanes (e.g., nonadecane) (CAS No. 629-
92-5) In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell
expansion composition" may be used interchangeably with a media comprising one or more of the
following components: octanoic acid (CAS No. 124-07-2), nicotinamide (CAS No. 98-92-0), 2,4,79-
tetramethyl-5-decyn-4,7-diol (TMDD) (CAS No. 126-86-3), diisopropyl adipate (DIPA) (CAS No. 6938-
94-9), n-butyl-benzenesulfonamide (CAS No. 3622-84-2), 1,2-benzenedicarboxylic acid, bis(2-
methylpropyl) ester (CAS No. 84-69-5), palmitic acid (CAS No. 57-10-3), linoleic acid (CAS No. 60-33-
3), oleic acid (CAS No. 112-80-1), stearic acid hydrazide (CAS No. 4130-54-5), oleamide (CAS No 3322-
62-1), sterol (e.g., cholesterol) (CAS No. 57-88-5), alkanes (e.g., nonadecane) (CAS No. 629-92-5), and
phenol red (CAS No. 143-74-8). In certain embodiments, the terms "supplemented T-cell expansion
composition" or "T-cell expansion composition" may be used interchangeably with a media comprising
one or more of the following components: octanoic acid (CAS No. 124-07-2), nicotinamide (CAS No. 98-
92-0), 2,4,7,9-tetramethyl-5-decyn-4,7-diol (TMDD) (CAS No. 126-86-3), diisopropyl adipate (DIPA)
(CAS No. 6938-94-9), n-butyl-benzenesulfonamide (CAS No. 3622-84-2), 1,2-benzenedicarboxylic acid,
bis(2-methylpropyl) ester (CAS No. 84-69-5), palmitic acid (CAS No. 57-10-3), linoleic acid (CAS No.
60-33-3), oleic acid (CAS No. 112-80-1), stearic acid hydrazide (CAS No. 4130-54-5), oleamide (CAS
No. 3322-62-1), phenol red (CAS No. 143-74-8) and lanolin alcohol.
(373) In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell
expansion composition" may be used interchangeably with a media comprising one or more of human
serum albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, and an expansion
supplement at 37° C. Alternatively, or in addition, the terms "supplemented T-cell expansion composition"
or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of
the following ions: sodium, ammonium, potassium, magnesium, calcium, chloride, sulfate and phosphate.
(374) As used herein, the terms "supplemented T-cell expansion composition" or "T-cell expansion
composition" may be used interchangeably with a media comprising one or more of human serum
albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, and an expansion supplement
at 37° C. Alternatively, or in addition, the terms "supplemented T-cell expansion composition" or "T-cell
expansion composition" may be used interchangeably with a media comprising one or more of the
following free amino acids, histidine, asparagine, serine, glutamate, arginine, glycine, aspartic acid,
glutamic acid, threonine, alanine, proline, cysteine, lysine, tyrosine, methionine, valine, isoleucine,
leucine, phenylalanine and tryptophan. In certain embodiments, the terms "supplemented T-cell expansion
composition" or "T-cell expansion composition" may be used interchangeably with a media comprising
one or more of the following free amino acids in the corresponding average mole percentages: histidine
(about 1%), asparagine (about 0.5%), serine (about 1.5%), glutamine (about 67%), arginine (about 1.5%),
glycine (about 1.5%), aspartic acid (about 1%), glutamic acid (about 2%), threonine (about 2%), alanine
(about 1%), proline (about 1.5%), cysteine (about 1.5%), lysine (about 3%), tyrosine (about 1.5%),
methionine (about 1%), valine (about 3.5%), isoleucine (about 3%), leucine (about 3.5%), phenylalanine
(about 1.5%) and tryptophan (about 0.5%). In certain embodiments, the terms "supplemented T-cell
expansion composition" or "T-cell expansion composition" may be used interchangeably with a media
comprising one or more of the following free amino acids in the corresponding average mole percentages:
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histidine (about 0.78%), asparagine (about 0.4%), serine (about 1.6%), glutamine (about 67.01%), arginine (about 1.67%), glycine (about 1.72%), aspartic acid (about 1.00%), glutamic acid (about 1.93%), threonine (about 2.38%), alanine (about 1.11%), proline (about 1.49%), cysteine (about 1.65%), lysine (about 2.84%), tyrosine (about 1.62%), methionine (about 0.85%), valine (about 3.45%), isoleucine (about 3.14%), leucine (about 3.3%), phenylalanine (about 1.64%) and tryptophan (about 0.37%). (375) As used herein, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of human serum albumin, recombinant human insulin, human transferrin, 2-Mercaptoethanol, Iscove's MDM, and an expansion supplement at 37° C. Alternatively, or in addition, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of phosphorus, an octanoic fatty acid, a palmitic fatty acid, a linoleic fatty acid and an oleic acid. In certain embodiments, the media comprises an amount of phosphorus that is 10-fold higher than may be found in, for example, Iscove's Modified Dulbecco's Medium ((IMDM); available at ThermoFisher Scientific as Catalog number 12440053).

(376) In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid, palmitic acid, linoleic acid, oleic acid and a sterol (e.g. cholesterol). In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of between 0.9 mg/kg to 90 mg/kg, inclusive of the endpoints; palmitic acid at a concentration of between 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; linoleic acid at a concentration of between 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; oleic acid at a concentration of 0.2 mg/kg to 20 mg/kg, inclusive of the endpoints; and a sterol at a concentration of about 0.1 mg/kg to 10 mg/kg, inclusive of the endpoints (wherein mg/kg=parts per million). In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of about 9 mg/kg, palmitic acid at a concentration of about 2 mg/kg, linoleic acid at a concentration of about 2 mg/kg, oleic acid at a concentration of about 2 mg/kg, and a sterol at a concentration of about 1 mg/kg (wherein mg/kg=parts per million). In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of 9.19 mg/kg, palmitic acid at a concentration of 1.86 mg/kg, linoleic acid at a concentration of about 2.12 mg/kg, oleic acid at a concentration of about 2.13 mg/kg, and a sterol at a concentration of about 1.01 mg/kg (wherein mg/kg=parts per million). In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of 9.19 mg/kg, palmitic acid at a concentration of 1.86 mg/kg, linoleic acid at a concentration of 2.12 mg/kg, oleic acid at a concentration of about 2.13 mg/kg, and a sterol at a concentration of 1.01 mg/kg (wherein mg/kg=parts per million). In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of between 6.4 µmol/kg and 640 µmol/kg, inclusive of the endpoints; palmitic acid at a concentration of between 0.7 µmol/kg and 70 µmol/kg, inclusive of the endpoints; linoleic acid at a concentration of between 0.75 µmol/kg and 75 µmol/kg, inclusive of the endpoints; oleic acid at a concentration of between 0.75 µmol/kg and 75 µmol/kg, inclusive of the endpoints; and a sterol at a concentration of between 0.25 μmol/kg and 25 μmol/kg, inclusive of the endpoints. In certain embodiments, the terms "supplemented Tcell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of about 64 µmol/kg, palmitic acid at a concentration of about 7 µmol/kg, linoleic acid at a concentration of about 7.5 µmol/kg, oleic acid at a concentration of about 7.5 µmol/kg and a sterol at a concentration of about 2.5 µmol/kg. (377) In certain embodiments, the terms "supplemented T-cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of about 63.75 µmol/kg, palmitic acid at a concentration of about 7.27 µmol/kg, linoleic acid at a concentration of about 7.57 µmol/kg, oleic acid at a concentration of about 7.56 µmol/kg

and a sterol at a concentration of about 2.61 µmol/kg. In certain embodiments, the terms "supplemented T-

cell expansion composition" or "T-cell expansion composition" may be used interchangeably with a media comprising one or more of octanoic acid at a concentration of about 63.75 µmol/kg, palmitic acid at a concentration of about 7.27 µmol/kg, linoleic acid at a concentration of about 7.57 µmol/kg, oleic acid at a concentration of 7.56 µmol/kg and a sterol at a concentration of 2.61 µmol/kg.

(378) In certain embodiments of the methods of producing a modified T cell (e.g. a stem cell-like T cell, a T.sub.SCM and/or a T.sub.CM) of the disclosure, the method comprises contacting a modified T cell and an inhibitor of the PI3K-Akt-mTOR pathway. Modified T-cells of the disclosure, including modified stem cell-like T cells, T.sub.SCM and/or T.sub.CM of the disclosure, may be incubated, cultured, grown, stored, or otherwise, combined at any step in the methods of the procedure with a growth medium comprising one or more inhibitors a component of a PI3K pathway. Exemplary inhibitors a component of a PI3K pathway include, but are not limited to, an inhibitor of GSK30 such as TWS119 (also known as GSK 3B inhibitor XII; CAS Number 601514-19-6 having a chemical formula C.sub.18H.sub.14N.sub.4O.sub.2). Exemplary inhibitors of a component of a PI3K pathway include, but are not limited to, bb007 (BLUEBIRDBIO™). Additional Exemplary inhibitors of a component of a PI3K pathway include, but are not limited to, an allosteric Akt inhibitor VIII (also referred to as Akti-1/2 having Compound number 10196499), ATP competitive inhibitors (Orthosteric inhibitors targeting the ATP-binding pocket of the protein kinase B (Akt)), Isoquinoline-5-sulfonamides (H-8, H-89, and NL-71-101), Azepane derivatives (A series of structures derived from (-)-balanol), Aminofurazans (GSK690693). Heterocyclic rings (7-azaindole, 6phenylpurine derivatives, pyrrolo[2,3-d]pyrimidine derivatives, CCT128930, 3-aminopyrrolidine, anilinotriazole derivatives, spiroindoline derivatives, AZD5363, ipatasertib (GDC-0068, RG7440), A-674563, and A-443654). Phenylpyrazole derivatives (AT7867 and AT13148), Thiophenecarboxamide derivatives (Afuresertib (GSK2110183), 2-pyrimidyl-5-amidothiophene derivative (DC120), uprosertib (GSK2141795)), Allosteric inhibitors (Superior to orthosteric inhibitors providing greater specificity, reduced side-effects and less toxicity). 2,3-diphenylquinoxaline analogues (2,3-diphenylquinoxaline derivatives, triazolo[3,4-f][1,6]naphthyridin-3(2H)-one derivative (MK-2206)), Alkylphospholipids (Edelfosine (1-O-octadecyl-2-O-methyl-rac-glycero-3-phosphocholine, ET-8-OCH.sub.3) ilmofosine (BM 41.440), miltefosine (hexadecylphosphocholine, HePC), perifosine (D-21266), erucylphosphocholine (ErPC), erufosine (ErPC3, erucylphosphohomocholine), Indole-3-carbinol analogues (Indole-3-carbinol, 3-chloroacetylindole, diindolylmethane, diethyl 6-methoxy-5,7-dihydroindolo [2,3-b]carbazole-2,10dicarboxylate (SR13668), OSU-A9), Sulfonamide derivatives (PH-316 and PHT-427), Thiourea derivatives (PIT-1, PIT-2, DM-PIT-1, N-[(1-methyl-1H-pyrazol-4-yl)carbonyl]-N'-(3-bromophenyl)thiourea), Purine derivatives (Triciribine (TCN, NSC 154020), triciribine mono-phosphate active analogue (TCN-P), 4-amino-pyrido[2,3-d]pyrimidine derivative API-1, 3-phenyl-3H-imidazo[4,5-b]pyridine derivatives, ARQ 092). BAY 1125976, 3-methyl-xanthine, quinoline-4-carboxamide and 2-[4-(cyclohexa-1,3-dien-1-yl)-1H-pyrazol-3-yl]phenol, 3-oxo-tirucallic acid,  $3\alpha$ - and  $3\beta$ -acetoxy-tirucallic acids, acetoxytirucallic acid, and irreversible inhibitors (antibiotics, Lactoquinomycin, Frenolicin B, kalafungin, medermycin, Boc-Phe-vinyl ketone, 4-hydroxynonenal (4-HNE), 1,6-naphthyridinone derivatives, and imidazo-1,2-pyridine derivatives).

- (379) In certain embodiments of the methods of producing a modified T cell (e.g. a stem cell-like T cell, a T.sub.SCM and/or a T.sub.CM) of the disclosure, the method comprises contacting a modified T cell and an inhibitor of T cell effector differentiation. Exemplary inhibitors of T cell effector differentiation include, but are not limited to, a BET inhibitor (e.g. JQ1, a hienotriazolodiazepine) and/or an inhibitor of the BET family of proteins (e.g. BRD2, BRD3, BRD4, and BRDT).
- (380) In certain embodiments of the methods of producing a modified T cell (e.g. a stem cell-like T cell, a T.sub.SCM and/or a T.sub.CM) of the disclosure, the method comprises contacting a modified T cell and an agent that reduces nucleo-cytoplasmic Acetyl-CoA. Exemplary agents that reduce nucleo-cytoplasmic Acetyl-CoA include, but are not limited to, 2-hydroxy-citrate (2-HC) as well as agents that increase expression of Acss1.
- (381) In certain embodiments of the methods of producing a modified T cell (e.g. a stem cell-like T cell, a TSCM and/or a TCM) of the disclosure, the method comprises contacting a modified T cell and a composition comprising a histone deacetylase (HDAC) inhibitor. In some embodiments, the composition comprising an HDAC inhibitor comprises or consists of valproic acid, Sodium Phenylbutyrate (NaPB) or a combination thereof. In some embodiments, the composition comprising an HDAC inhibitor comprises or

consists of valproic acid. In some embodiments, the composition comprising an HDAC inhibitor comprises or consists of Sodium Phenylbutyrate (NaPB).

- (382) In certain embodiments of the methods of producing a modified T cell (e.g. a stem cell-like T cell, a T.sub.SCM and/or a T.sub.CM) of the disclosure, the activation supplement may comprise one or more cytokine(s). The one or more cytokine(s) may comprise any cytokine, including but not limited to, lymphokines. Exemplary lymphokines include, but are not limited to, interleukin-2 (IL-2), interleukin-3 (IL-3), interleukin-4 (IL-4), interleukin-5 (IL-5), interleukin-6 (IL-6), interleukin-7 (IL-7), interleukin-15 (IL-15), interleukin-21 (IL-21), granulocyte-macrophage colony-stimulating factor (GM-CSF) and interferon-gamma (INFγ). The one or more cytokine(s) may comprise IL-2.
- (383) In certain embodiments of the methods of producing a modified T cell (e.g. a stem cell-like T cell, a T.sub.SCM and/or a T.sub.CM) of the disclosure, the activation supplement may comprise one or more activator complexes. Exemplary and nonlimiting activator complexes may comprise a monomeric, dimeric, trimeric or tetrameric antibody complex that binds one or more of CD3, CD28, and CD2. In some embodiments, the activation supplement comprises or consists of an activator complex that comprises a human, a humanized or a recombinant or a chimeric antibody. In some embodiments, the activation supplement comprises or consists of an activator complex that binds CD3 and CD28. In some embodiments, the activation supplement comprises or consists of an activator complex that binds CD3, CD28 and CD2.
- (384) Natural Killer (NK) Cells
- (385) In certain embodiments, the modified immune or immune precursor cells of the disclosure are natural killer (NK) cells. In certain embodiments, NK cells are cytotoxic lymphocytes that differentiate from lymphoid progenitor cells.
- (386) Modified NK cells of the disclosure may be derived from modified hematopoietic stem and progenitor cells (HSPCs) or modified HSCs.
- (387) In certain embodiments, non-activated NK cells are derived from CD3-depleted leukopheresis (containing CD14/CD19/CD56+ cells).
- (388) In certain embodiments, NK cells are electroporated using a Lonza 4D nucleofector or BTX ECM 830 (500V, 700 usec pulse length, 0.2 mm electrode gap, one pulse). All Lonza 4D nucleofector programs are contemplated as within the scope of the methods of the disclosure.
- (389) In certain embodiments, 5×10E6 cells were electroporated per electroporation in 100 μL P3 buffer in cuvettes. However, this ratio of cells per volume is scalable for commercial manufacturing methods. (390) In certain embodiments, NK cells were stimulated by co-culture with an additional cell line. In
- certain embodiments, the additional cell line comprises artificial antigen presenting cells (aAPCs). In certain embodiments, stimulation occurs at day 1, 2, 3, 4, 5, 6, or 7 following electroporation. In certain embodiments, stimulation occurs at day 2 following electroporation.
- (391) In certain embodiments, NK cells express CD56.
- (392) B Cells
- (393) In certain embodiments, the modified immune or immune precursor cells of the disclosure are B cells. B cells are a type of lymphocyte that express B cell receptors on the cell surface. B cell receptors bind to specific antigens.
- (394) Modified B cells of the disclosure may be derived from modified hematopoietic stem and progenitor cells (HSPCs) or modified HSCs.
- (395) In certain embodiments, HSPCs are modified using the methods of the disclosure, and then primed for B cell differentiation in presence of human IL-3, Flt3L, TPO, SCF, and G-CSF for at least 3 days, at least 4 days, at least 5 days, at least 6 days or at least 7 days. In certain embodiments. HSPCs are modified using the methods of the disclosure, and then primed for B cell differentiation in presence of human IL-3, Flt3L, TPO, SCF, and G-CSF for 5 days.
- (396) In certain embodiments, following priming, modified HSPC cells are transferred to a layer of feeder cells and fed bi-weekly, along with transfer to a fresh layer of feeders once per week. In certain embodiments, the feeder cells are MS-5 feeder cells.
- (397) In certain embodiments, modified HSPC cells are cultured with MS-5 feeder cells for at least 7, 14, 21, 28, 30, 33, 35, 42 or 48 days. In certain embodiments, modified HSPC cells are cultured with MS-5 feeder cells for 33 days.

- (398) Methods of Cell Modification
- (399) In some embodiments of the methods of the disclosure, a composition comprises a scalable ratio of 250×10.sup.6 primary human T cells per milliliter of buffer or other media during a delivery or an introduction step.
- (400) In some embodiments of the methods of the disclosure, a composition is delivered or introduced to a cell by electroporation or nucleofection. In some embodiments, a delivery or introduction step comprises electroporation or nucleofection.
- (401) In some embodiments of the methods of the disclosure, a composition is delivered or introduced to a cell by a method other than electroporation or nucleofection.
- (402) In some embodiments of the methods of the disclosure, a composition is delivered or introduced by one or more of topical delivery, adsorption, absorption, electroporation, spin-fection, co-culture, transfection, mechanical delivery, sonic delivery, vibrational delivery, magnetofection or by nanoparticle-mediated delivery. In some embodiments, a delivery or introduction step comprises one or more of topical delivery, adsorption, absorption, electroporation, spin-fection, co-culture, transfection, mechanical delivery, sonic delivery, vibrational delivery, magnetofection or by nanoparticle-mediated delivery. (403) In some embodiments of the methods of the disclosure, a composition is delivered or introduced by liposomal transfection, calcium phosphate transfection, fugene transfection, and dendrimer-mediated transfection. In some embodiments, a delivery or introduction step comprises one or more of liposomal transfection, calcium phosphate transfection, fugene transfection, and dendrimer-mediated transfection. (404) In some embodiments of the methods of the disclosure, a composition is delivered or introduced by mechanical transfection comprises cell squeezing, cell bombardment, or gene gun techniques. In some embodiments, a delivery or introduction step comprises one or more of mechanical transfection comprises cell squeezing, cell bombardment, or gene gun techniques.
- (405) In some embodiments of the methods of the disclosure, a composition is delivered or introduced by nanoparticle-mediated transfection comprises liposomal delivery, delivery by micelles, and delivery by polymerosomes. In some embodiments, a delivery or introduction step comprises one or more of liposomal delivery, delivery by micelles, and delivery by polymerosomes.
- (406) Non-Transposition Methods of Delivery
- (407) In some embodiments of the compositions and methods of the disclosure, a modified cell of the disclosure may be produced by introducing a sequence into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure. The introducing step may comprise delivery of a sequence and/or a gene editing composition via a non-transposition delivery system. The introduction step may be performed ex vivo, in vivo, in vitro or in situ.
- (408) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure comprises one or more of topical delivery, adsorption, absorption, electroporation, spin-fection, co-culture, transfection, mechanical delivery, sonic delivery, vibrational delivery, magnetofection and nanoparticle-mediated delivery.
- (409) In some embodiments of the compositions and methods of the disclosure, introducing a nucleic acid sequence and/or a gene editing construct into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure comprises liposomal transfection, calcium phosphate transfection, fugene transfection, and dendrimer-mediated transfection.
- (410) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure by mechanical transfection comprises cell squeezing, cell bombardment, or gene gun techniques.
- (411) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure by nanoparticle-mediated transfection comprises one or more of a liposome, a micelle, a polymer and a polymerosome.
- (412) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure comprises a non-viral vector. In some embodiments, the non-viral vector

comprises the sequence and/or the gene editing composition. In some embodiments, the non-viral vector comprises plasmid DNA, linear double-stranded DNA (dsDNA), linear single-stranded DNA (ssDNA). DoggyBone™ DNA, nanoplasmids, minicircle DNA, single-stranded oligodeoxynucleotides (ssODN), DDNA oligonucleotides, single-stranded mRNA (ssRNA), and double-stranded mRNA (dsRNA). (413) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure comprises a viral vector. In some embodiments, the viral vector is a nonintegrating and/or non-chromosomal vector. Exemplary non-integrating non-chromosomal vectors include, but are not limited to, adeno-associated virus (AAV), adenovirus, and herpes viruses. In some embodiments, the viral vector is an integrating chromosomal vector. Integrating chromosomal vectors include, but are not limited to, adeno-associated vectors (AAV), Lentiviruses, and gamma-retroviruses. In some embodiments, the viral vector comprises the sequence and/or the gene editing composition. (414) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure comprises a combination of vectors of the disclosure. Exemplary, nonlimiting vector combinations include: viral and non-viral vectors, a plurality of non-viral vectors, or a plurality of viral vectors. Exemplary but non-limiting vectors combinations include: a combination of a DNA-derived and an RNA-derived vector, a combination of non-viral expression vector and a viral delivery vector, a combination of a non-viral expression vector and a nanoparticle delivery vector, a combination of two distinct non-viral expression vectors, a combination of a non-viral expression vector and a mechanical or chemical method of transfection.

(415) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure stably integrates a sequence, transiently integrates a sequence, produces site-specific integration of a sequence, or produces a biased integration of a sequence. In some embodiments, the sequence is a nucleic acid sequence. In some embodiments, the nucleic acid sequence comprises a transgene.

(416) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure stably integrates a sequence. In some embodiments, the sequence is a nucleic acid sequence. In some embodiments, the stable chromosomal integration can be a random integration, a site-specific integration, or a biased integration. In some embodiments, the site-specific integration can be non-assisted or assisted. In some embodiments, the assisted site-specific integration is co-delivered with a site-directed nuclease. In some embodiments, the site-directed nuclease comprises a transgene with 5' and 3' nucleotide sequence extensions that contain a percentage homology to upstream and downstream regions of the site of genomic integration. In some embodiments, the transgene with homologous nucleotide extensions enable genomic integration by homologous recombination, microhomology-mediated end joining, or nonhomologous end-joining. In some embodiments the sitespecific integration occurs at a safe harbor site. Genomic safe harbor sites are able to accommodate the integration of new genetic material in a manner that ensures that the newly inserted genetic elements function reliably (for example, are expressed at a therapeutically effective level of expression) and do not cause deleterious alterations to the host genome that cause a risk to the host organism. Potential genomic safe harbors include, but are not limited to, intronic sequences of the human albumin gene, the adenoassociated virus site 1 (AAVS1), a naturally occurring site of integration of AAV virus on chromosome 19, the site of the chemokine (C-C motif) receptor 5 (CCR5) gene and the site of the human ortholog of the mouse Rosa26 locus.

(417) In some embodiments, the site-specific transgene integration occurs at a site that disrupts expression of a target gene. In some embodiments, disruption of target gene expression occurs by site-specific integration at introns, exons, promoters, genetic elements, enhancers, suppressors, start codons, stop codons, and response elements. In some embodiments, exemplary target genes targeted by site-specific integration include but are not limited to TRAC, TRAB, PD1, any immunosuppressive gene, and genes involved in allo-rejection.

(418) In some embodiments of the compositions and methods of the disclosure, introducing a sequence

and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure site-specific transgene integration occurs at a site that results in enhanced expression of a target gene. In some embodiments, enhancement of target gene expression occurs by site-specific integration at introns, exons, promoters, genetic elements, enhancers, suppressors, start codons, stop codons, and response elements.

- (419) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure enzymes may be used to create strand breaks in the host genome to facilitate delivery or integration of the transgene. In some embodiments, enzymes create single-strand breaks. In some embodiments, enzymes create double-strand breaks. In some embodiments, examples of break-inducing enzymes include but are not limited to: transposases, integrases, endonucleases, CRISPR-Cas9, transcription activator-like effector nucleases (TALEN), zinc finger nucleases (ZFN), Cas-CLOVER™, and CPF1. In some embodiments, break-inducing enzymes can be delivered to the cell encoded in DNA, encoded in mRNA, as a protein, as a nucleoprotein complex with a guide RNA (gRNA). (420) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure site-specific transgene integration is controlled by a vector-mediated integration site bias. In some embodiments vector-mediated integration site bias is controlled by the chosen lentiviral vector. In some embodiments vector-mediated integration site bias is controlled by the chosen gamma-retroviral vector.
- (421) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure site-specific transgene integration site is a non-stable chromosomal insertion. In some embodiments, the integrated transgene may become silenced, removed, excised, or further modified.
- (422) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure genome modification comprises a non-stable integration of a transgene. In some embodiments, the non-stable integration can be a transient non-chromosomal integration, a semi-stable non chromosomal integration, a semi-persistent non-chromosomal insertion, or a non-stable chromosomal insertion. In some embodiments, the transient non-chromosomal insertion can be epi-chromosomal or cytoplasmic.
- (423) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure the transient non-chromosomal insertion of a transgene does not integrate into a chromosome and the modified genetic material is not replicated during cell division.
- (424) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure genome modification comprises a semi-stable or persistent non-chromosomal integration of a transgene. In some embodiments, a DNA vector encodes a Scaffold/matrix attachment region (S-MAR) module that binds to nuclear matrix proteins for episomal retention of a non-viral vector allowing for autonomous replication in the nucleus of dividing cells.
- (425) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure genome modification is a non-stable chromosomal integration of a transgene. In some embodiments, the integrated transgene may become silenced, removed, excised, or further modified.
- (426) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC, an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure modification to the genome by transgene insertion can occur via host cell-directed double-strand breakage repair (homology-directed repair) by homologous recombination (HR), microhomology-mediated end joining (MMEJ), nonhomologous end joining (NHEJ), transposase enzyme-mediated modification, integrase enzyme-mediated

modification, or recombinant enzyme-mediated modification. In some embodiments, the modification to the genome by transgene insertion can occur via CRISPR-Cas9, TALEN, ZFNs, Cas-CLOVER, and cpf1. (427) Nanoparticle Delivery

- (428) In some embodiments of the compositions and methods of the disclosure, introducing a sequence and/or a gene editing composition into an HSC an HSC descendent cell, an immune cell or an immune precursor cell of the disclosure comprise a nanoparticle vector. Nanoparticle vectors may encapsulate a composition of the disclosure. Alternatively, or in addition, a surface of a nanoparticle vector may comprise a composition of the disclosure. In some embodiments, the surface is an interior surface. In some embodiments, the surface comprises a composition of the disclosure integrated therein or thereon.
- (429) Nonlimiting examples of nanoparticle vectors of the disclosure may comprise one or more of a hydrophilic block, a hydrophobic block, and a charged block. In some embodiments, the hydrophilic block may be poly(ethylene oxide) (PEO), and the charged block may be poly(L-histidine).
- (430) The disclosure provides nanoparticle vectors comprising di-block and tri-block co-polymers. Exemplary di-block co-polymers may comprise one or more of a hydrophilic block, a hydrophobic block, and a charged block. In some embodiments, the hydrophilic block may be poly(ethylene oxide) (PEO), and the charged block may be poly(L-histidine). Exemplary tri-block co-polymers may comprise one or more of a hydrophilic block, a hydrophobic block, and a charged block. In some embodiments, the hydrophilic block may be poly(ethylene oxide) (PEO), and the charged block may be poly(L-histidine). (431) An exemplary tri-block copolymer that may be used in various embodiments is a PEO-b-PLA-b-PHIS, with variable numbers of repeating units in each block varying by design.
- (432) Poly(histidine) (i.e., poly(L-histidine)), is a pH-sensitive polymer due to the imidazole ring providing an electron lone pair on the unsaturated nitrogen. That is, poly(histidine) has amphoteric properties through protonation-deprotonation. The various embodiments enable intracellular delivery of compositions of the disclosure, including gene editing compositions, by, for example, complexing with poly(histidine)-based micelles.
- (433) Diblock copolymers that may be used as intermediates for making triblock copolymers of the embodiment micelles may have hydrophilic biocompatible poly(ethylene oxide) (PEO), which is chemically synonymous with PEG, coupled to various hydrophobic aliphatic poly(anhydrides), poly(nucleic acids), poly(esters), poly(ortho esters), poly(peptides), poly(phosphazenes) and poly(saccharides), including but not limited by poly(lactide) (PLA), poly(glycolide)(PLGA), poly(lactic-co-glycolic acid) (PLGA), poly(ε-caprolactone) (PCL), and poly (trimethylene carbonate) (PTMC). (434) Polymeric micelles comprised of 100% PEGylated surfaces possess improved in vitro chemical stability, augmented in vivo bioavailability, and prolonged blood circulatory half-lives. For example, aliphatic polyesters, constituting the polymeric micelle's membrane portions, are degraded by hydrolysis of their ester linkages in physiological conditions such as in the human body. Because of their biodegradable nature, aliphatic polyesters have received a great deal of attention for use as implantable biomaterials in drug delivery devices, bioresorbable sutures, adhesion barriers, and as scaffolds for injury repair via tissue engineering.
- (435) Without wishing to be bound by a particular theory, it is believed that believed that in the micelles that are formed by the various embodiment triblock copolymers, the hydrophobic blocks aggregate to form a core, leaving the hydrophilic blocks and poly(histidine) blocks on the ends to form one or more surrounding layer.
- (436) Scaffold Proteins
- (437) Protein scaffolds of the disclosure may be derived from a fibronectin type III (FN3) repeat protein, encoding or complementary nucleic acids, vectors, host cells, compositions, combinations, formulations, devices, and methods of making and using them. In a preferred embodiment, the protein scaffold is comprised of a consensus sequence of multiple FN3 domains from human Tenascin-C (hereinafter "Tenascin"). In a further preferred embodiment, the protein scaffold of the present disclosure is a consensus sequence of 15 FN3 domains. The protein scaffolds of the disclosure can be designed to bind various molecules, for example, a cellular target protein. In a preferred embodiment, the protein scaffolds of the disclosure can be designed to bind an epitope of a wild type and/or variant form of a ligand or an antigen.

- (438) Protein scaffolds of the disclosure may include additional molecules or moieties, for example, the Fc region of an antibody, albumin binding domain, or other moiety influencing half-life. In further embodiments, the protein scaffolds of the disclosure may be bound to a nucleic acid molecule that may encode the protein scaffold.
- (439) The disclosure provides at least one method for expressing at least one protein scaffold based on a consensus sequence of multiple FN3 domains, in a host cell, comprising culturing a host cell as described herein under conditions wherein at least one protein scaffold is expressed in detectable and/or recoverable amounts.
- (440) The disclosure provides at least one composition comprising (a) a protein scaffold based on a consensus sequence of multiple FN3 domains and/or encoding nucleic acid as described herein; and (b) a suitable and/or pharmaceutically acceptable carrier or diluent.
- (441) The disclosure provides a method of generating libraries of a protein scaffold based on a fibronectin type III (FN3) repeat protein, preferably, a consensus sequence of multiple FN3 domains and, more preferably, a consensus sequence of multiple FN3 domains from human Tenascin. The library is formed by making successive generations of scaffolds by altering (by mutation) the amino acids or the number of amino acids in the molecules in particular positions in portions of the scaffold, e.g., loop regions. Libraries can be generated by altering the amino acid composition of a single loop or the simultaneous alteration of multiple loops or additional positions of the scaffold molecule. The loops that are altered can be lengthened or shortened accordingly. Such libraries can be generated to include all possible amino acids at each position, or a designed subset of amino acids. The library members can be used for screening by display, such as in vitro or CIS display (DNA, RNA, ribosome display, etc.), yeast, bacterial, and phage display.
- (442) Protein scaffolds of the disclosure provide enhanced biophysical properties, such as stability under reducing conditions and solubility at high concentrations; they may be expressed and folded in prokaryotic systems, such as *E. coli*, in eukaryotic systems, such as yeast, and in in vitro transcription/translation systems, such as the rabbit reticulocyte lysate system.
- (443) The disclosure provides a method of generating a scaffold molecule that binds to a particular target by panning the scaffold library of the invention with the target and detecting binders. In other related aspects, the disclosure comprises screening methods that may be used to generate or affinity mature protein scaffolds with the desired activity, e.g., capable of binding to target proteins with a certain affinity. Affinity maturation can be accomplished by iterative rounds of mutagenesis and selection using systems, such as phage display or in vitro display. Mutagenesis during this process may be the result of site directed mutagenesis to specific scaffold residues, random mutagenesis due to error-prone PCR, DNA shuffling, and/or a combination of these techniques.
- (444) The disclosure provides an isolated, recombinant and/or synthetic protein scaffold based on a consensus sequence of fibronectin type III (FN3) repeat protein, including, without limitation, mammalian-derived scaffold, as well as compositions and encoding nucleic acid molecules comprising at least one polynucleotide encoding protein scaffold based on the consensus FN3 sequence. The disclosure further includes, but is not limited to, methods of making and using such nucleic acids and protein scaffolds, including diagnostic and therapeutic compositions, methods and devices.
- (445) The protein scaffolds of the disclosure offer advantages over conventional therapeutics, such as ability to administer locally, orally, or cross the blood-brain barrier, ability to express in *E. Coli* allowing for increased expression of protein as a function of resources versus mammalian cell expression ability to be engineered into bispecific or tandem molecules that bind to multiple targets or multiple epitopes of the same target, ability to be conjugated to drugs, polymers, and probes, ability to be formulated to high concentrations, and the ability of such molecules to effectively penetrate diseased tissues and tumors. (446) Moreover, the protein scaffolds possess many of the properties of antibodies in relation to their fold that mimics the variable region of an antibody. This orientation enables the FN3 loops to be exposed similar to antibody complementarity determining regions (CDRs). They should be able to bind to cellular targets and the loops can be altered, e.g., affinity matured, to improve certain binding or related properties. (447) Three of the six loops of the protein scaffold of the disclosure correspond topologically to the complementarity determining regions (CDRs 1-3), i.e., antigen-binding regions, of an antibody, while the remaining three loops are surface exposed in a manner similar to antibody CDRs. These loops span at or

about residues 13-16, 22-28, 38-43, 51-54, 60-64, and 75-81 of the consensus sequence. Preferably, the loop regions at or about residues 22-28, 51-54, and 75-81 are altered for binding specificity and affinity. One or more of these loop regions are randomized with other loop regions and/or other strands maintaining their sequence as backbone portions to populate a library and potent binders can be selected from the library having high affinity for a particular protein target. One or more of the loop regions can interact with a target protein similar to an antibody CDR interaction with the protein.

(448) Discovery of Antigen/Ligand Recognition Region Sequences

- (449) The disclosure provides a method of generating libraries of antigen/ligand recognition region (ARR/LRR) sequences for binding antigens and/or ligands of the disclosure. The library is formed by making successive generations of ARR/LRR sequences by altering (by mutation) the amino acids or the number of amino acids in the sequences at particular positions of the ARR/LRR In some embodiments, the ARR/LRR comprises one or more of a protein scaffold, an antibody mimetic, a Centyrin, a single chain antibody (scFv), a single domain antibody, a VHH and a VH of the disclosure. In some embodiments, the library is formed by making successive generations of ARR/LRR sequences by altering (by mutation) the amino acids or the number of amino acids in the sequences at particular positions of an antibody, an ScFv, VHH or VH, e.g., one or more complementarity determining regions (CDR) and/or framework regions of a variable domain.
- (450) Libraries can be generated by altering the amino acid composition of a single CDR or the simultaneous alteration of multiple CDRs or additional positions of an antibody, an scFv, VHH or VH (e.g. a framework sequence of the variable region). The CDR and/or framework sequence of the variable domain that are altered can be lengthened or shortened accordingly.
- (451) Libraries can be generated by altering the amino acid composition of a loop of a scaffold protein or a Centyrin. The loop sequences that are altered can be lengthened or shortened accordingly.
- (452) Libraries can be generated by altering the amino acid composition of an antigen or ligand-binding or specificity-determining region of an antibody mimetic.
- (453) Such libraries can be generated to include all possible amino acids at each position, or a designed subset of amino acids. The library members can be used for screening by display, such as in vitro or CIS display (DNA, RNA, ribosome display, etc.), yeast, bacterial, and phage display.
- (454) ARRs/LRRs of the disclosure provide enhanced biophysical properties, such as stability under reducing conditions and solubility at high concentrations; they may be expressed and folded in prokaryotic systems, such as *E. coli*, in eukaryotic systems, such as yeast, and in in vitro transcription/translation systems, such as the rabbit reticulocyte lysate system.
- (455) The disclosure provides a method of generating an ARR/LRR or a portion thereof that binds to a particular target by panning a library of the invention with the target and detecting binders. In other related aspects, the disclosure comprises screening methods that may be used to generate or affinity mature ARRs/LRRs with the desired activity. e.g., capable of binding to target proteins with a certain affinity. Affinity maturation can be accomplished by iterative rounds of mutagenesis and selection using systems, such as phage display or in vitro display. Mutagenesis during this process may be the result of site directed mutagenesis to specific protein residues, random mutagenesis due to error-prone PCR, DNA shuffling, and/or a combination of these techniques.
- (456) The disclosure provides an isolated, recombinant and/or synthetic protein scaffold comprising at least one VHH. The disclosure further includes, but is not limited to, methods of making and using such nucleic acids and protein scaffolds, including diagnostic and therapeutic compositions, methods and devices.
- (457) The compositions of the disclosure offer advantages over conventional therapeutics, such as ability to administer locally, orally, or cross the blood-brain barrier, ability to express in *E. Coli* allowing for increased expression of protein as a function of resources versus mammalian cell expression ability to be engineered into bispecific or tandem molecules that bind to multiple targets or multiple epitopes of the same target, ability to be conjugated to drugs, polymers, and probes, ability to be formulated to high concentrations, and the ability of such molecules to effectively penetrate diseased tissues and tumors. (458) Production and Generation of Proteins
- (459) Proteins of the disclosure can be optionally produced by a cell line, a mixed cell line, an immortalized cell or clonal population of immortalized cells, as well known in the art See, e.g., Ausubel, et

al., ed., Current Protocols in Molecular Biology, John Wiley & Sons, Inc., NY, N.Y. (1987-2001); Sambrook, et al., Molecular Cloning: A Laboratory Manual. 2nd Edition, Cold Spring Harbor, N.Y. (1989); Harlow and Lane, Antibodies, a Laboratory Manual, Cold Spring Harbor, N.Y. (1989); Colligan, et al., eds., Current Protocols in Immunology, John Wiley & Sons, Inc., NY (1994-2001); Colligan et al., Current Protocols in Protein Science, John Wiley & Sons, NY, N.Y., (1997-2001).

- (460) Amino acids encoding a protein can be altered, added and/or deleted to reduce immunogenicity or reduce, enhance or modify binding, affinity, on-rate, off-rate, avidity, specificity, half-life, stability, solubility or any other suitable characteristic, as known in the art.
- (461) Proteins can be engineered with retention of high affinity for an antigen or a ligand as well as other favorable biological properties. To achieve this goal, the proteins can be optionally prepared by a process of analysis of the parental sequences and various conceptual engineered products using three-dimensional models of the parental and engineered sequences. Three-dimensional models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate sequences and can measure possible immunogenicity (e.g., Immunofilter program of Xencor, Inc. of Monrovia, Calif.). Inspection of these displays permits analysis of the likely role of the residues in the functioning of the candidate sequence, i.e., the analysis of residues that influence the ability of the protein to bind its antigen. In this way, residues can be selected and combined from the parent and reference sequences so that the desired characteristic, such as affinity for the target antigen(s)/ligand(s), is achieved. Alternatively, or in addition to, the above procedures, other suitable methods of engineering can be used.

(462) Screening of ARRs/LRRs

(463) Screening protein ARRs/LRRs or any portion thereof for specific binding to similar proteins or fragments can be conveniently achieved using nucleotide (DNA or RNA display) or peptide display libraries, for example, in vitro display. This method involves the screening of large collections of peptides for individual members having the desired function or structure. The displayed nucleotide or peptide sequences can be from 3 to 5000 or more nucleotides or amino acids in length, frequently from 5-100 amino acids long, and often from about 8 to 25 amino acids long. In addition to direct chemical synthetic methods for generating peptide libraries, several recombinant DNA methods have been described. One type involves the display of a peptide sequence on the surface of a bacteriophage or cell. Each bacteriophage or cell contains the nucleotide sequence encoding the particular displayed peptide sequence. Such methods are described in PCT Patent Publication Nos. 91/17271, 91/18980, 91/19818, and 93/08278. (464) Other systems for generating libraries of peptides have aspects of both in vitro chemical synthesis and recombinant methods. See, PCT Patent Publication Nos. 92/05258, 92/14843, and 96/19256. See also, U.S. Pat. Nos. 5,658,754; and 5,643,768. Peptide display libraries, vector, and screening kits are commercially available from such suppliers as Invitrogen (Carlsbad, Calif.), and Cambridge Antibody Technologies (Cambridgeshire, UK). See, e.g., U.S. Pat. Nos. 4,704,692, 4,939,666, 4,946,778, 5,260,203, 5,455,030, 5,518,889, 5,534,621, 5,656,730, 5,763,733, 5,767,260, 5,856,456, assigned to Enzon; U.S. Pat. Nos. 5,223,409, 5,403,484, 5,571,698, 5,837,500, assigned to Dyax, U.S. Pat. Nos. 5,427,908, 5,580,717, assigned to Affymax; U.S. Pat. No. 5,885,793, assigned to Cambridge Antibody Technologies; U.S. Pat. No. 5,750,373, assigned to Genentech, U.S. Pat. Nos. 5,618,920, 5,595,898, 5,576,195, 5,698,435, 5,693,493, 5,698,417, assigned to Xoma, Colligan, supra; Ausubel, supra; or Sambrook, supra. (465) The ARRs/LRRs of the disclosure comprising one or more of a protein scaffold, an antibody, an ScFv, a Centyrin, a single domain antibody, a VHH or a VH of the disclosure can bind human or other mammalian proteins with a wide range of affinities (KD). In a preferred embodiment, at least one ARR/LRR can optionally bind to a target protein with high affinity, for example, with a KD equal to or less than about 10.sup.-7 M, such as but not limited to, 0.1-9.9 (or any range or value therein)×10.sup.-8, 10.sup.-9, 10.sup.-10, 10.sup.-11, 10.sup.-12, 10.sup.-13, 10.sup.-14, 10.sup.-15 or any range or value therein, as determined by surface plasmon resonance or the Kinexa method, as practiced by those of skill in the art. In a preferred embodiment, at least one protein scaffold, antibody, ScFv, Centyrin, single domain antibody, VHH or VH of the disclosure can optionally bind to a target protein with high affinity, for example, with a KD equal to or less than about 10.sup. –7 M, such as but not limited to, 0.1-9.9 (or any range or value therein)×10.sup.-8, 10.sup.-9, 10.sup.-10, 10.sup.-11, 10.sup.-12, 10.sup.-13, 10.sup. -14, 10.sup.-15 or any range or value therein, as determined by surface plasmon resonance or the Kinexa

method, as practiced by those of skill in the art.

(466) The affinity or avidity of a protein scaffold, an antibody, an ScFv, a Centyrin, a single domain antibody, a VHH or a VH of the disclosure for an antigen/ligand can be determined experimentally using any suitable method. (See, for example, Berzofsky, et al., "Antibody-Antigen Interactions," In Fundamental Immunology. Paul, W. E., Ed., Raven Press: New York. N.Y. (1984); Kuby, Janis Immunology, W.H. Freeman and Company: New York, N.Y. (1992); and methods described herein). The measured affinity of a particular protein-antigen/ligand interaction can vary if measured under different conditions (e.g., salt concentration. pH). Thus, measurements of affinity and other antigen-binding parameters (e.g., KD, K.sub.on, K.sub.off) are preferably made with standardized solutions of protein scaffold (e.g. VHH) and antigen, and a standardized buffer, such as the buffer described herein. (467) Competitive assays can be performed with the protein scaffold, antibody, ScFv. Centyrin, single domain antibody, VHH or VH of the disclosure in order to determine what proteins, antibodies, and other antagonists compete for binding to a target protein and/or share the epitope region. These assays as readily known to those of ordinary skill in the art evaluate competition between antagonists or ligands for a limited number of binding sites on a protein. The protein and/or antibody is immobilized or insolubilized before or after the competition and the sample bound to the target protein is separated from the unbound sample, for example, by decanting (where the protein/antibody was preinsolubilized) or by centrifuging (where the protein/antibody was precipitated after the competitive reaction). Also, the competitive binding may be determined by whether function is altered by the binding or lack of binding of the protein scaffold, antibody, ScFv, Centyrin, single domain antibody, VHH or VH to the target protein, e.g., whether protein scaffold, antibody, ScFv, Centyrin, single domain antibody, VHH or VH inhibits or potentiates the enzymatic activity of, for example, a label. ELISA and other functional assays may be used, as well known in the art.

## (468) Therapeutic Proteins

(469) In certain embodiments of the disclosure. T cells are modified to express therapeutic proteins, including secreted human proteins. These secreted proteins may be used as a monotherapy or in combination with another therapy in the treatment or prevention of any disease or disorder. These secreted proteins may be used as a monotherapy or in combination with another therapy for enzyme replacement and/or administration of biologic therapeutics. A database of human secreted proteins can be found at proteinatlas.org/search/protein\_class:Predicted%20secreted%20proteins, the contents of which are incorporated herein by reference. Exemplary human therapeutic proteins can be found, but are not limited to the human proteins in Table 1.

(470) TABLE-US-00132 TABLE 1 Exemplary therapeutic proteins (and proteins to enhance CAR-T efficacy). Compositions of the disclosure may comprise a promoter of one or more of the proteins of Table 1 driving expression of any sequence of the disclosure. Gene Name Gene Description Protein SEQ ID NO A1BG Alpha-1-B glycoprotein SEQ ID NOS: 1-2 A2M Alpha-2-macroglobulin SEQ ID NOS: 3-6 A2ML1 Alpha-2-macroglobulin-like 1 SEQ ID NOS: 7-12 A4GNT Alpha-1,4-Nacetylglucosaminyltransferase SEQ ID NO: 13 AADACL2 Arylacetamide deacetylase-like 2 SEQ ID NOS: 14-15 AANAT Aralkylamine N-acetyltransferase SEQ ID NOS: 16-19 ABCG1 ATP-binding cassette, sub-family G (WHITE), SEQ ID NOS: 20-26 member 1 ABHD1 Abhydrolase domain containing 1 SEQ ID NOS: 27-31 ABHD10 Abhydrolase domain containing 10 SEQ ID NOS: 32-35 ABHD14A Abhydrolase domain containing 14A SEQ ID NOS: 36-40 ABHD15 Abhydrolase domain containing 15 SEQ ID NO: 41 ABI3BP ABI family, member 3 (NESH) binding protein SEQ ID NOS: 42-63 FAM175A Family with sequence similarity 175, member A SEQ ID NOS: 64-71 LA16c- SEQ ID NO: 72 380H5.3 AC008641.1 SEQ ID NO: 73 CTB- SEQ ID NOS: 74-75 60B18.6 AC009133.22 SEQ ID NO: 76 AC009491.2 SEQ ID NO: 77 RP11- SEQ ID NOS: 78-80 977G19.10 CTD- SEQ ID NOS: 81-84 2370N5.3 RP11- SEO ID NOS: 85-87 196G11.1 AC136352.5 SEO ID NO: 88 RP11- SEO ID NO: 89 812E19.9 AC145212.4 MaFF-interacting protein SEQ ID NO: 90 AC233755.1 SEQ ID NO: 91 AC011513.3 SEQ ID NOS: 92-93 ACACB Acetyl-CoA carboxylase beta SEQ ID NOS: 94-100 ACAN Aggrecan SEQ ID NOS: 101-108 ACE Angiotensin I converting enzyme SEQ ID NOS: 109-121 ACHE Acetylcholinesterase (Yt blood group) SEQ ID NOS: 122-134 ACP2 Acid phosphatase 2, lysosomal SEQ ID NOS: 135-142 ACP5 Acid phosphatase 5, tartrate resistant SEQ ID NOS: 143-151 ACP6 Acid phosphatase 6, lysophosphatidic SEQ ID NOS: 152-158 PAPL Iron/zinc purple acid phosphatase-like

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protein SEQ ID NOS: 159-162 ACPP Acid phosphatase, prostate SEQ ID NOS: 163-167 ACR Acrosin
SEQ ID NOS: 168-169 ACRBP Acrosin binding protein SEQ ID NOS: 170-174 ACRV1 Acrosomal
vesicle protein 1 SEQ ID NOS: 175-178 ACSF2 Acyl-CoA synthetase family member 2 SEQ ID NOS:
179-187 ACTL10 Actin-like 10 SEQ ID NO: 188 ACVR1 Activin A receptor, type I SEQ ID NOS: 189-
197 ACVR1C Activin A receptor, type IC SEQ ID NOS: 198-201 ACVRL1 Activin A receptor type II-like
1 SEQ ID NOS: 202-207 ACYP1 Acylphosphatase 1, erythrocyte (common) type SEQ ID NOS: 208-213
ACYP2 Acylphosphatase 2, muscle type SEQ ID NOS: 214-221 CECR1 Cat eye syndrome chromosome
region, candidate 1 SEQ ID NOS: 222-229 ADAM10 ADAM metallopeptidase domain 10 SEQ ID NOS:
230-237 ADAM12 ADAM metallopeptidase domain 12 SEQ ID NOS: 238-240 ADAM15 ADAM
metallopeptidase domain 15 SEQ ID NOS: 241-252 ADAM17 ADAM metallopeptidase domain 17 SEQ
ID NOS: 253-255 ADAM18 ADAM metallopeptidase domain 18 SEQ ID NOS: 256-260 ADAM22
ADAM metallopeptidase domain 22 SEQ ID NOS: 261-269 ADAM28 ADAM metallopeptidase domain
28 SEQ ID NOS: 270-275 ADAM29 ADAM metallopeptidase domain 29 SEQ ID NOS: 276-284
ADAM32 ADAM metallopeptidase domain 32 SEQ ID NOS: 285-291 ADAM33 ADAM
metallopeptidase domain 33 SEQ ID NOS: 292-296 ADAM7 ADAM metallopeptidase domain 7 SEQ ID
NOS: 297-300 ADAM8 ADAM metallopeptidase domain 8 SEQ ID NOS: 301-305 ADAM9 ADAM
metallopeptidase domain 9 SEQ ID NOS: 306-311 ADAMDEC1 ADAM-like, decysin 1 SEQ ID NOS:
312-314 ADAMTS1 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 315-318 type 1 motif,
1 ADAMTS10 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 319-324 type 1 motif, 10
ADAMTS12 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 325-327 type 1 motif, 12
ADAMTS13 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 328-335 type 1 motif, 13
ADAMTS14 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 336-337 type 1 motif, 14
ADAMTS15 ADAM metallopeptidase with thrombospondin SEQ ID NO: 338 type 1 motif, 15
ADAMTS16 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 339-340 type 1 motif, 16
ADAMTS17 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 341-344 type 1 motif, 17
ADAMTS18 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 345-348 type 1 motif, 18
ADAMTS19 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 349-352 type 1 motif, 19
ADAMTS2 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 353-355 type 1 motif, 2
ADAMTS20 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 356-359 type 1 motif, 20
ADAMTS3 ADAM metallopeptidase with thrombospondin SEQ ID NOS: 360-361 type 1 motif, 3
ADAMTS5 ADAM metallopeptidase with thrombospondin SEQ ID NO: 362 type 1 motif, 5 ADAMTS6
ADAM metallopeptidase with thrombospondin SEQ ID NOS: 363-364 type 1 motif, 6 ADAMTS7 ADAM
metallopeptidase with thrombospondin SEQ ID NO: 365 type 1 motif, 7 ADAMTS8 ADAM
metallopeptidase with thrombospondin SEQ ID NO: 366 type 1 motif, 8 ADAMTS9 ADAM
metallopeptidase with thrombospondin SEQ ID NOS: 367-371 type 1 motif, 9 ADAMTSL1 ADAMTS-
like 1 SEQ ID NOS: 372-382 ADAMTSL2 ADAMTS-like 2 SEQ ID NOS: 383-385 ADAMTSL3
ADAMTS-like 3 SEQ ID NOS: 386-387 ADAMTSL4 ADAMTS-like 4 SEQ ID NOS: 388-391
ADAMTSL5 ADAMTS-like 5 SEQ ID NOS: 392-397 ADCK1 AarF domain containing kinase 1 SEQ ID
NOS: 398-402 ADCYAP1 Adenylate cyclase activating polypeptide 1 SEQ ID NOS: 403-404 (pituitary)
ADCYAP1R1 Adenylate cyclase activating polypeptide 1 SEQ ID NOS: 405-411 (pituitary) receptor type
I ADGRA3 Adhesion G protein-coupled receptor A3 SEQ ID NOS: 412-416 ADGRB2 Adhesion G
protein-coupled receptor B2 SEQ ID NOS: 417-425 ADGRD1 Adhesion G protein-coupled receptor D1
SEQ ID NOS: 426-431 ADGRE3 Adhesion G protein-coupled receptor E3 SEQ ID NOS: 432-436
ADGRE5 Adhesion G protein-coupled receptor E5 SEQ ID NOS: 437-442 ADGRF1 Adhesion G protein-
coupled receptor F1 SEQ ID NOS: 443-447 ADGRG1 Adhesion G protein-coupled receptor G1 SEQ ID
NOS: 448-512 ADGRG5 Adhesion G protein-coupled receptor G5 SEQ ID NOS: 513-515 ADGRG6
Adhesion G protein-coupled receptor G6 SEQ ID NOS: 516-523 ADGRV1 Adhesion G protein-coupled
receptor V1 SEQ ID NOS: 524-540 ADI1 Acireductone dioxygenase 1 SEQ ID NOS: 541-543 ADIG
Adipogenin SEQ ID NOS: 544-547 ADIPOQ Adiponectin, C1Q and collagen domain containing SEQ ID
NOS: 548-549 ADM Adrenomedullin SEQ ID NOS: 550-557 ADM2 Adrenomedullin 2 SEQ ID NOS:
558-559 ADM5 Adrenomedullin 5 (putative) SEQ ID NO: 560 ADPGK ADP-dependent glucokinase SEQ
ID NOS: 561-570 ADPRHL2 ADP-ribosylhydrolase like 2 SEQ ID NO: 571 AEBP1 AE binding protein 1
SEQ ID NOS: 572-579 LACE1 Lactation elevated 1 SEQ ID NOS: 580-583 AFM Afamin SEQ ID NO:
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584 AFP Alpha-fetoprotein SEQ ID NOS: 585-586 AGA Aspartylglucosaminidase SEQ ID NOS: 587-589
AGER Advanced glycosylation end product-specific SEQ ID NOS: 590-600 receptor AGK Acylglycerol
kinase SEQ ID NOS: 601-606 AGPS Alkylglycerone phosphate synthase SEQ ID NOS: 607-610 AGR2
Anterior gradient 2, protein disulphide isomerase SEQ ID NOS: 611-614 family member AGR3 Anterior
gradient 3, protein disulphide isomerase SEQ ID NOS: 615-617 family member AGRN Agrin SEQ ID
NOS: 618-621 AGRP Agouti related neuropeptide SEQ ID NO: 622 AGT Angiotensinogen (serpin
peptidase inhibitor, clade SEQ ID NO: 623 A, member 8) AGTPBP1 ATP/GTP binding protein 1 SEQ ID
NOS: 624-627 AGTRAP Angiotensin II receptor-associated protein SEQ ID NOS: 628-635 AHCYL2
Adenosylhomocysteinase-like 2 SEQ ID NOS: 636-642 AHSG Alpha-2-HS-glycoprotein SEQ ID NOS:
643-644 AIG1 Androgen-induced 1 SEQ ID NOS: 645-653 AK4 Adenylate kinase 4 SEQ ID NOS: 654-
657 AKAP10 A kinase (PRKA) anchor protein 10 SEQ ID NOS: 658-666 AKR1C1 Aldo-keto reductase
family 1, member C1 SEQ ID NOS: 667-669 RP4- SEQ ID NOS: 670-672 576H24.4 SERPINA3 Serpin
peptidase inhibitor, clade A (alpha-1 SEQ ID NO: 673 antiproteinase, antitrypsin), member 3 RP11- SEQ
ID NOS: 674-675 14J7.7 RP11- SEQ ID NO: 676 903H12.5 AL356289.1 SEQ ID NO: 677 AL589743.1
SEQ ID NO: 678 XXbac- SEQ ID NOS: 679-680 BPG116M5.17 XXbac- SEQ ID NO: 681
BPG181M17.5 XXbac- SEQ ID NO: 682 BPG32J3.20 RP11- SEQ ID NO: 683 350O14.18 ALAS2 5'-
aminolevulinate synthase 2 SEQ ID NOS: 684-691 ALB Albumin SEQ ID NOS: 692-701 ALDH9A1
Aldehyde dehydrogenase 9 family, member A1 SEQ ID NO: 702 ALDOA Aldolase A, fructose-
bisphosphate SEQ ID NOS: 703-717 ALG1 ALG1, chitobiosyldiphosphodolichol beta- SEQ ID NOS:
718-723 mannosyltransferase ALG5 ALG5, dolichyl-phosphate beta- SEQ ID NOS: 724-725
glucosyltransferase ALG9 ALG9, alpha-1,2-mannosyltransferase SEQ ID NOS: 726-736 FAM150A
Family with sequence similarity 150, member A SEQ ID NOS: 737-738 FAM150B Family with sequence
similarity 150, member B SEQ ID NOS: 739-745 ALKBH1 AlkB homolog 1, histone H2A dioxygenase
SEQ ID NOS: 746-748 ALKBH5 AlkB homolog 5, RNA demethylase SEQ ID NOS: 749-750 ALP1
Alkaline phosphatase, intestinal SEQ ID NOS: 751-752 ALPL Alkaline phosphatase, liver/bone/kidney
SEQ ID NOS: 753-757 ALPP Alkaline phosphatase, placental SEQ ID NO: 758 ALPPL2 Alkaline
phosphatase, placental-like 2 SEQ ID NO: 759 AMBN Ameloblastin (enamel matrix protein) SEQ ID
NOS: 760-762 AMBP Alpha-1-microglobulin/bikunin precursor SEQ ID NOS: 763-765 AMELX
Amelogenin, X-linked SEQ ID NOS: 766-768 AMELY Amelogenin, Y-linked SEQ ID NOS: 769-770
AMH Anti-Mnllerian hormone SEQ ID NO: 771 AMPD1 Adenosine monophosphate deaminase 1 SEQ
ID NOS: 772-774 AMTN Amelotin SEQ ID NOS: 775-776 AMY1A Amylase, alpha 1A (salivary) SEQ
ID NOS: 777-779 AMY1B Amylase, alpha 1B (salivary) SEQ ID NOS: 780-783 AMY1C Amylase, alpha
1C (salivary) SEQ ID NO: 784 AMY2A Amylase, alpha 2A (pancreatic) SEQ ID NOS: 785-787 AMY2B
Amylase, alpha 2B (pancreatic) SEQ ID NOS: 788-792 ANG Angiogenin, ribonuclease, RNase A family,
5 SEQ ID NOS: 793-794 ANGEL1 Angel homolog 1 (Drosophila) SEQ ID NOS: 795-798 ANGPT1
Angiopoietin 1 SEQ ID NOS: 799-803 ANGPT2 Angiopoietin 2 SEQ ID NOS: 804-807 ANGPT4
Angiopoietin 4 SEQ ID NO: 808 ANGPTL1 Angiopoietin-like 1 SEQ ID NOS: 809-811 ANGPTL2
Angiopoietin-like 2 SEQ ID NOS: 812-813 ANGPTL3 Angiopoietin-like 3 SEQ ID NO: 814 ANGPTL4
Angiopoietin-like 4 SEQ ID NOS: 815-822 ANGPTL5 Angiopoietin-like 5 SEQ ID NOS: 823-824
ANGPTL6 Angiopoietin-like 6 SEQ ID NOS: 825-827 ANGPTL7 Angiopoietin-like 7 SEQ ID NO: 828
C19orf80 Chromosome 19 open reading frame 80 SEQ ID NOS: 829-832 ANK1 Ankyrin 1, erythrocytic
SEQ ID NOS: 833-843 ANKDD1A Ankyrin repeat and death domain containing 1A SEQ ID NOS: 844-
850 ANKRD54 Ankyrin repeat domain 54 SEQ ID NOS: 851-859 ANKRD60 Ankyrin repeat domain 60
SEQ ID NO: 860 ANO7 Anoctamin 7 SEQ ID NOS: 861-864 ANOS1 Anosmin 1 SEQ ID NO: 865
ANTXR1 Anthrax toxin receptor 1 SEQ ID NOS: 866-869 AOAH Acyloxyacyl hydrolase (neutrophil)
SEQ ID NOS: 870-874 AOC1 Amine oxidase, copper containing 1 SEQ ID NOS: 875-880 AOC2 Amine
oxidase, copper containing 2 (retina- SEQ ID NOS: 881-882 specific) AOC3 Amine oxidase, copper
containing 3 SEQ ID NOS: 883-889 AP000721.4 SEQ ID NO: 890 APBB1 Amyloid beta (A4) precursor
protein-binding, SEQ ID NOS: 891-907 family B, member I (Fe65) APCDD1 Adenomatosis polyposis
coli down-regulated 1 SEQ ID NOS: 908-913 APCS Amyloid P component, serum SEQ ID NO: 914
APELA Apelin receptor early endogenous ligand SEQ ID NOS: 915-917 APLN Apelin SEQ ID NO: 918
APLP2 Amyloid beta (A4) precursor-like protein 2 SEQ ID NOS: 919-928 APOA1 Apolipoprotein A-I
SEQ ID NOS: 929-933 APOA2 Apolipoprotein A-II SEQ ID NOS: 934-942 APOA4 Apolipoprotein A-IV
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SEQ ID NO: 943 APOA5 Apolipoprotein A-V SEQ ID NOS: 944-946 APOB Apolipoprotein B SEQ ID
NOS: 947-948 APOC1 Apolipoprotein C-I SEQ ID NOS: 949-957 APOC2 Apolipoprotein C-II SEQ ID
NOS: 958-962 APOC3 Apolipoprotein C-III SEQ ID NOS: 963-966 APOC4 Apolipoprotein C-IV SEQ ID
NOS: 967-968 APOC4- APOC4-APOC2 readthrough (NMD candidate) SEQ ID NOS: 969-970 APOC2
APOD Apolipoprotein D SEQ ID NOS: 971-974 APOE Apolipoprotein E SEQ ID NOS: 975-978 APOF
Apolipoprotein F SEQ ID NO: 979 APOH Apolipoprotein H (beta-2-glycoprotein I) SEQ ID NOS: 980-
983 APOL1 Apolipoprotein L, 1 SEQ ID NOS: 984-994 APOL3 Apolipoprotein L, 3 SEQ ID NOS: 995-
1009 APOM Apolipoprotein M SEQ ID NOS: 1010-1012 APOOL Apolipoprotein O-like SEQ ID NOS:
1013-1015 ARCN1 Archain 1 SEQ ID NOS: 1016-1020 ARFIP2 ADP-ribosylation factor interacting
protein 2 SEQ ID NOS: 1021-1027 ARHGAP36 Rho GTPase activating protein 36 SEQ ID NOS: 1028-
1033 HMHA1 Histocompatibility (minor) HA-1 SEQ ID NOS: 1034-1042 ARHGAP6 Rho GTPase
activating protein 6 SEQ ID NOS: 1043-1048 ARIIGEF4 Rho guanine nucleotide exchange factor (GEF)
4 SEQ ID NOS: 1049-1059 ARL16 ADP-ribosylation factor-like 16 SEQ ID NOS: 1060-1068 ARMC5
Armadillo repeat containing 5 SEQ ID NOS: 1069-1075 ARNTL Aryl hydrocarbon receptor nuclear
translocator-like SEQ ID NOS: 1076-1090 ARSA Arylsulfatase A SEQ ID NOS: 1091-1096 ARSB
Arylsulfatase B SEQ ID NOS: 1097-1100 ARSE Arylsulfatase E (chondrodysplasia punctata 1) SEQ ID
NOS: 1101-1104 ARSG Arylsulfatase G SEQ ID NOS: 1105-1108 ARSI Arylsulfatase family, member I
SEQ ID NOS: 1109-1111 ARSK Arylsulfatase family, member K SEQ ID NOS: 1112-1116 ARTS ADP-
ribosyltransferase 3 SEQ ID NOS: 1117-1124 ART4 ADP-ribosyltransferase 4 (Dombrock blood group)
SEQ ID NOS: 1125-1128 ART5 ADP-ribosyltransferase 5 SEQ ID NOS: 1129-1133 ARTN Artemin SEQ
ID NOS: 1134-1144 ASAH1 N-acylsphingosine amidohydrolase (acid SEQ ID NOS: 1145-1195
ceramidase) 1 ASAH2 N-acylsphingosine amidohydrolase (non-lysosomal SEQ ID NOS: 1196-1201
ceramidase) 2 ASCL1 Achaete-scute family bHLH transcription factor 1 SEQ ID NO: 1202 ASIP Agouti
signaling protein SEQ ID NOS: 1203-1204 ASPN Asporin SEQ ID NOS: 1205-1206 ASTL Astacin-like
metallo-endopeptidase (M12 family) SEQ ID NO: 1207 ATAD5 ATPase family, AAA domain containing 5
SEQ ID NOS: 1208-1209 ATAT1 Alpha tubulin acetyltransferase 1 SEQ ID NOS: 1210-1215 ATG2A
Autophagy related 2A SEQ ID NOS: 1216-1218 ATG5 Autophagy related 5 SEQ ID NOS: 1219-1227
ATMIN ATM interactor SEQ ID NOS: 1228-1231 ATP13A1 ATPase type 13A1 SEQ ID NOS: 1232-1234
ATP5F1 ATP synthase, H+ transporting, mitochondrial Fo SEQ ID NOS: 1235-1236 complex, subunit B1
ATP6AP1 ATPase, H+ transporting, lysosomal accessory SEQ ID NOS: 1237-1244 protein 1 ATP6AP2
ATPase, H+ transporting, lysosomal accessory SEQ ID NOS: 1245-1267 protein 2 ATPAF1 ATP synthase
mitochondrial F1 complex assembly SEQ ID NOS: 1268-1278 factor 1 AUH AU RNA binding
protein/enoyl-CoA hydratase SEQ ID NOS: 1279-1280 AVP Arginine vasopressin SEQ ID NO: 1281
AX1N2 Axin 2 SEQ ID NOS: 1282-1289 AZGP1 Alpha-2-glycoprotein 1, zinc-binding SEQ ID NOS:
1290-1292 AZU1 Azurocidin 1 SEQ ID NOS: 1293-1294 B2M Beta-2-microglobulin SEQ ID NOS:
1295-1301 B3GALNT1 Beta-1,3-N-acetylgalactosaminyltransferase 1 SEQ ID NOS: 1302-1314
(globoside blood group) B3GALNT2 Beta-1,3-N-acetylgalactosaminyltransferase 2 SEQ ID NOS: 1315-
1317 B3GALT1 UDP-Gal:betaGlcNAc beta 1,3- SEQ ID NO: 1318 galactosyltransferase, polypeptide 1
B3GALT4 UDP-Gal:betaGlcNAc beta 1,3- SEQ ID NO: 1319 galactosyltransferase, polypeptide 4
B3GALT5 UDP-Gal:betaGlcNAc beta 1,3- SEQ ID NOS: 1320-1324 galactosyltransferase, polypeptide 5
B3GALT6 UDP-Gal:betaGal beta 1,3-galactosyltransferase SEQ ID NO: 1325 polypeptide 6 B3GAT3
Beta-1,3-glucuronyltransferase 3 SEQ ID NOS: 1326-1330 B3GLCT Beta 3-glucosyltransferase SEQ ID
NO: 1331 B3GNT3 UDP-GlcNAc:betaGal beta-1,3-N- SEQ ID NOS: 1332-1335
acetylglucosaminyltransferase 3 B3GNT4 UDP-GlcNAc:betaGal beta-1,3-N- SEQ ID NOS: 1336-1339
acetylglucosaminyltransferase 4 B3GNT6 UDP-GlcNAc:betaGal beta-1,3-N- SEQ ID NOS: 1340-1341
acetylglucosaminyltransferase 6 B3GNT7 UDP-GlcNAc:betaGal beta-1,3-N- SEQ ID NO: 1342
acetylglucosaminyltransferase 7 B3GNT8 UDP-GlcNAc:betaGal beta-1,3-N- SEQ ID NO: 1343
acetylglucosaminyltransferase 8 B3GNT9 UDP-GlcNAc:betaGal beta-1,3-N- SEQ ID NO: 1344
acetylglucosaminyltransferase 9 B4GALNT1 Beta-1,4-N-acetyl-galactosaminyl transferase 1 SEQ ID
NOS: 1345-1356 B4GALNT3 Beta-1,4-N-acetyl-galactosaminyl transferase 3 SEQ ID NOS: 1357-1358
B4GALNT4 Beta-1,4-N-acetyl-galactosaminyl transferase 4 SEQ ID NOS: 1359-1361 B4GALT4 UDP-
Gal:betaGlcNAc beta 1,4- SEQ ID NOS: 1362-1375 galactosyltransferase, polypeptide 4 B4GALT5 UDP-
Gal:betaGlcNAc beta 1,4- SEQ ID NO: 1376 galactosyltransferase, polypeptide 5 B4GALT6 UDP-
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Gal:betaGlcNAc beta 1,4- SEQ ID NOS: 1377-1380 galactosyltransferase, polypeptide 6 B4GAT1 Beta-
1,4-glucuronyltransferase 1 SEQ ID NO: 1381 B9D1 B9 protein domain 1 SEQ ID NOS: 1382-1398
BACE2 Beta-site APP-cleaving enzyme 2 SEQ ID NOS: 1399-1401 BAGE5 B melanoma antigen family,
member 5 SEQ ID NO: 1402 BCAM Basal cell adhesion molecule (Lutheran blood SEQ ID NOS: 1403-
1406 group) BCAN Brevican SEQ ID NOS: 1407-1413 BCAP29 B-cell receptor-associated protein 29
SEQ ID NOS: 1414-1426 BCAR1 Breast cancer anti-estrogen resistance 1 SEQ ID NOS: 1427-1444
BCHE Butyrylcholinesterase SEQ ID NOS: 1445-1449 BCKDHB Branched chain keto acid
dehydrogenase E1, beta SEQ ID NOS: 1450-1452 polypeptide BDNF Brain-derived neurotrophic factor
SEQ ID NOS: 1453-1470 BGLAP Bone gamma-carboxyglutamate (gla) protein SEQ ID NO: 1471 BGN
Biglycan SEQ ID NOS: 1472-1473 BLVRB Biliverdin reductase B SEQ ID NOS: 1474-1478 BMP1 Bone
morphogenetic protein 1 SEQ ID NOS: 1479-1490 BMP10 Bone morphogenetic protein 10 SEQ ID NO:
1491 BMP15 Bone morphogenetic protein 15 SEQ ID NO: 1492 BMP2 Bone morphogenetic protein 2
SEQ ID NO: 1493 BMP3 Bone morphogenetic protein 3 SEQ ID NO: 1494 BMP4 Bone morphogenetic
protein 4 SEQ ID NOS: 1495-1502 BMP6 Bone morphogenetic protein 6 SEQ ID NO: 1503 BMP7 Bone
morphogenetic protein 7 SEQ ID NOS: 1504-1507 BMP8A Bone morphogenetic protein 8a SEQ ID NO:
1508 BMP8B Bone morphogenetic protein 8b SEQ ID NO: 1509 BMPER BMP binding endothelial
regulator SEO ID NOS: 1510-1513 BNC1 Basonuclin 1 SEO ID NOS: 1514-1515 BOC BOC cell
adhesion associated, oncogene regulated SEQ ID NOS: 1516-1526 BOD1 Biorientation of chromosomes
in cell division 1 SEQ ID NOS: 1527-1531 BOLA1 BolA family member 1 SEQ ID NOS: 1532-1534 BPI
Bactericidal/permeability-increasing protein SEQ ID NOS: 1535-1538 BPIFA1 BPI fold containing family
A, member 1 SEQ ID NOS: 1539-1542 BPIFA2 BPI fold containing family A, member 2 SEQ ID NOS:
1543-1544 BPIFA3 BPI fold containing family A. member 3 SEQ ID NOS: 1545-1546 BPIFB1 BPI fold
containing family B, member 1 SEQ ID NOS: 1547-1548 BPIFB2 BPI fold containing family B, member
2 SEQ ID NO: 1549 BPIFB3 BPI fold containing family B, member 3 SEQ ID NO: 1550 BPIFB4 BPI
fold containing family B, member 4 SEQ ID NOS: 1551-1552 BPIFB6 BPI fold containing family B,
member 6 SEQ ID NOS: 1553-1554 BPIFC BPI fold containing family C SEQ ID NOS: 1555-1558 BRF1
BRF1, RNA polymerase III transcription initiation SEQ ID NOS: 1559-1574 factor 90 kDa subunit
BRINP1 Bone morphogenetic protein/retinoic acid inducible SEQ ID NOS: 1575-1576 neural-specific 1
BRINP2 Bone morphogenetic protein/retinoic acid inducible SEQ ID NO: 1577 neural-specific 2 BRINP3
Bone morphogenetic protein/retinoic acid inducible SEQ ID NOS: 1578-1580 neural-specific 3 BSG
Basigin (Ok blood group) SEQ ID NOS: 1581-1591 BSPH1 Binder of sperm protein homolog 1 SEQ ID
NO: 1592 BST1 Bone marrow stromal cell antigen 1 SEQ ID NOS: 1593-1597 BTBD17 BTB (POZ)
domain containing 17 SEQ ID NO: 1598 BTD Biotinidase SEQ ID NOS: 1599-1608 BTN2A2
Butyrophilin, subfamily 2, member A2 SEQ ID NOS: 1609-1622 BTN3A1 Butyrophilin, subfamily 3,
member A1 SEQ ID NOS: 1623-1629 BTN3A2 Butyrophilin, subfamily 3, member A2 SEQ ID NOS:
1630-1640 BTN3A3 Butyrophilin, subfamily 3, member A3 SEQ ID NOS: 1641-1649 RP4- Complement
factor H-related protein 2 SEQ ID NO: 1650 608O15.3 C10orf99 Chromosome 10 open reading frame 99
SEQ ID NO: 1651 C11orf1 Chromosome 11 open reading frame 1 SEQ ID NOS: 1652-1656 C11orf24
Chromosome 11 open reading frame 24 SEQ ID NOS: 1657-1659 C11orf45 Chromosome 11 open reading
frame 45 SEQ ID NOS: 1660-1661 C11orf94 Chromosome 11 open reading frame 94 SEQ ID NO: 1662
C12orf10 Chromosome 12 open reading frame 10 SEQ ID NOS: 1663-1666 C12orf49 Chromosome 12
open reading frame 49 SEQ ID NOS: 1667-1670 C12orf73 Chromosome 12 open reading frame 73 SEQ
ID NOS: 1671-1680 C12orf76 Chromosome 12 open reading frame 76 SEQ ID NOS: 1681-1688
C14orf93 Chromosome 14 open reading frame 93 SEQ ID NOS: 1689-1704 C16orf89 Chromosome 16
open reading frame 89 SEQ ID NOS: 1705-1707 C16orf90 Chromosome 16 open reading frame 90 SEQ
ID NOS: 1708-1709 C17orf67 Chromosome 17 open reading frame 67 SEQ ID NO: 1710 C17orf75
Chromosome 17 open reading frame 75 SEQ ID NOS: 1711-1719 C17orf99 Chromosome 17 open
reading frame 99 SEQ ID NOS: 1720-1722 C18orf54 Chromosome 18 open reading frame 54 SEQ ID
NOS: 1723-1727 C19orf47 Chromosome 19 open reading frame 47 SEQ ID NOS: 1728-1735 C19orf70
Chromosome 19 open reading frame 70 SEQ ID NOS: 1736-1739 C1GALT1 Core 1 synthase,
glycoprotein-N- SEQ ID NOS: 1740-1744 acetylgalactosamine 3-beta-galactosyltransferase 1 C1orf127
Chromosome 1 open reading frame 127 SEQ ID NOS: 1745-1748 C1orf159 Chromosome 1 open reading
frame 159 SEQ ID NOS: 1749-1761 C1orf198 Chromosome 1 open reading frame 198 SEQ ID NOS:
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1762-1766 C1orf54 Chromosome 1 open reading frame 54 SEQ ID NOS: 1767-1769 C1orf56
Chromosome 1 open reading frame 56 SEQ ID NO: 1770 C1QA Complement component 1, q
subcomponent, A SEQ ID NOS: 1771-1773 chain C1QB Complement component 1, q subcomponent, B
SEQ ID NOS: 1774-1777 chain C1QC Complement component 1, q subcomponent, C SEQ ID NOS:
1778-1780 chain C1QL1 Complement component 1, q subcomponent-like 1 SEQ ID NO: 1781 C1QL2
Complement component 1, q subcomponent-like 2 SEQ ID NO: 1782 C1QL3 Complement component 1,
g subcomponent-like 3 SEQ ID NOS: 1783-1784 C1QL4 Complement component 1, g subcomponent-like
4 SEQ ID NO: 1785 C1QTNF1 C1g and tumor necrosis factor related protein 1 SEQ ID NOS: 1786-1795
FAM132A Family with sequence similarity 132, member A SEQ ID NO: 1796 C1QTNF2 C1g and tumor
necrosis factor related protein 2 SEQ ID NO: 1797 C1QTNF3 C1q and tumor necrosis factor related
protein 3 SEQ ID NOS: 1798-1799 C1QTNF4 C1q and tumor necrosis factor related protein 4 SEQ ID
NOS: 1800-1801 C1QTNF5 C1q and tumor necrosis factor related protein 5 SEQ ID NOS: 1802-1804
C1QTNF7 C1q and tumor necrosis factor related protein 7 SEQ ID NOS: 1805-1809 C1QTNF8 C1q and
tumor necrosis factor related protein 8 SEQ ID NOS: 1810-1811 C1QTNF9 C1g and tumor necrosis factor
related protein 9 SEQ ID NOS: 1812-1813 C1QTNF9B C1q and tumor necrosis factor related protein 9B
SEQ ID NOS: 1814-1816 C1R Complement component 1, r subcomponent SEQ ID NOS: 1817-1825
C1RL Complement component 1, r subcomponent-like SEQ ID NOS: 1826-1834 C1S Complement
component 1, s subcomponent SEQ ID NOS: 1835-1844 C2 Complement component 2 SEQ ID NOS:
1845-1859 C21orf33 Chromosome 21 open reading frame 33 SEQ ID NOS: 1860-1868 C21orf62
Chromosome 21 open reading frame 62 SEQ ID NOS: 1869-1872 C22orf15 Chromosome 22 open
reading frame 15 SEQ ID NOS: 1873-1875 C22orf46 Chromosome 22 open reading frame 46 SEQ ID
NO: 1876 C2CD2 C2 calcium-dependent domain containing 2 SEQ ID NOS: 1877-1879 C2orf40
Chromosome 2 open reading frame 40 SEQ ID NOS: 1880-1882 C2orf66 Chromosome 2 open reading
frame 66 SEQ ID NO: 1883 C2orf69 Chromosome 2 open reading frame 69 SEQ ID NO: 1884 C2orf78
Chromosome 2 open reading frame 78 SEQ ID NO: 1885 C3 Complement component 3 SEQ ID NOS:
1886-1890 C3orf33 Chromosome 3 open reading frame 33 SEQ ID NOS: 1891-1895 C3orf58
Chromosome 3 open reading frame 58 SEQ ID NOS: 1896-1899 C4A Complement component 4A
(Rodgers blood SEQ ID NOS: 1900-1901 group) C4B Complement component 4B (Chido blood group)
SEQ ID NOS: 1902-1903 C4BPA Complement component 4 binding protein, alpha SEQ ID NOS: 1904-
1906 C4BPB Complement component 4 binding protein, beta SEQ ID NOS: 1907-1911 C4orf48
Chromosome 4 open reading frame 48 SEQ ID NOS: 1912-1913 C5 Complement component 5 SEQ ID
NO: 1914 C5orf46 Chromosome 5 open reading frame 46 SEQ ID NOS: 1915-1916 C6 Complement
component 6 SEQ ID NOS: 1917-1920 C6orf120 Chromosome 6 open reading frame 120 SEQ ID NO:
1921 C6orf15 Chromosome 6 open reading frame 15 SEQ ID NO: 1922 C6orf58 Chromosome 6 open
reading frame 58 SEQ ID NO: 1923 C7 Complement component 7 SEQ ID NO: 1924 C7orf57
Chromosome 7 open reading frame 57 SEQ ID NOS: 1925-1929 C8A Complement component 8, alpha
polypeptide SEQ ID NO: 1930 C8B Complement component 8, beta polypeptide SEQ ID NOS: 1931-
1933 C8G Complement component 8, gamma polypeptide SEQ ID NOS: 1934-1935 C9 Complement
component 9 SEQ ID NO: 1936 C9orf47 Chromosome 9 open reading frame 47 SEQ ID NOS: 1937-1939
CA10 Carbonic anhydrase X SEQ ID NOS: 1940-1946 CA11 Carbonic anhydrase XI SEQ ID NOS: 1947-
1948 CA6 Carbonic anhydrase VI SEQ ID NOS: 1949-1953 CA9 Carbonic anhydrase IX SEQ ID NOS:
1954-1955 CABLES1 Cdk5 and Abl enzyme substrate 1 SEQ ID NOS: 1956-1961 CABP1 Calcium
binding protein 1 SEQ ID NOS: 1962-1965 CACNA2D1 Calcium channel, voltage-dependent, alpha
2/delta SEQ ID NOS: 1966-1969 subunit 1 CACNA2D4 Calcium channel, voltage-dependent, alpha
2/delta SEQ ID NOS: 1970-1983 subunit 4 CADM3 Cell adhesion molecule 3 SEQ ID NOS: 1984-1986
CALCA Calcitonin-related polypeptide alpha SEQ ID NOS: 1987-1991 CALCB Calcitonin-related
polypeptide beta SEQ ID NOS: 1992-1994 CALCR Calcitonin receptor SEQ ID NOS: 1995-2001
CALCRL Calcitonin receptor-like SEQ ID NOS: 2002-2006 FAM26D Family with sequence similarity
26, member D SEQ ID NOS: 2007-2011 CALR Calreticulin SEQ ID NOS: 2012-2015 CALR3
Calreticulin 3 SEO ID NOS: 2016-2017 CALU Calumenin SEO ID NOS: 2018-2023 CAMK2D
Calcium/calmodulin-dependent protein kinase II SEQ ID NOS: 2024-2035 delta CAMP Cathelicidin
antimicrobial peptide SEQ ID NO: 2036 CANX Calnexin SEQ ID NOS: 2037-2051 CARM1 Coactivator-
associated arginine methyltransferase SEQ ID NOS: 2052-2059 1 CARNS1 Carnosine synthase 1 SEQ ID
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NOS: 2060-2062 CARTPT CART prepropeptide SEQ ID NO: 2063 CASQ1 Calsequestrin 1 (fast-twitch,
skeletal muscle) SEQ ID NOS: 2064-2065 CASQ2 Calsequestrin 2 (cardiac muscle) SEQ ID NO: 2066
CATSPERG Catsper channel auxiliary subunit gamma SEQ ID NOS: 2067-2074 CBLN1 Cerebellin 1
precursor SEQ ID NOS: 2075-2077 CBLN2 Cerebellin 2 precursor SEQ ID NOS: 2078-2081 CBLN3
Cerebellin 3 precursor SEQ ID NOS: 2082-2083 CBLN4 Cerebellin 4 precursor SEQ ID NO: 2084
CCBE1 Collagen and calcium binding EGF domains 1 SEQ ID NOS: 2085-2087 CCDC112 Coiled-coil
domain containing 112 SEQ ID NOS: 2088-2091 CCDC129 Coiled-coil domain containing 129 SEQ ID
NOS: 2092-2099 CCDC134 Coiled-coil domain containing 134 SEQ ID NOS: 2100-2101 CCDC149
Coiled-coil domain containing 149 SEQ ID NOS: 2102-2105 CCDC3 Coiled-coil domain containing 3
SEQ ID NOS: 2106-2107 CCDC80 Coiled-coil domain containing 80 SEQ ID NOS: 2108-2111
CCDC85A Coiled-coil domain containing 85A SEQ ID NO: 2112 CCDC88B Coiled-coil domain
containing 88B SEQ ID NOS: 2113-2115 CCER2 Coiled-coil glutamate-rich protein 2 SEQ ID NOS:
2116-2117 CCK Cholecystokinin SEQ ID NOS: 2118-2120 CCL1 Chemokine (C-C motif) ligand 1 SEQ
ID NO: 2121 CCL11 Chemokine (C-C motif) ligand 11 SEQ ID NO: 2122 CCL13 Chemokine (C-C
motif) ligand 13 SEQ ID NOS: 2123-2124 CCL14 Chemokine (C-C motif) ligand 14 SEQ ID NOS: 2125-
2128 CCL15 Chemokine (C-C motif) ligand 15 SEQ ID NOS: 2129-2130 CCL16 Chemokine (C-C motif)
ligand 16 SEQ ID NOS: 2131-2133 CCL17 Chemokine (C-C motif) ligand 17 SEQ ID NOS: 2134-2135
CCL18 Chemokine (C-C motif) ligand 18 (pulmonary and SEQ ID NO: 2136 activation-regulated)
CCL19 Chemokine (C-C motif) ligand 19 SEQ ID NOS: 2137-2138 CCL2 Chemokine (C-C motif) ligand
2 SEQ ID NOS: 2139-2140 CCL20 Chemokine (C-C motif) ligand 20 SEQ ID NOS: 2141-2143 CCL21
Chemokine (C-C motif) ligand 21 SEQ ID NOS: 2144-2145 CCL22 Chemokine (C-C motif) ligand 22
SEQ ID NO: 2146 CCL23 Chemokine (C-C motif) ligand 23 SEQ ID NOS: 2147-2149 CCL24
Chemokine (C-C motif) ligand 24 SEQ ID NOS: 2150-2151 CCL25 Chemokine (C-C motif) ligand 25
SEQ ID NOS: 2152-2155 CCL26 Chemokine (C-C motif) ligand 26 SEQ ID NOS: 2156-2157 CCL27
Chemokine (C-C motif) ligand 27 SEQ ID NO: 2158 CCL28 Chemokine (C-C motif) ligand 28 SEQ ID
NOS: 2159-2161 CCL3 Chemokine (C-C motif) ligand 3 SEQ ID NO: 2162 CCL3L3 Chemokine (C-C
motif) ligand 3-like 3 SEQ ID NO: 2163 CCL4 Chemokine (C-C motif) ligand 4 SEQ ID NOS: 2164-
2165 CCL4L2 Chemokine (C-C motif) ligand 4-like 2 SEQ ID NOS: 2166-2175 CCL5 Chemokine (C-C
motif) ligand 5 SEQ ID NOS: 2176-2178 CCL7 Chemokine (C-C motif) ligand 7 SEQ ID NOS: 2179-
2181 CCL8 Chemokine (C-C motif) ligand 8 SEQ ID NO: 2182 CCNB1IP1 Cyclin B1 interacting protein
1, E3 ubiquitin SEQ ID NOS: 2183-2194 protein ligase CCNL1 Cyclin L1 SEQ ID NOS: 2195-2203
CCNL2 Cyclin L2 SEQ ID NOS: 2204-2211 CD14 CD14 molecule SEQ ID NOS: 2212-2216 CD160
CD160 molecule SEQ ID NOS: 2217-2221 CD164 CD164 molecule, sialomucin SEQ ID NOS: 2222-
2227 CD177 CD177 molecule SEQ ID NOS: 2228-2230 CD1E CD1e molecule SEQ ID NOS: 2231-2244
CD2 CD2 molecule SEQ ID NOS: 2245-2246 CD200 CD200 molecule SEQ ID NOS: 2247-2253
CD200R1 CD200 receptor 1 SEQ ID NOS: 2254-2258 CD22 CD22 molecule SEQ ID NOS: 2259-2276
CD226 CD226 molecule SEQ ID NOS: 2277-2284 CD24 CD24 molecule SEQ ID NOS: 2285-2291
CD276 CD276 molecule SEQ ID NOS: 2292-2307 CD300A CD300a molecule SEQ ID NOS: 2308-2312
CD300LB CD300 molecule-like family member b SEQ ID NOS: 2313-2314 CD300LF CD300 molecule-
like family member f SEQ ID NOS: 2315-2323 CD300LG CD300 molecule-like family member g SEQ
ID NOS: 2324-2329 CD3D CD3d molecule, delta (CD3-TCR complex) SEQ ID NOS: 2330-2333 CD4
CD4 molecule SEQ ID NOS: 2334-2336 CD40 CD40 molecule, TNF receptor superfamily SEQ ID NOS:
2337-2340 member 5 CD44 CD44 molecule (Indian blood group) SEQ ID NOS: 2341-2367 CD48 CD48
molecule SEQ ID NOS: 2368-2370 CD5 CD5 molecule SEQ ID NOS: 2371-2372 CD55 CD55 molecule,
decay accelerating factor for SEQ ID NOS: 2373-2383 complement (Cromer blood group) CD59 CD59
molecule, complement regulatory protein SEQ ID NOS: 2384-2394 CD5L CD5 molecule-like SEQ ID
NO: 2395 CD6 CD6 molecule SEQ ID NOS: 2396-2403 CD68 CD68 molecule SEQ ID NOS: 2404-2407
CD7 CD7 molecule SEQ ID NOS: 2408-2413 CD79A CD79a molecule, immunoglobulin-associated SEQ
ID NOS: 2414-2416 alpha CD80 CD80 molecule SEQ ID NOS: 2417-2419 CD86 CD86 molecule SEQ
ID NOS: 2420-2426 CD8A CD8a molecule SEQ ID NOS: 2427-2430 CD8B CD8b molecule SEQ ID
NOS: 2431-2436 CD99 CD99 molecule SEQ ID NOS: 2437-2445 CDC23 Cell division cycle 23 SEQ ID
NOS: 2446-2450 CDC40 Cell division cycle 40 SEQ ID NOS: 2451-2453 CDC45 Cell division cycle 45
SEQ ID NOS: 2454-2460 CDCP1 CUB domain containing protein 1 SEQ ID NOS: 2461-2462 CDCP2
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CUB domain containing protein 2 SEQ ID NOS: 2463-2464 CDH1 Cadherin 1, type 1 SEQ ID NOS:
2465-2472 CDH11 Cadherin 11, type 2, OB-cadherin (osteoblast) SEQ ID NOS: 2473-2482 CDH13
Cadherin 13 SEQ ID NOS: 2483-2492 CDH17 Cadherin 17, LI cadherin (liver-intestine) SEQ ID NOS:
2493-2497 CDH18 Cadherin 18, type 2 SEQ ID NOS: 2498-2504 CDH19 Cadherin 19, type 2 SEQ ID
NOS: 2505-2509 CDH23 Cadherin-related 23 SEQ ID NOS: 2510-2525 CDH5 Cadherin 5, type 2
(vascular endothelium) SEQ ID NOS: 2526-2533 CDHR1 Cadherin-related family member 1 SEQ ID
NOS: 2534-2539 CDHR4 Cadherin-related family member 4 SEQ ID NOS: 2540-2544 CDHR5
Cadherin-related family member 5 SEQ ID NOS: 2545-2551 CDKN2A Cyclin-dependent kinase inhibitor
2A SEQ ID NOS: 2552-2562 CONF Cerebral dopamine neurotrophic factor SEQ ID NOS: 2563-2564
CDON Cell adhesion associated, oncogene regulated SEQ ID NOS: 2565-2572 CDSN Corneodesmosin
SEQ ID NO: 2573 CEACAM16 Carcinoembryonic antigen-related cell adhesion SEQ ID NOS: 2574-
2575 molecule 16 CEACAM18 Carcinoembryonic antigen-related cell adhesion SEQ ID NO: 2576
molecule 18 CEACAM19 Carcinoembryonic antigen-related cell adhesion SEQ ID NOS: 2577-2583
molecule 19 CEACAM5 Carcinoembryonic antigen-related cell adhesion SEQ ID NOS: 2584-2591
molecule 5 CEACAM7 Carcinoembryonic antigen-related cell adhesion SEQ ID NOS: 2592-2594
molecule 7 CEACAM8 Carcinoembryonic antigen-related cell adhesion SEQ ID NOS: 2595-2596
molecule 8 CEL Carboxyl ester lipase SEQ ID NO: 2597 CELA2A Chymotrypsin-like elastase family,
member 2A SEQ ID NO: 2598 CELA2B Chymotrypsin-like elastase family, member 2B SEQ ID NOS:
2599-2600 CELA3A Chymotrypsin-like elastase family, member 3A SEQ ID NOS: 2601-2603 CELA3B
Chymotrypsin-like elastase family, member 3B SEQ ID NOS: 2604-2606 CEMIP Cell migration inducing
protein, hyaluronan SEQ ID NOS: 2607-2611 binding CEP89 Centrosomal protein 89 kDa SEQ ID NOS:
2612-2617 CER1 Cerberus 1, DAN family BMP antagonist SEQ ID NO: 2618 CERCAM Cerebral
endothelial cell adhesion molecule SEQ ID NOS: 2619-2626 CERS1 Ceramide synthase 1 SEQ ID NOS:
2627-2631 CES1 Carboxylesterase 1 SEQ ID NOS: 2632-2637 CES3 Carboxylesterase 3 SEQ ID NOS:
2638-2642 CES4A Carboxylesterase 4A SEQ ID NOS: 2643-2648 CES5A Carboxylesterase 5A SEQ ID
NOS: 2649-2656 CETP Cholesteryl ester transfer protein, plasma SEQ ID NOS: 2657-2659 CCDC108
Coiled-coil domain containing 108 SEQ ID NOS: 2660-2669 CFB Complement factor B SEQ ID NOS:
2670-2674 CFC1 Cripto, FRL-1, cryptic family 1 SEQ ID NOS: 2675-2677 CFC1B Cripto, FRL-1,
cryptic family 1B SEQ ID NOS: 2678-2680 CFD Complement factor D (adipsin) SEQ ID NOS: 2681-
2682 CFDP1 Craniofacial development protein 1 SEQ ID NOS: 2683-2686 CFH Complement factor H
SEQ ID NOS: 2687-2689 CFHR1 Complement factor H-related 1 SEQ ID NOS: 2690-2691 CFHR2
Complement factor H-related 2 SEQ ID NOS: 2692-2693 CFHR3 Complement factor H-related 3 SEQ ID
NOS: 2694-2698 CFHR4 Complement factor H-related 4 SEQ ID NOS: 2699-2702 CFHR5 Complement
factor H-related 5 SEQ ID NO: 2703 CFI Complement factor I SEQ ID NOS: 2704-2708 CFP
Complement factor properdin SEQ ID NOS: 2709-2712 CGA Glycoprotein hormones, alpha polypeptide
SEQ ID NOS: 2713-2717 CGB1 Chorionic gonadotropin, beta polypeptide 1 SEQ ID NOS: 2718-2719
CGB2 Chorionic gonadotropin, beta polypeptide 2 SEQ ID NOS: 2720-2721 CGB Chorionic
gonadotropin, beta polypeptide SEQ ID NO: 2722 CGB5 Chorionic gonadotropin, beta polypeptide 5 SEQ
ID NO: 2723 CGB7 Chorionic gonadotropin, beta polypeptide 7 SEQ ID NOS: 2724-2726 CGB8
Chorionic gonadotropin, beta polypeptide 8 SEQ ID NO: 2727 CGREF1 Cell growth regulator with EF-
hand domain 1 SEQ ID NOS: 2728-2735 CHAD Chondroadherin SEQ ID NOS: 2736-2738 CHADL
Chondroadherin-like SEQ ID NOS: 2739-2741 CHEK2 Checkpoint kinase 2 SEQ ID NOS: 2742-2763
CHGA Chromogranin A SEQ ID NOS: 2764-2766 CHGB Chromogranin B SEQ ID NOS: 2767-2768
CHI3L1 Chitinase 3-like 1 (cartilage glycoprotein-39) SEQ ID NOS: 2769-2770 CHI3L2 Chitinase 3-like
2 SEQ ID NOS: 2771-2784 CHIA Chitinase, acidic SEQ ID NOS: 2785-2793 CHID1 Chitinase domain
containing 1 SEQ ID NOS: 2794-2812 CHIT1 Chitinase 1 (chitotriosidase) SEQ ID NOS: 2813-2816
CHL1 Cell adhesion molecule L1-like SEQ ID NOS: 2817-2825 CHN1 Chimerin 1 SEQ ID NOS: 2826-
2836 CHPF Chondroitin polymerizing factor SEQ ID NOS: 2837-2839 CHPF2 Chondroitin polymerizing
factor 2 SEQ ID NOS: 2840-2843 CHRD Chordin SEQ ID NOS: 2844-2849 CHRDL1 Chordin-like 1
SEQ ID NOS: 2850-2854 CHRDL2 Chordin-like 2 SEQ ID NOS: 2.855-2863 CHRNA2 Cholinergic
receptor, nicotinic, alpha 2 (neuronal) SEQ ID NOS: 2864-2872 CHRNA5 Cholinergic receptor, nicotinic,
alpha 5 (neuronal) SEQ ID NOS: 2873-2876 CHRNB1 Cholinergic receptor, nicotinic, beta 1 (muscle)
SEQ ID NOS: 2877-2882 CHRND Cholinergic receptor, nicotinic, delta (muscle) SEQ ID NOS: 2883-
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2888 CHST1 Carbohydrate (keratan sulfate Gal-6) SEQ ID NO: 2889 sulfotransferase 1 CHST10
Carbohydrate sulfotransferase 10 SEQ ID NOS: 2890-2897 CHST11 Carbohydrate (chondroitin 4)
sulfotransferase 11 SEQ ID NOS: 2898-2902 CHST13 Carbohydrate (chondroitin 4) sulfotransferase 13
SEQ ID NOS: 2903-2904 CHST4 Carbohydrate (N-acetylglucosamine 6-O) SEQ ID NOS: 2905-2906
sulfotransferase 4 CHST5 Carbohydrate (N-acetylglucosamine 6-O) SEQ ID NOS: 2907-2908
sulfotransferase 5 CHST6 Carbohydrate (N-acetylglucosamine 6-O) SEQ ID NOS: 2909-2910
sulfotransferase 6 CHST7 Carbohydrate (N-acetylglucosamine 6-O) SEQ ID NO: 2911 sulfotransferase 7
CHST8 Carbohydrate (N-acetylgalactosamine 4-0) SEQ ID NOS: 2912-2915 sulfotransferase 8 CHSY1
Chondroitin sulfate synthase 1 SEQ ID NOS: 2916-2917 CHSY3 Chondroitin sulfate synthase 3 SEQ ID
NO: 2918 CHTF8 Chromosome transmission fidelity factor 8 SEQ ID NOS: 2919-2929 CILP Cartilage
intermediate layer protein, nucleotide SEQ ID NO: 2930 pyrophosphohydrolase CILP2 Cartilage
intermediate layer protein 2 SEQ ID NOS: 2931-2932 CKLF Chemokine-like factor SEQ ID NOS: 2933-
2938 CKMT1A Creatine kinase, mitochondrial 1A SEQ ID NOS: 2939-2944 CKMT1B Creatine kinase,
mitochondrial 1B SEQ ID NOS: 2945-2954 CLCA1 Chloride channel accessory 1 SEQ ID NOS: 2955-
2956 CLCF1 Cardiotrophin-like cytokine factor 1 SEQ ID NOS: 2957-2958 CLDN15 Claudin 15 SEQ ID
NOS: 2959-2964 CLDN7 Claudin 7 SEQ ID NOS: 2.965-2971 CLDND1 Claudin domain containing 1
SEQ ID NOS: 2972-2997 CLEC11A C-type lectin domain family 11, member A SEQ ID NOS: 2998-3000
CLEC16A C-type lectin domain family 16, member A SEQ ID NOS: 3001-3006 CLEC18A C-type lectin
domain family 18, member A SEQ ID NOS: 3007-3012 CLEC18B C-type lectin domain family 18,
member B SEQ ID NOS: 3013-3016 CLEC18C C-type lectin domain family 18, member C SEQ ID NOS:
3017-3023 CLEC19A C-type lectin domain family 19, member A SEQ ID NOS: 3024-3027 CLEC2B C-
type lectin domain family 2, member B SEQ ID NOS: 3028-3029 CLEC3A C-type lectin domain family
3, member A SEQ ID NOS: 3030-3031 CLEC3B C-type lectin domain family 3, member B SEQ ID NOS:
3032-3033 CLGN Calmegin SEQ ID NOS: 3034-3036 CLN5 Ceroid-lipofuscinosis, neuronal 5 SEQ ID
NOS: 3037-3048 CLPS Colipase, pancreatic SEQ ID NOS: 3049-3051 CLPSL1 Colipase-like 1 SEQ ID
NOS: 3052-3053 CLPSL2 Colipase-like 2 SEQ ID NOS: 3054-3055 CLPX Caseinolytic mitochondrial
matrix peptidase SEQ ID NOS: 3056-3058 chaperone subunit CLSTN3 Calsyntenin 3 SEQ ID NOS:
3059-3065 CLU Clusterin SEQ ID NOS: 3066-3079 CLUL1 Clusterin-like 1 (retinal) SEQ ID NOS:
3080-3087 CMA1 Chymase 1, mast cell SEQ ID NOS: 3088-3089 CMPK1 Cytidine monophosphate
(UMP-CMP) kinase 1, SEQ ID NOS: 3090-3093 cytosolic CNBD1 Cyclic nucleotide binding domain
containing 1 SEQ ID NOS: 3094-3097 CNDP1 Carnosine dipeptidase 1 (metallopeptidase M20 SEQ ID
NOS: 3098-3100 family) RQCD1 RCD1 required for cell differentiation1 homolog SEQ ID NOS: 3101-
3107 (S. pombe) CNPY2 Canopy FGF signaling regulator 2 SEQ ID NOS: 3108-3112 CNPY3 Canopy
FGF signaling regulator 3 SEQ ID NOS: 3113-3114 CNPY4 Canopy FGF signaling regulator 4 SEQ ID
NOS: 3115-3117 CNTFR Ciliary neurotrophic factor receptor SEQ ID NOS: 3118-3121 CNTN1
Contactin 1 SEQ ID NOS: 3122-3131 CNTN2 Contactin 2 (axonal) SEQ ID NOS: 3132-3143 CNTN3
Contactin 3 (plasmacytoma associated) SEQ ID NO: 3144 CNTN4 Contactin 4 SEQ ID NOS: 3145-3153
CNTN5 Contactin 5 SEQ ID NOS: 3154-3159 CNTNAP2 Contactin associated protein-like 2 SEQ ID
NOS: 3160-3163 CNTNAP3 Contactin associated protein-like 3 SEQ ID NOS: 3164-3168 CNTNAP3B
Contactin associated protein-like 3B SEQ ID NOS: 3169-3177 COASY CoA synthase SEQ ID NOS:
3178-3187 COCH Cochlin SEQ ID NOS: 3188-3199 COG3 Component of oligomeric golgi complex 3
SEQ ID NOS: 3200-3203 COL10A1 Collagen, type X, alpha 1 SEQ ID NOS: 3204-3207 COL11A1
Collagen, type XI, alpha 1 SEQ ID NOS: 3208-3218 COL11A2 Collagen, type XI, alpha 2 SEQ ID NOS:
3219-3223 COL12A1 Collagen, type XII, alpha 1 SEQ ID NOS: 3224-3231 COL14A1 Collagen, type
XIV, alpha 1 SEQ ID NOS: 3232-3239 COL15A1 Collagen, type XV, alpha 1 SEQ ID NOS: 3240-3241
COL16A1 Collagen, type XVI, alpha 1 SEQ ID NOS: 3242-3246 COL18A1 Collagen, type XVIII, alpha
1 SEQ ID NOS: 3247-3251 COL19A1 Collagen, type XIX, alpha 1 SEQ ID NOS: 3252-3254 COL1A1
Collagen, type I, alpha 1 SEQ ID NOS: 3255-3256 COL1A2 Collagen, type I, alpha 2 SEQ ID NOS:
3257-3258 COL20A1 Collagen, type XX, alpha 1 SEQ ID NOS: 3259-3262 COL21A1 Collagen, type
XXI, alpha 1 SEQ ID NOS: 3263-3268 COL22A1 Collagen, type XXII, alpha 1 SEQ ID NOS: 3269-3271
COL24A1 Collagen, type XXIV, alpha 1 SEQ ID NOS: 3272-3275 COL26A1 Collagen, type XXVI,
alpha 1 SEQ ID NOS: 3276-3277 COL27A1 Collagen, type XXVII, alpha 1 SEQ ID NOS: 3278-3280
COL28A1 Collagen, type XXVIII, alpha 1 SEQ ID NOS: 3281-3285 COL2A1 Collagen, type II, alpha 1
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SEQ ID NOS: 3286-3287 COL3A1 Collagen, type III, alpha 1 SEQ ID NOS: 3288-3290 COL4A1
Collagen, type IV, alpha 1 SEQ ID NOS: 3291-3293 COL4A2 Collagen, type IV, alpha 2 SEQ ID NOS:
3294-3296 COL4A3 Collagen, type IV, alpha 3 (Goodpasture antigen) SEQ ID NOS: 3297-3300 COL4A4
Collagen, type IV, alpha 4 SEQ ID NOS: 3301-3302 COL4A5 Collagen, type IV, alpha 5 SEQ ID NOS:
3303-3309 COL4A6 Collagen, type IV, alpha 6 SEQ ID NOS: 3310-3315 COL5A1 Collagen, type V,
alpha 1 SEQ ID NOS: 3316-3318 COL5A2 Collagen, type V, alpha 2 SEQ ID NOS: 3319-3320 COL5A3
Collagen, type V, alpha 3 SEQ ID NO: 3321 COL6A1 Collagen, type VI, alpha 1 SEQ ID NOS: 3322-
3323 COL6A2 Collagen, type VI, alpha 2 SEQ ID NOS: 3324-3329 COL6A3 Collagen, type VI, alpha 3
SEQ ID NOS: 3330-3338 COL6A5 Collagen, type VI, alpha 5 SEQ ID NOS: 3339-3343 COL6A6
Collagen, type VI, alpha 6 SEQ ID NOS: 3344-3346 COL7A1 Collagen, type VII, alpha 1 SEQ ID NOS:
3347-3348 COL8A1 Collagen, type VIII, alpha 1 SEQ ID NOS: 3349-3352 COL8A2 Collagen, type VIII,
alpha 2 SEQ ID NOS: 3353-3355 COL9A1 Collagen, type IX, alpha 1 SEQ ID NOS: 3356-3359 COL9A2
Collagen, type IX, alpha 2 SEQ ID NOS: 3360-3363 COL9A3 Collagen, type IX, alpha 3 SEQ ID NOS:
3364-3365 COLEC10 Collectin sub-family member 10 (C-type lectin) SEQ ID NO: 3366 COLEC11
Collectin sub-family member 11 SEQ ID NOS: 3367-3376 COLGALT1 Collagen beta(1-
O)galactosyltransferase 1 SEQ ID NOS: 3377-3379 COLGALT2 Collagen beta(1-O)galactosyltransferase
2 SEQ ID NOS: 3380-3382 COLQ Collagen-like tail subunit (single strand of SEQ ID NOS: 3383-3387
homotrimer) of asymmetric acetylcholinesterase COMP Cartilage oligomeric matrix protein SEQ ID NOS:
3388-3390 COPS6 COP9 signalosome subunit 6 SEQ ID NOS: 3391-3394 COQ6 Coenzyme Q6
monooxygenase SEO ID NOS: 3395-3402 CORT Cortistatin SEO ID NO: 3403 CP Ceruloplasmin
(ferroxidase) SEQ ID NOS: 3404-3408 CPA1 Carboxypeptidase A1 (pancreatic) SEQ ID NOS: 3409-3413
CPA2 Carboxy peptidase A2 (pancreatic) SEQ ID NOS: 3414-3415 CPA3 Carboxypeptidase A3 (mast
cell) SEQ ID NO: 3416 CPA4 Carboxypeptidase A4 SEQ ID NOS: 3417-3422 CPA6 Carboxypeptidase
A6 SEQ ID NOS: 3423-3425 CPAMD8 C3 and PZP-like, alpha-2-macroglobulin domain SEQ ID NOS:
3426-3431 containing 8 CPB1 Carboxypeptidase B1 (tissue) SEQ ID NOS: 3432-3436 CPB2
Carboxypeptidase B2 (plasma) SEQ ID NOS: 3437-3439 CPE Carboxypeptidase E SEQ ID NOS: 3440-
3444 CPM Carboxypeptidase M SEQ ID NOS: 3445-3454 CPN1 Carboxypeptidase N, polypeptide 1 SEQ
ID NOS: 3455-3456 CPN2 Carboxypeptidase N, polypeptide 2 SEQ ID NOS: 3457-3458 CPO
Carboxypeptidase O SEQ ID NO: 3459 CPQ Carboxypeptidase Q SEQ ID NOS: 3460-3465 CPVL
Carboxypeptidase, vitellogenic-like SEQ ID NOS: 3466-3476 CPXM1 Carboxypeptidase X (M14 family),
member 1 SEQ ID NO: 3477 CPXM2 Carboxypeptidase X (M14 family), member 2 SEQ ID NOS: 3478-
3479 CPZ Carboxypeptidase Z SEQ ID NOS: 3480-3483 CR1L Complement component (3b/4b) receptor
1-like SEQ ID NOS: 3484-3485 CRB2 Crumbs family member 2 SEQ ID NOS: 3486-3488 CREG1
Cellular repressor of E1A-stimulated genes 1 SEQ ID NO: 3489 CREG2 Cellular repressor of E1A-
stimulated genes 2 SEQ ID NO: 3490 CRELD1 Cysteine-rich with EGF-like domains 1 SEQ ID NOS:
3491-3496 CRELD2 Cysteine-rich with EGF-like domains 2 SEQ ID NOS: 3497-3501 CRH
Corticotropin releasing hormone SEQ ID NO: 3502 CRHBP Corticotropin releasing hormone binding
protein SEQ ID NOS: 3503-3504 CRHR1 Corticotropin releasing hormone receptor 1 SEQ ID NOS:
3505-3516 CRHR2 Corticotropin releasing hormone receptor 2 SEQ ID NOS: 3517-3523 CRISP1
Cysteine-rich secretory protein 1 SEQ ID NOS: 3524-3527 CRISP2 Cysteine-rich secretory protein 2 SEQ
ID NOS: 3528-3530 CRISP3 Cysteine-rich secretory protein 3 SEQ ID NOS: 3531-3534 CRISPLD2
Cysteine-rich secretory protein LCCL domain SEQ ID NOS: 3535-3542 containing 2 CRLF1 Cytokine
receptor-like factor 1 SEQ ID NOS: 3543-3544 CRP C-reactive protein, pentraxin-related SEQ ID NOS:
3545-3549 CRTAC1 Cartilage acidic protein 1 SEQ ID NOS: 3550-3554 CRTAP Cartilage associated
protein SEQ ID NOS: 3555-3556 CRY2 Cryptochrome circadian clock 2 SEQ ID NOS: 3557-3560 CSAD
Cysteine sulfinic acid decarboxylase SEQ ID NOS: 3561-3573 CSF1 Colony stimulating factor 1
(macrophage) SEQ ID NOS: 3574-3581 CSF1R Colony stimulating factor 1 receptor SEQ ID NOS: 3582-
3586 CSF2 Colony stimulating factor 2 (granulocyte- SEQ ID NO: 3587 macrophage) CSF2RA Colony
stimulating factor 2 receptor, alpha, low- SEQ ID NOS: 3588-3599 affinity (granulocyte-macrophage)
CSF3 Colony stimulating factor 3 (granulocyte) SEQ ID NOS: 3600-3606 CSGALNACT1 Chondroitin
sulfate N- SEQ ID NOS: 3607-3615 acetylgalactosaminyltransferase 1 CSH1 Chorionic
somatomammotropin hormone 1 SEQ ID NOS: 3616-3619 (placental lactogen) CSH2 Chorionic
somatomammotropin hormone 2. SEQ ID NOS: 3620-3624 CSHL1 Chorionic somatomammotropin
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hormone-like 1 SEQ ID NOS: 3625-3631 CSN1S1 Casein alpha s1 SEQ ID NOS: 3632-3637 CSN2
Casein beta SEQ ID NO: 3638 CSN3 Casein kappa SEQ ID NO: 3639 CST1 Cystatin SN SEQ ID NOS:
3640-3641 CST11 Cystatin 11 SEQ ID NOS: 3642-3643 CST2 Cystatin SA SEQ ID NO: 3644 CST3
Cystatin C SEQ ID NOS: 3645-3647 CST4 Cystatin S SEQ ID NO: 3648 CST5 Cystatin D SEQ ID NO:
3649 CST6 Cystatin E/M SEQ ID NO: 3650 CST7 Cystatin F (leukocystatin) SEQ ID NO: 3651 CST8
Cystatin 8 (cystatin-related epididymal specific) SEQ ID NOS: 3652-3653 CST9 Cystatin 9 (testatin) SEQ
ID NO: 3654 CST9L Cystatin 9-like SEQ ID NO: 3655 CSTL1 Cystatin-like 1 SEQ ID NOS: 3656-3658
CT55 Cancer/testis antigen 55 SEQ ID NOS: 3659-3660 CTBS Chitobiase, di-N-acetyl- SEQ ID NOS:
3661-3663 CTGF Connective tissue growth factor SEQ ID NO: 3664 CTHRC1 Collagen triple helix
repeat containing 1 SEQ ID NOS: 3665-3668 CTLA4 Cytotoxic T-lymphocyte-associated protein 4 SEQ
ID NOS: 3669-3672 CTNS Cystinosin, lysosomal cystine transporter SEQ ID NOS: 3673-3680 CTRB1
Chymotrypsinogen B1 SEQ ID NOS: 3681-3683 CTRB2 Chymotrypsinogen B2 SEQ ID NOS: 3684-
3687 CTRC Chymotrypsin C (caldecrin) SEQ ID NOS: 3688-3689 CTRL Chymotrypsin-like SEQ ID
NOS: 3690-3692 CTSA Cathepsin A SEQ ID NOS: 3693-3701 CTSB Cathepsin B SEQ ID NOS: 3702-
3726 CTSC Cathepsin C SEQ ID NOS: 3727-3731 CTSD Cathepsin D SEQ ID NOS; 3732-3742 CTSE
Cathepsin E SEQ ID NOS: 3743-3744 CTSF Cathepsin F SEQ ID NOS: 3745-3748 CTSG Cathepsin G
SEQ ID NO: 3749 CTSH Cathepsin H SEQ ID NOS: 3750-3755 CTSK Cathepsin K SEQ ID NOS: 3756-
3757 CTSL Cathepsin L SEQ ID NOS: 3758-3760 CTSO Cathepsin O SEQ ID NO: 3761 CTSS
Cathepsin S SEQ ID NOS: 3762-3766 CTSV Cathepsin V SEQ ID NOS: 3767-3768 CTSW Cathepsin W
SEO ID NOS: 3769-3771 CTSZ Cathepsin Z SEO ID NO: 3772 CUBN Cubilin (intrinsic factor-
cobalamin receptor) SEQ ID NOS: 3773-3776 CUTA CutA divalent cation tolerance homolog (E. coli)
SEQ ID NOS: 3777-3786 CX3CL1 Chemokine (C-X3-C motif) ligand 1 SEQ ID NOS: 3787-3790
CXADR Coxsackie virus and adenovirus receptor SEQ ID NOS: 3791-3795 CXCL1 Chemokine (C-X-C
motif) ligand 1 (melanoma SEQ ID NO: 3796 growth stimulating activity, alpha) CXCL10 Chemokine (C-
X-C motif) ligand 10 SEQ ID NO: 3797 CXCL11 Chemokine (C-X-C motif) ligand 11 SEQ ID NOS:
3798-3799 CXCL12 Chemokine (C-X-C motif) ligand 12 SEQ ID NOS: 3800-3805 CXCL13 Chemokine
(C-X-C motif) ligand 13 SEQ ID NO: 3806 CXCL14 Chemokine (C-X-C motif) ligand 14 SEQ ID NOS:
3807-3808 CXCL17 Chemokine (C-X-C motif) ligand 17 SEQ ID NOS: 3809-3810 CXCL2 Chemokine
(C-X-C motif) ligand 2 SEQ ID NO: 3811 CXCL3 Chemokine (C-X-C motif) ligand 3 SEQ ID NO: 3812
CXCL5 Chemokine (C-X-C motif) ligand 5 SEQ ID NO: 3813 CXCL6 Chemokine (C-X-C motif) ligand
6 SEQ ID NOS: 3814-3815 CXCL8 Chemokine (C-X-C motif) ligand 8 SEQ ID NOS: 3816-3817
CXCL9 Chemokine (C-X-C motif) ligand 9 SEQ ID NO: 3818 CXorf36 Chromosome X open reading
frame 36 SEQ ID NOS: 3819-3820 CYB5D2 Cytochrome b5 domain containing 2 SEQ ID NOS: 3821-
3824 CYHR1 Cysteine/histidine-rich 1 SEQ ID NOS: 3825-3832 CYP17A1 Cytochrome P450, family 17,
subfamily A, SEQ ID NOS: 3833-3837 polypeptide 1 CYP20A1 Cytochrome P450, family 20, subfamily
A, SEQ ID NOS: 3838-3844 polypeptide 1 CYP21A2 Cytochrome P450, family 21, subfamily A, SEQ ID
NOS: 3845-3852 polypeptide 2 CYP26B1 Cytochrome P450, family 26, subfamily B, SEQ ID NOS:
3853-3857 polypeptide 1 CYP2A6 Cytochrome P450, family 2, subfamily A, SEQ ID NOS: 3858-3859
polypeptide 6 CYP2A7 Cytochrome P450, family 2, subfamily A, SEQ ID NOS: 3860-3862 polypeptide 7
CYP2B6 Cytochrome P450, family 2, subfamily B, SEQ ID NOS: 3863-3866 polypeptide 6 CYP2C18
Cytochrome P450, family 2, subfamily C, SEQ ID NOS: 3867-3868 polypeptide 18 CYP2C19
Cytochrome P450, family 2, subfamily C, SEQ ID NOS: 3869-3870 polypeptide 19 CYP2C8 Cytochrome
P450, family 2, subfamily C, SEQ ID NOS: 3871-3878 polypeptide 8 CYP2C9 Cytochrome P450, family
2, subfamily C, SEQ ID NOS: 3879-3881 polypeptide 9 CYP2E1 Cytochrome P450, family 2, subfamily
E, SEQ ID NOS: 3882-3887 polypeptide 1 CYP2F1 Cytochrome P450, family 2, subfamily F, SEQ ID
NOS: 3888-3891 polypeptide 1 CYP2J2 Cytochrome P450, family 2, subfamily J, SEQ ID NO: 3892
polypeptide 2 CYP2R1 Cytochrome P450, family 2, subfamily R, SEQ ID NOS: 3893-3898 polypeptide 1
CYP2S1 Cytochrome P450, family 2, subfamily S, SEQ ID NOS: 3899-3904 polypeptide 1 CYP2W1
Cytochrome P450, family 2, subfamily W, SEQ ID NOS: 3905-3907 polypeptide 1 CYP46A1
Cytochrome P450, family 46, subfamily A, SEQ ID NOS: 3908-3912 polypeptide 1 CYP4F11
Cytochrome P450, family 4, subfamily F, SEQID NOS: 3913-3917 polypeptide 11 CYP4F2 Cytochrome
P450, family 4, subfamily F, SEQ ID NOS: 3918-3922 polypeptide 2 CYR61 Cysteine-rich, angiogenic
inducer, 61 SEQ ID NO: 3923 CYTL1 Cytokine-like 1 SEQ ID NOS: 3924-3926 D2HGDH D-2-
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hydroxyglutarate dehydrogenase SEQ ID NOS: 3927-3935 DAG1 Dystroglycan 1 (dystrophin-associated
SEQ ID NOS: 3936-3950 glycoprotein 1) DAND5 DAN domain family member 5, BMP antagonist SEQ
ID NOS: 3951-3952 DAO D-amino-acid oxidase SEQ ID NOS: 3953-3958 DAZAP2 DAZ associated
protein 2 SEQ ID NOS: 3959-3967 DBH Dopamine beta-hydroxylase (dopamine beta- SEQ ID NOS:
3968-3969 monooxygenase) DBNL Drebrin-like SEQ ID NOS: 3970-3987 DCD Dermcidin SEQ ID
NOS: 3988-3990 DCN Decorin SEQ ID NOS: 3991-4009 DD1AS DNA damage-induced apoptosis
suppressor SEQ ID NOS: 4010-4019 DDOST Dolichyl-diphosphooligosaccharide--protein SEQ ID NOS:
4020-4023 glycosyltransferase subunit (non-catalytic) DDR1 Discoidin domain receptor tyrosine kinase 1
SEQ ID NOS: 4024-4069 DDR2 Discoidin domain receptor tyrosine kinase 2 SEQ ID NOS: 4070-4075
DDT D-dopachrome tautomerase SEQ ID NOS: 4076-4081 DDX17 DEAD (Asp-Glu-Ala-Asp) box
helicase 17 SEQ ID NOS: 4082-4086 DDX20 DEAD (Asp-Glu-Ala-Asp) box polypeptide 20 SEQ ID
NOS: 4087-4089 DDX25 DEAD (Asp-Glu-Ala-Asp) box helicase 25 SEQ ID NOS: 4090-4096 DDX28
DEAD (Asp-Glu-Ala-Asp) box polypeptide 28 SEQ ID NO: 4097 DEAE1 DEAF1 transcription factor
SEQ ID NOS: 4098-4100 DEF8 Differentially expressed in FDCP 8 homolog SEQ ID NOS: 4101-4120
(mouse) DEFA1 Defensin, alpha 1 SEQ ID NOS: 4121-4122 DEFA1B Defensin, alpha 1B SEQ ID NO:
4123 DEFA3 Defensin, alpha 3, neutrophil-specific SEQ ID NO: 4124 DEFA4 Defensin, alpha 4,
corticostatin SEQ ID NO: 4125 DEFA5 Defensin, alpha 5, Paneth cell-specific SEQ ID NO: 4126 DEFA6
Defensin, alpha 6, Paneth cell-specific SEQ ID NO: 4127 DEFB1 Defensin, beta 1 SEQ ID NO: 4128
DEFB103A Defensin, beta 103A SEQ ID NO: 4129 DEFB103B Defensin, beta 103B SEQ ID NO: 4130
DEFB104A Defensin, beta 104A SEO ID NO: 4131 DEFB104B Defensin, beta 104B SEO ID NO: 4132
DEFB105A Defensin, beta 105A SEQ ID NO: 4133 DEFB105B Defensin, beta 105B SEQ ID NO: 4134
DEFB106A Defensin, beta 106A SEQ ID NO: 4135 DEFB106B Defensin, beta 106B SEQ ID NO: 4136
DEFB107A Defensin, beta 107A SEQ ID NO: 4137 DEFB107B Defensin, beta 107B SEQ ID NO: 4138
DEFB108B Defensin, beta 108B SEQ ID NO: 4139 DEFB110 Defensin, beta 110 SEQ ID NOS: 4140-
4141 DEFB113 Defensin, beta 113 SEQ ID NO: 4142 DEFB114 Defensin, beta 114 SEQ ID NO: 4143
DEFB115 Defensin, beta 115 SEQ ID NO: 4144 DEFB116 Defensin, beta 116 SEQ ID NO: 4145
DEFB118 Defensin, beta 118 SEQ ID NO: 4146 DEFB119 Defensin, beta 119 SEQ ID NOS: 4147-4149
DEFB121 Defensin, beta 121 SEQ ID NO: 4150 DEEB123 Defensin, beta 123 SEQ ID NO: 4151
DEFB124 Defensin, beta 124 SEQ ID NO: 4152 DEFB125 Defensin, beta 125 SEQ ID NO: 4153
DEFB126 Defensin, beta 126 SEQ ID NO: 4154 DEFB127 Defensin, beta 127 SEQ ID NO: 4155
DEEB128 Defensin, beta 128 SEQ ID NO: 4156 DEFB129 Defensin, beta 129 SEQ ID NO: 4157
DEFB130 Defensin, beta 130 SEQ ID NO: 4158 RP11- SEQ ID NO: 4159 1236K1.1 DEFB131 Defensin,
beta 131 SEQ ID NO: 4160 CTD- SEQ ID NO: 4161 2313N18.7 DEFB132 Defensin, beta 132 SEQ ID
NO: 4162 DEFB133 Defensin, beta 133 SEQ ID NO: 4163 DEFB134 Defensin, beta 134 SEQ ID NOS:
4164-4165 DEFB135 Defensin, beta 135 SEQ ID NO: 4166 DEEB136 Defensin, beta 136 SEQ ID NO:
4167 DEFB4A Defensin, beta 4A SEQ ID NO: 4168 DEFB4B Defensin, beta 4B SEQ ID NO: 4169
C10orf10 Chromosome 10 open reading frame 10 SEQ ID NOS: 4170-4171 DGCR2 DiGeorge syndrome
critical region gene 2 SEQ ID NOS: 4172-4175 DHH Desert hedgehog SEQ ID NO: 4176 DHRS4
Dehydrogenase/reductase (SDR family) member 4 SEQ ID NOS: 4177-4184 DHRS4L2
Dehydrogenase/reductase (SDR family) member 4 SEQ ID NOS: 4185-4194 like 2 DHRS7
Dehydrogenase/reductase (SDR family) member 7 SEQ ID NOS: 4195-4202 DHRS7C
Dehvdrogenase/reductase (SDR family) member SEQ ID NOS: 4203-4205 7C DHRS9
Dehydrogenase/reductase (SDR family) member 9 SEQ ID NOS: 4206-4213 DHRSX
Dehydrogenase/reductase (SDR family) X-linked SEQ ID NOS: 4214-4218 DHX29 DEAH (Asp-Glu-
Ala-His) box polypeptide 29 SEQ ID NOS: 4219-4221 DHX30 DEAH (Asp-Glu-Ala-His) box helicase 30
SEQ ID NOS: 4222-4229 DHX8 DEAH (Asp-Glu-Ala-His) box polypeptide 8 SEQ ID NOS: 4230-4234
DIO2 Deiodinase, iodothyronine, type II SEQ ID NOS: 4235-4244 DIXDC1 DIX domain containing 1
SEQ ID NOS: 4245-4248 DKK1 Dickkopf WNT signaling pathway inhibitor 1 SEQ ID NO: 4249 DKK2
Dickkopf WNT signaling pathway inhibitor 2 SEQ ID NOS: 4250-4252 DKK3 Dickkopf WNT signaling
pathway inhibitor 3 SEO ID NOS: 4253-4258 DKK4 Dickkopf WNT signaling pathway inhibitor 4 SEO
ID NO: 4259 DKKL1 Dickkopf-like 1 SEQ ID NOS: 4260-4265 DLG4 Discs, large homolog 4
(Drosophila) SEQ ID NOS: 4266-4274 DLK1 Delta-like 1 homolog (Drosophila) SEQ ID NOS: 4275-
4278 DLL1 Delta-like 1 (Drosophila) SEQ ID NOS: 4279-4280 DLL3 Delta-like 3 (Drosophila) SEQ ID
```

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NOS: 4281-4283 DMBT1 Deleted in malignant brain tumors 1 SEQ ID NOS: 4284-4290 DMKN
Dermokine SEQ ID NOS: 4291-4337 DMP1 Dentin matrix acidic phosphoprotein 1 SEQ ID NOS: 4338-
4339 DMRTA2 DMRT-like family A2 SEQ ID NOS: 4340-4341 DNAAF5 Dynein, axonemal, assembly
factor 5 SEQ ID NOS: 4342-4345 DNAH14 Dynein, axonemal, heavy chain 14 SEQ ID NOS: 4346-4360
DNAJB11 DnaJ (Hsp40) homolog, subfamily B, member 11 SEQ ID NOS: 4361-4362 DNAJB9 DnaJ
(Hsp40) homolog, subfamily B, member 9 SEQ ID NO: 4363 DNAJC25- DNAJC25-GNG10 readthrough
SEQ ID NO: 4364 GNG10 DNAJC3 DnaJ (Hsp40) homolog, subfamily C, member 3 SEQ ID NOS:
4365-4366 DNASE1 Deoxyribonuclease I SEQ ID NOS: 4367-4377 DNASE1L1 Deoxyribonuclease I-
like 1 SEQ ID NOS: 4378-4388 DNASE1L2 Deoxyribonuclease I-like 2 SEQ ID NOS: 4389-4394
DNASE1L3 Deoxyribonuclease I-like 3 SEQ ID NOS: 4395-4400 DNASE2 Deoxyribonuclease II,
lysosomal SEQ ID NOS: 4401-4402 DNASE2B Deoxyribonuclease II beta SEQ ID NOS: 4403-4404
DPEP1 Dipeptidase 1 (renal) SEQ ID NOS: 4405-4409 DPEP2 Dipeptidase 2 SEQ ID NOS: 4410-4416
DPEP3 Dipeptidase 3 SEQ ID NO: 4417 DPF3 D4, zinc and double PHD fingers, family 3 SEQ ID NOS:
4418-4424 DPP4 Dipeptidyl-peptidase 4 SEQ ID NOS: 4425-4429 DPP7 Dipeptidyl-peptidase 7 SEQ ID
NOS: 4430-4435 DPT Dermatopontin SEQ ID NO: 4436 DRAXIN Dorsal inhibitory axon guidance
protein SEQ ID NO: 4437 DSE Dermatan sulfate epimerase SEQ ID NOS: 4438-4446 DSG2 Desmoglein
2 SEQ ID NOS: 4447-4448 DSPP Dentin sialophosphoprotein SEQ ID NOS: 4449-4450 DST Dystonin
SEQ ID NOS: 4451-4469 DUOX1 Dual oxidase 1 SEQ ID NOS: 4470-4474 DYNLT3 Dynein, light
chain, Tctex-type 3 SEQ ID NOS: 4475-4477 E2F5 E2F transcription factor 5, p130-binding SEQ ID
NOS: 4478-4484 EBAG9 Estrogen receptor binding site associated, antigen, SEQ ID NOS: 4485-4493 9
EBI3 Epstein-Barr virus induced 3 SEQ ID NO: 4494 ECHDC1 Ethylmalonyl-CoA decarboxylase 1 SEQ
ID NOS: 4495-4513 ECM1 Extracellular matrix protein 1 SEQ ID NOS: 4514-4516 ECM2 Extracellular
matrix protein 2, female organ and SEQ ID NOS: 4517-4520 adipocyte specific ECSIT ECSIT signalling
integrator SEQ ID NOS: 4521-4532 EDDM3A Epididymal protein 3A SEQ ID NO: 4533 EDDM3B
Epididymal protein 3B SEQ ID NO: 4534 EDEM2 ER degradation enhancer, mamiosidase alpha-like
SEQ ID NOS: 4535-4536 2 EDEM3 ER degradation enhancer, mannosidase alpha-like SEQ ID NOS:
4537-4539 3 EDIL3 EGF-like repeats and discoidin I-like domains 3 SEQ ID NOS: 4540-4541 EDN1
Endothelin 1 SEQ ID NO: 4542 EDN2 Endothelin 2 SEQ ID NO: 4543 EDN3 Endothelin 3 SEQ ID
NOS: 4544-4549 EDNRB Endothelin receptor type B SEQ ID NOS: 4550-4558 EFEMP1 EGF containing
fibulin-like extracellular matrix SEQ ID NOS: 4559-4569 protein 1 EFEMP2 EGF containing fibulin-like
extracellular matrix SEQ ID NOS: 4570-4581 protein 2 EFNA1 Ephrin-A1 SEQ ID NOS: 4582-4583
EFNA2 Ephrin-A2 SEQ ID NO: 4584 EFNA4 Ephrin-A4 SEQ ID NOS: 4585-4587 EGFL6 EGF-like-
domain, multiple 6 SEQ ID NOS: 4588-4589 EGFL7 EGF-like-domain, multiple 7 SEQ ID NOS: 4590-
4594 EGFL8 EGF-like-domain, multiple 8 SEQ ID NOS: 4595-4597 EGFLAM EGF-like, fibronectin
type III and laminin G SEQ ID NOS: 4598-4606 domains EGFR Epidermal growth factor receptor SEQ
ID NOS: 4607-4614 EHBP1 EH domain binding protein 1 SEQ ID NOS: 4615-4626 EHF Ets
homologous factor SEQ ID NOS: 4627-4636 EHMT1 Euchromatic histone-lysine N-methyltransferase 1
SEQ ID NOS: 4637-4662 EHMT2 Euchromatic histone-lysine N-methyltransferase 2 SEQ ID NOS:
4663-4667 EIF2AK1 Eukaryotic translation initiation factor 2-alpha SEQ ID NOS: 4668-4671 kinase 1
ELANE Elastase, neutrophil expressed SEQ ID NOS: 4672-4673 ELN Elastin SEQ ID NOS: 4674-4696
ELP2 Elongator acetyltransferase complex subunit 2 SEQ ID NOS: 4697-4709 ELSPBP1 Epididymal
sperm binding protein 1 SEQ ID NOS: 4710-4715 EMC1 ER membrane protein complex subunit 1 SEQ
ID NOS: 4716-4722 EMC10 ER membrane protein complex subunit 10 SEQ ID NOS: 4723-4729 EMC9
ER membrane protein complex subunit 9 SEQ ID NOS: 4730-4733 EMCN Endomucin SEQ ID NOS:
4734-4738 EMID1 EMI domain containing 1 SEQ ID NOS: 4739-4745 EMILIN1 Elastin microfibril
interfacer 1 SEQ ID NOS: 4746-4747 EMILIN2 Elastin microfibril interfacer 2 SEQ ID NO: 4748
EMILIN3 Elastin microfibril interfacer 3 SEQ ID NO: 4749 ENAM Enamelin SEQ ID NO: 4750 ENDOG
Endonuclease G SEQ ID NO: 4751 ENDOU Endonuclease, polyU-specific SEQ ID NOS: 4752-4754
ENHO Energy homeostasis associated SEQ ID NO: 4755 ENO4 Enolase family member 4 SEQ ID NOS:
4756-4760 ENPP6 Ectonucleotide pyrophosphatase/phosphodiesterase SEQ ID NOS: 4761-4762 6
ENPP7 Ectonucleotide pyrophosphatase/phosphodiesterase SEQ ID NOS: 4763-4764 7 ENTPD5
Ectonucleoside triphosphate diphosphohydrolase 5 SEQ ID NOS: 4765-4769 ENTPD8 Ectonucleoside
triphosphate diphosphohydrolase 8 SEQ ID NOS: 4770-4773 EOGT EGF domain-specific O-linked N-
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SEQ ID NOS: 4774-4781 acetylglucosamine (GlcNAc) transferase EPCAM Epithelial cell adhesion
molecule SEQ ID NOS: 4782-4785 EPDR1 Ependymin related 1 SEQ ID NOS: 4786-4789 EPGN
Epithelial mitogen SEQ ID NOS: 4790-4798 EPHA10 EPH receptor A10 SEQ ID NOS: 4799-4806
EPHA3 EPH receptor A3 SEQ ID NOS: 4807-4809 EPHA4 EPH receptor A4 SEQ ID NOS: 4810-4819
EPHA7 EPH receptor A7 SEQ ID NOS: 4820-4821 EPHA8 EPH receptor A8 SEQ ID NOS: 4822-4823
EPHB2 EPH receptor B2 SEQ ID NOS: 4824-4828 EPHB4 EPH receptor B4 SEQ ID NOS: 4829-4831
EPHX3 Epoxide hydrolase 3 SEQ ID NOS: 4832-4835 EPO Erythropoietin SEQ ID NO: 4836 EPPIN
Epididymal peptidase inhibitor SEQ ID NOS: 4837-4839 EPPIN- EPPIN-WFDC6 readthrough SEQ ID
NO: 4840 WFDC6 EPS15 Epidermal growth factor receptor pathway SEQ ID NOS: 4841-4843 substrate
15 EPS8L1 EPS8-like 1 SEQ ID NOS: 4844-4849 EPX Eosinophil peroxidase SEQ ID NO: 4850 EPYC
Epiphycan SEQ ID NOS: 4851-4852 EQTN Equatorin, sperm acrosome associated SEQ ID NOS: 4853-
4855 ERAP1 Endoplasmic reticulum aminopeptidase 1 SEQ ID NOS: 4856-4861 ERAP2 Endoplasmic
reticulum aminopeptidase 2 SEQ ID NOS: 4862-4869 ERBB3 Erb-b2 receptor tyrosine kinase 3 SEQ ID
NOS: 4870-4883 FAM132B Family with sequence similarity 132, member B SEQ ID NOS: 4884-4886
ERLIN1 ER lipid raft associated 1 SEQ ID NOS: 4887-4889 ERLIN2 ER lipid raft associated 2 SEQ ID
NOS: 4890-4898 ERN1 Endoplasmic reticulum to nucleus signaling 1 SEQ ID NOS: 4899-4900 ERN2
Endoplasmic reticulum to nucleus signaling 2 SEQ ID NOS: 4901-4905 ERO1A Endoplasmic reticulum
oxidoreductase alpha SEQ ID NOS: 4906-4912 ERO1B Endoplasmic reticulum oxidoreductase beta SEQ
ID NOS: 4913-4915 ERP27 Endoplasmic reticulum protein 27 SEQ ID NOS: 4916-4917 ERP29
Endoplasmic reticulum protein 29 SEO ID NOS: 4918-4921 ERP44 Endoplasmic reticulum protein 44
SEQ ID NO: 4922 ERV3-1 Endogenous retrovirus group 3, member 1 SEQ ID NO: 4923 ESM1
Endothelial cell-specific molecule 1 SEQ ID NOS: 4924-4926 ESRP1 Epithelial splicing regulatory
protein 1 SEQ ID NOS: 4927-4935 EXOG Endo/exomiclease (5'-3'), endonuclease G-like SEQ ID NOS:
4936-4949 EXTL1 Exostosin-like glycosyltransferase 1 SEQ ID NO: 4950 EXTL2 Exostosin-like
glycosyltransferase 2 SEQ ID NOS: 4951-4955 F10 Coagulation factor X SEQ ID NOS: 4956-4959 F11
Coagulation factor XI SEQ ID NOS: 4960-4964 F12 Coagulation factor XII (Hageman factor) SEQ ID
NO: 4965 F13B Coagulation factor XIII, B polypeptide SEQ ID NO: 4966 F2 Coagulation factor II
(thrombin) SEQ ID NOS: 4967-4969 F2R Coagulation factor II (thrombin) receptor SEQ ID NOS: 4970-
4971 F2RL3 Coagulation factor II (thrombin) receptor-like 3 SEQ ID NOS: 4972-4973 F5 Coagulation
factor V (proaccelerin, labile factor) SEQ ID NOS: 4974-4975 F7 Coagulation factor VII (serum
prothrombin SEQ ID NOS: 4976-4979 conversion accelerator) F8 Coagulation factor VIII, procoagulant
component SEQ ID NOS: 4980-4985 F9 Coagulation factor IX SEQ ID NOS: 4986-4987 FABP6 Fatty
acid binding protein 6, ileal SEQ ID NOS: 4988-4990 FAM107B Family with sequence similarity 107,
member B SEQ ID NOS: 4991-5012 FAM131A Family with sequence similarity 131, member A SEQ ID
NOS: 5013-5021 FAM171A1 Family with sequence similarity 171, member A1 SEQ ID NOS: 5022-5023
FAM171B Family with sequence similarity 171, member B SEQ ID NOS: 5024-5025 FAM172A Family
with sequence similarity 172, member A SEQ ID NOS: 5026-5030 FAM177A1 Family with sequence
similarity 177, member A1 SEQ ID NOS: 5031-5040 FAM180A Family with sequence similarity 180,
member A SEQ ID NOS: 5041-5043 FAM189A1 Family with sequence similarity 189, member A1 SEQ
ID NOS: 5044-5045 FAM198A Family with sequence similarity 198, member A SEQ ID NOS: 5046-5048
FAM19A1 Family with sequence similarity 19 (chemokine (C- SEQ ID NOS: 5049-5051 C motif)-like),
member A1 FAM19A2 Family with sequence similarity 19 (chemokine (C- SEQ ID NOS: 5052-5059 C
motif)-like), member A2 FAM19A3 Family with sequence similarity 19 (chemokine (C- SEQ ID NOS:
5060-5061 C motif)-like), member A3 FAM19A4 Family with sequence similarity 19 (chemokine (C-
SEQ ID NOS: 5062-5064 C motif)-like), member A4 FAM19A5 Family with sequence similarity 19
(chemokine (C- SEQ ID NOS: 5065-5068 C motif)-like), member A5 FAM20A Family with sequence
similarity 20, member A SEQ ID NOS: 5069-5072 FAM20C Family with sequence similarity 20, member
C SEQ ID NO: 5073 FAM213A Family with sequence similarity 213, member A SEQ ID NOS: 5074-
5079 FAM46B Family with sequence similarity 46, member B SEQ ID NO: 5080 FAM57A Family with
sequence similarity 57, member A SEQ ID NOS: 5081-5086 FAM78A Family with sequence similarity 78,
member A SEQ ID NOS: 5087-5089 FAM96A Family with sequence similarity 96, member A SEQ ID
NOS: 5090-5094 FAM9B Family with sequence similarity 9, member B SEQ ID NOS: 5095-5098 FAP
Fibroblast activation protein, alpha SEQ ID NOS: 5099-5105 FAS Fas cell surface death receptor SEQ ID
```

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NOS: 5106-5115 FAT1 FAT atypical cadherin 1 SEQ ID NOS: 5116-5122 FBLN1 Fibulin 1 SEQ ID NOS:
5123-5135 FBLN2 Fibulin 2 SEQ ID NOS: 5136-5141 FBLN5 Fibulin 5 SEQ ID NOS: 5142-5147
FBLN7 Fibulin 7 SEQ ID NOS: 5148-5153 FBN1 Fibrillin 1 SEQ ID NOS: 5154-5157 FBN2 Fibrillin 2
SEQ ID NOS: 5158-5163 FBN3 Fibrillin 3 SEQ ID NOS: 5164-5168 FBXW7 F-box and WD repeat
domain containing 7, E3 SEQ ID NOS: 5169-5179 ubiquitin protein ligase FCAR Fc fragment of IgA
receptor SEQ ID NOS: 5180-5189 FCGBP Fc fragment of IgG binding protein SEQ ID NOS: 5190-5192
FCGR1B Fc fragment of IgG, high affinity Ib, receptor SEQ ID NOS: 5193-5198 (CD64) FCGR3A Fc
fragment of IgG, low affinity IIIa, receptor SEQ ID NOS: 5199-5205 (CD16a) FCGRT Fc fragment of
IgG, receptor, transporter, alpha SEQ ID NOS: 5206-5216 FCMR Fc fragment of IgM receptor SEQ ID
NOS: 5217-5223 FCN1 Ficolin (collagen/fibrinogen domain containing) 1 SEQ ID NOS: 5224-5225
FCN2 Ficolin (collagen/fibrinogen domain containing SEQ ID NOS: 5226-5227 lectin) 2 FCN3 Ficolin
(collagen/fibrinogen domain containing) 3 SEQ ID NOS: 5228-5229 FCRL1 Fc receptor-like 1 SEQ ID
NOS: 5230-5232 FCRL3 Fc receptor-like 3 SEQ ID NOS: 5233-5238 FCRL5 Fc receptor-like 5 SEQ ID
NOS: 5239-5241 FCRLA Fc receptor-like A SEQ ID NOS: 5242-5253 FCRLB Fc receptor-like B SEQ ID
NOS: 5254-5258 FDCSP Follicular dendritic cell secreted protein SEQ ID NO: 5259 FETUB Fetuin B
SEQ ID NOS: 5260-5266 FGA Fibrinogen alpha chain SEQ ID NOS: 5267-5269 FGB Fibrinogen beta
chain SEQ ID NOS: 5270-5272 FGF10 Fibroblast growth factor 10 SEQ ID NOS: 5273-5274 FGF17
Fibroblast growth factor 17 SEQ ID NOS: 5275-5276 FGF18 Fibroblast growth factor 18 SEQ ID NO:
5277 FGF19 Fibroblast growth factor 19 SEQ ID NO: 5278 FGF21 Fibroblast growth factor 21 SEQ ID
NOS: 5279-5280 FGF22 Fibroblast growth factor 22 SEO ID NOS: 5281-5282 FGF23 Fibroblast growth
factor 23 SEQ ID NO: 5283 FGF3 Fibroblast growth factor 3 SEQ ID NO: 5284 FGF4 Fibroblast growth
factor 4 SEQ ID NO: 5285 FGF5 Fibroblast growth factor 5 SEQ ID NOS: 5286-5288 FGF7 Fibroblast
growth factor 7 SEQ ID NOS: 5289-5293 FGF8 Fibroblast growth factor 8 (androgen-induced) SEQ ID
NOS: 5294-5299 FGFBP1 Fibroblast growth factor binding protein 1 SEQ ID NO: 5300 FGFBP2
Fibroblast growth factor binding protein 2 SEQ ID NO: 5301 FGFBP3 Fibroblast growth factor binding
protein 3 SEQ ID NO: 5302 FGFR1 Fibroblast growth factor receptor 1 SEQ ID NOS: 5303-5325 FGFR2
Fibroblast growth factor receptor 2 SEQ ID NOS: 5326-5347 FGFR3 Fibroblast growth factor receptor 3
SEQ ID NOS: 5348-5355 FGFR4 Fibroblast growth factor receptor 4 SEQ ID NOS: 5356-5365 FGFRL1
Fibroblast growth factor receptor-like 1 SEQ ID NOS: 5366-5371 FGG Fibrinogen gamma chain SEQ ID
NOS: 5372-5377 FGL1 Fibrinogen-like 1 SEQ ID NOS: 5378-5384 FGL2 Fibrinogen-like 2 SEQ ID
NOS: 5385-5386 FHL1 Four and a half LIM domains 1 SEQ ID NOS: 5387-5414 FHOD3 Formin
homology 2 domain containing 3 SEQ ID NOS: 5415-5421 FIBIN Fin bud initiation factor homolog
(zebrafish) SEQ ID NO: 5422 FICD FIC domain containing SEQ ID NOS: 5423-5426 FJX1 Four jointed
box 1 SEQ ID NO: 5427 FKBP10 FK506 binding protein 10, 65 kDa SEQ ID NOS: 5428-5433 FKBP11
FK506 binding protein 11, 19 kDa SEQ ID NOS: 5434-5440 FKBP14 FK506 binding protein 14, 22 kDa
SEQ ID NOS: 5441-5443 FKBP2 FK506 binding protein 2, 13 kDa SEQ ID NOS: 5444-5447 FKBP7
FK506 binding protein 7 SEQ ID NOS: 5448-5453 FKBP9 FK506 binding protem 9, 63 kDa SEQ ID
NOS: 5454-5457 FLT1 Fms-related tyrosine kinase 1 SEQ ID NOS: 5458-5466 FLT4 Fms-related
tyrosine kinase 4 SEQ ID NOS: 5467-5471 FMO1 Flavin containing monooxygenase 1 SEQ ID NOS:
5472-5476 FMO2 Flavin containing monooxygenase 2 (non- SEQ ID NOS: 5477-5479 functional) FMO3
Flavin containing monooxygenase 3 SEQ ID NOS: 5480-5482 FMO5 Flavin containing monooxygenase 5
SEQ ID NOS: 5483-5489 FMOD Fibromodulin SEQ ID NO: 5490 FN1 Fibronectin 1 SEQ ID NOS:
5491-5503 FNDC1 Fibronectin type III domain containing 1 SEQ ID NOS: 5504-5505 FNDC7
Fibronectin type III domain containing 7 SEQ ID NOS: 5506-5507 FOCAD Focadhesin SEQ ID NOS:
5508-5514 FOLR2 Folate receptor 2 (fetal) SEQ ID NOS: 5515-5524 FOLR3 Folate receptor 3 (gamma)
SEQ ID NOS: 5525-5529 FOXRED2 FAD-dependent oxidoreductase domain containing SEQ ID NOS:
5530-5533 2 FP325331.1 Uncharacterized protein UNQ6126/PRO20091 SEQ ID NO: 5534 CH507- SEQ
ID NOS: 5535-5541 9B2.3 FPGS Folylpolyglutamate synthase SEQ ID NOS: 5542-5548 FRAS1 Fraser
extracellular matrix complex subunit 1 SEQ ID NOS: 5549-5554 FREM1 FRAS1 related extracellular
matrix 1 SEO ID NOS: 5555-5559 FREM3 FRAS1 related extracellular matrix 3 SEO ID NO: 5560
FRMPD2 FERM and PDZ domain containing 2 SEQ ID NOS: 5561-5564 FRZB Frizzled-related protein
SEQ ID NO: 5565 FSHB Follicle stimulating hormone, beta polypeptide SEQ ID NOS: 5566-5568 FSHR
Follicle stimulating hormone receptor SEQ ID NOS: 5569-5572 FST Follistatin SEQ ID NOS: 5573-5576
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FSTL1 Follistatin-like 1 SEQ ID NOS: 5577-5580 FSTL3 Follistatin-like 3 (secreted glycoprotein) SEQ
ID NOS: 5581-5586 FSTL4 Follistatin-like 4 SEQ ID NOS: 5587-5589 FSTL5 Follistatin-like 5 SEQ ID
NOS: 5590-5592 FTCDNL1 Formiminotransferase cyclodeaminase N-terminal SEQ ID NOS: 5593-5596
like FUCA1 Fucosidase, alpha-L-1, tissue SEQ ID NO: 5597 FUCA2 Fucosidase, alpha-L-2, plasma SEQ
ID NOS: 5598-5599 FURIN Furin (paired basic amino acid cleaving enzyme) SEQ ID NOS: 5600-5606
FUT10 Fucosyltransferase 10 (alpha (1,3) SEQ ID NOS: 5607-5609 fucosyltransferase) FUT11
Fucosyltransferase 11 (alpha (1,3) SEQ ID NOS: 5610-5611 fucosyltransferase) FXN Frataxin SEQ ID
NOS: 5612-5619 FXR1 Fragile X mental retardation, autosomal homolog 1 SEQ ID NOS: 5620-5632
FXYD3 FXYD domain containing ion transport regulator 3 SEQ ID NOS: 5633-5645 GABBR1 Gamma-
aminobutyric acid (GABA) B receptor, 1 SEQ ID NOS: 5646-5657 GABRA1 Gamma-aminobutyric acid
(GABA) A receptor, SEQ ID NOS: 5658-5673 alpha 1 GABRA2 Gamma-aminobutyric acid (GABA) A
receptor, SEQ ID NOS: 5674-5688 alpha 2 GABRA5 Gamma-aminobutyric acid (GABA) A receptor,
SEQ ID NOS: 5689-5697 alpha 5 GABRG3 Gamma-aminobutyric acid (GABA) A receptor, SEQ ID
NOS: 5698-5703 gamma 3 GABRP Gamma-aminobutyric acid (GABA) A receptor, pi SEQ ID NOS:
5704-5712 GAL Galanin/GMAP prepropertide SEQ ID NO: 5713 GAL3ST1 Galactose-3-O-
sulfotransferase 1 SEQ ID NOS: 5714-5735 GAL3ST2 Galactose-3-O-sulfotransferase 2 SEQ ID NO:
5736 GAL3ST3 Galactose-3-O-sulfotransferase 3 SEQ ID NOS: 5737-5738 GALC Galactosylceramidase
SEQ ID NOS: 5739-5748 GALNS Galactosamine (N-acetyl)-6-sulfatase SEQ ID NOS: 5749-5754
GALNT10 Polypeptide N-acetylgalactosaminyltransferase 10 SEQ ID NOS: 5755-5758 GALNT12
Polypeptide N-acetylgalactosaminyltransferase 12 SEQ ID NOS: 5759-5760 GALNT15 Polypeptide N-
acetylgalactosaminyltransferase 15 SEQ ID NOS: 5761-5764 GALNT2 Polypeptide N-
acetylgalactosaminyltransferase 2 SEQ ID NO: 5765 GALNT6 Polypeptide N-
acetylgalactosaminyltransferase 6 SEQ ID NOS: 5766-5777 GALNT8 Polypeptide N-
acetylgalactosaminyltransferase 8 SEQ ID NOS: 5778-5781 GALNTL6 Polypeptide N-
acetylgalactosaminyltransferase- SEQ ID NOS: 5782-5785 like 6 GALP Galanin-like peptide SEQ ID
NOS: 5786-5788 GANAB Glucosidase, alpha; neutral AB SEQ ID NOS: 5789-5797 GARS Glycyl-tRNA
synthetase SEQ ID NOS: 5798-5801 GAS1 Growth arrest-specific 1 SEQ ID NO: 5802 GAS6 Growth
arrest-specific 6 SEQ ID NO: 5803 GAST Gastrin SEQ ID NO: 5804 PDDC1 Parkinson disease 7 domain
containing 1 SEQ ID NOS: 5805-5813 GBA Glucosidase, beta, acid SEQ ID NOS: 5814-5817 GBGT1
Globoside alpha-1,3-N- SEQ ID NOS: 5818-5826 acetylgalactosaminyltransferase 1 GC Group-specific
component (vitamin D binding SEQ ID NOS: 5827-5831 protein) GCG Glucagon SEQ ID NOS: 5832-
5833 GCGR Glucagon receptor SEQ ID NOS: 5834-5836 GCNT7 Glucosaminyl (N-acetyl) transferase
family SEQ ID NOS: 5837-5838 member 7 GCSH Glycine cleavage system protein H (aminomethyl SEQ
ID NOS: 5839-5847 carrier) GDF1 Growth differentiation factor 1 SEQ ID NO: 5848 GDF10 Growth
differentiation factor 10 SEQ ID NO: 5849 GDF11 Growth differentiation factor 11 SEQ ID NOS: 5850-
5851 GDF15 Growth differentiation factor 15 SEQ ID NOS: 5852-5854 GDF2 Growth differentiation
factor 2 SEQ ID NO: 5855 GDF3 Growth differentiation factor 3 SEQ ID NO: 5856 GDF5 Growth
differentiation factor 5 SEQ ID NOS: 5857-5858 GDF6 Growth differentiation factor 6 SEQ ID NOS:
5859-5861 GDF7 Growth differentiation factor 7 SEQ ID NO: 5862 GDF9 Growth differentiation factor 9
SEQ ID NOS: 5863-5867 GDNF Glial cell derived neurotrophic factor SEQ ID NOS: 5868-5875 GFOD2
Glucose-fructose oxidoreductase domain SEQ ID NOS: 5876-5881 containing 2 GFPT2 Glutamine-
fructose-6-phosphate transaminase 2 SEQ ID NOS: 5882-5884 GFRA2 GDNF family receptor alpha 2
SEQ ID NOS: 5885-5891 GFRA4 GDNF family receptor alpha 4 SEQ ID NOS: 5892-5894 GGA2 Golgi-
associated, gamma adaptin ear containing, SEQ ID NOS: 5895-5903 ARF binding protein 2 GGH
Gamma-glutamyl hydrolase (conjugase, SEQ ID NO: 5904 folylpolygammaglutamyl hydrolase) GGT1
Gamma-glutamyltransferase 1 SEQ ID NOS: 5905-5927 GGT5 Gamma-glutamyltransferase 5 SEQ ID
NOS: 5928-5932 GH1 Growth hormone 1 SEQ ID NOS: 5933-5937 GH2 Growth hormone 2 SEQ ID
NOS: 5938-5942 GHDC GH3 domain containing SEQ ID NOS: 5943-5950 GHRH Growth hormone
releasing hormone SEQ ID NOS: 5951-5953 GHRHR Growth hormone releasing hormone receptor SEQ
ID NOS: 5954-5959 GHRL Ghrelin/obestatin prepropeptide SEQ ID NOS: 5960-5970 GIF Gastric
intrinsic factor (vitamin B synthesis) SEQ ID NOS: 5971-5972 GIP Gastric inhibitory polypeptide SEQ ID
NO: 5973 GKN1 Gastrokine 1 SEQ ID NO: 5974 GKN2 Gastrokine 2 SEQ ID NOS: 5975-5976 GLA
Galactosidase, alpha SEQ ID NOS: 5977-5978 GLB1 Galactosidase, beta 1 SEQ ID NOS: 5979-5987
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GLB1L Galactosidase, beta 1-like SEQ ID NOS: 5988-5995 GLB1L2 Galactosidase, beta 1-like 2 SEQ ID
NOS: 5996-5997 GLCE Glucuronic acid epimerase SEQ ID NOS: 5998-5999 GLG1 Golgi glycoprotein 1
SEQ ID NOS: 6000-6007 GLIPR1 GLI pathogenesis-related 1 SEQ ID NOS: 6008-6011 GLIPR1L1 GLI
pathogenesis-related 1 like 1 SEQ ID NOS: 6012-6015 GLIS3 GLIS family zinc finger 3 SEQ ID NOS:
6016-6024 GLMP Glycosylated lysosomal membrane protein SEQ ID NOS: 6025-6033 GLRB Glycine
receptor, beta SEQ ID NOS: 6034-6039 GLS Glutaminase SEQ ID NOS: 6040-6047 GLT6D1
Glycosyltransferase 6 domain containing 1 SEQ ID NOS: 6048-6049 GLTPD2 Glycolipid transfer protein
domain containing 2 SEQ ID NO: 6050 GLUD1 Glutamate dehydrogenase 1 SEQ ID NO: 6051 GM2A
GM2 ganglioside activator SEQ ID NOS: 6052-6054 GML Glycosylphosphatidylinositol anchored
molecule SEQ ID NOS: 6055-6056 like GNAS GNAS complex locus SEQ ID NOS: 6057-6078 GNLY
Granulysin SEQ ID NOS: 6079-6082 GNPTG N-acetylglucosamine-1-phosphate transferase, SEQ ID
NOS: 6083-6087 gamma subunit GNRH1 Gonadotropin-releasing hormone 1 (luteinizing- SEQ ID NOS:
6088-6089 releasing hormone) GNRH2 Gonadotropin-releasing hormone 2 SEQ ID NOS: 6090-6093
GNS Glucosamine (N-acetyl)-6-sulfatase SEQ ID NOS: 6094-6099 GOLM1 Golgi membrane protein 1
SEQ ID NOS: 6100-6104 GORAB Golgin, RAB6-interacting SEQ ID NOS: 6105-6107 GOT2 Glutamic-
oxaloacetic transaminase 2, SEQ ID NOS: 6108-6110 mitochondrial GP2 Glycoprotein 2 (zymogen
granule membrane) SEQ ID NOS: 6111-6119 GP6 Glycoprotein VI (platelet) SEQ ID NOS: 6120-6123
GPC2 Glypican 2 SEQ ID NOS: 6124-6125 GPC5 Glypican 5 SEQ ID NOS: 6126-6128 GPC6 Glypican
6 SEQ ID NOS: 6129-6130 GPD2 Glycerol-3-phosphate dehydrogenase 2 SEQ ID NOS: 6131-6139
(mitochondrial) GPER1 G protein-coupled estrogen receptor 1 SEQ ID NOS: 6140-6146 GPHA2
Glycoprotein hormone alpha 2 SEQ ID NOS: 6147-6149 GPHB5 Glycoprotein hormone beta 5 SEQ ID
NOS: 6150-6151 GPIHBP1 Glycosylphosphatidylinositol anchored high SEQ ID NO: 6152 density
lipoprotein binding protein 1 GPLD1 Glycosylphosphatidylinositol specific SEQ ID NO: 6153
phospholipase D1 GPNMB Glycoprotein (transmembrane) nmb SEQ ID NOS: 6154-6156 GPR162 G
protein-coupled receptor 162 SEQ ID NOS: 6157-6160 GPX3 Glutathione peroxidase 3 SEQ ID NOS:
6161-6168 GPX4 Glutathione peroxidase 4 SEQ ID NOS: 6169-6179 GPX5 Glutathione peroxidase 5
SEQ ID NOS: 6180-6181 GPX6 Glutathione peroxidase 6 SEQ ID NOS: 6182-6184 GPX7 Glutathione
peroxidase 7 SEQ ID NO: 6185 GREM1 Gremlin 1, DAN family BMP antagonist SEQ ID NOS: 6186-
6188 GREM2 Gremlin 2, DAN family BMP antagonist SEQ ID NO: 6189 GRHL3 Grainyhead-like
transcription factor 3 SEQ ID NOS: 6190-6195 GRIA2 Glutamate receptor, ionotropic, AMPA 2 SEQ ID
NOS: 6196-6207 GRIA3 Glutamate receptor, ionotropic, AMPA 3 SEQ ID NOS: 6208-6213 GRIA4
Glutamate receptor, ionotropic, AMPA 4 SEQ ID NOS: 6214-6225 GRIK2 Glutamate receptor, ionotropic,
kainate 2 SEQ ID NOS: 6226-6234 GRIN2B Glutamate receptor, ionotropic, N-methyl D- SEQ ID NOS:
6235-6238 aspartate 2B GRM2 Glutamate receptor, metabotropic 2 SEQ ID NOS: 6239-6242 GRM3
Glutamate receptor, metabotropic 3 SEQ ID NOS: 6243-6247 GRM5 Glutamate receptor, metabotropic 5
SEQ ID NOS: 6248-6252 GRN Granulin SEQ ID NOS: 6253-6268 GRP Gastrin-releasing peptide SEQ
ID NOS: 6269-6273 DFNA5 Deafness, autosomal dominant 5 SEQ ID NOS: 6274-6282 GSG1 Germ cell
associated 1 SEQ ID NOS: 6283-6291 GSN Gelsolin SEQ ID NOS: 6292-6300 GTDC1
Glycosyltransferase-like domain containing 1 SEQ ID NOS: 6301-6314 GTPBP10 GTP-binding protein
10 (putative) SEQ ID NOS: 6315-6323 GUCA2A Guanylatc cyclase activator 2A (guanylin) SEQ ID NO:
6324 GUCA2B Guanylate cyclase activator 2B (uroguanylin) SEQ ID NO: 6325 GUSB Glucuronidase,
beta SEQ ID NOS: 6326-6330 GVQW1 GVQW motif containing 1 SEQ ID NO: 6331 GXYLT1
Glucoside xylosyltransferase 1 SEQ ID NOS: 6332-6333 GXYLT2 Glucoside xylosyltransferase 2 SEQ
ID NOS: 6334-6336 GYPB Glycophorin B (MNS blood group) SEQ ID NOS: 6337-6345 GZMA
Granzyme A (granzyme 1, cytotoxic T- SEQ ID NO: 6346 lymphocyte-associated serine esterase 3)
GZMB Granzyme B (granzyme 2, cytotoxic T- SEQ ID NOS: 6347-6355 lymphocyte-associated serine
esterase 1) GZMH Granzyme H (cathepsin G-like 2, protein h-CCPX) SEQ ID NOS: 6356-6358 GZMK
Granzyme K (granzyme 3; tryptase II) SEQ ID NO: 6359 GZMM Granzyme M (lymphocyte met-ase 1)
SEQ ID NOS: 6360-6361 H6PD Hexose-6-phosphate dehydrogenase (glucose 1- SEQ ID NOS: 6362-
6363 dehydrogenase) HABP2 Hyaluronan binding protein 2 SEQ ID NOS: 6364-6365 HADHB
Hydroxyacyl-CoA dehydrogenase/3-ketoacyl-CoA SEQ ID NOS: 6366-6372 thiolase/enoyl-CoA
hydratase (trifunctional protein), beta subunit HAMP Hepcidin antimicrobial peptide SEQ ID NOS: 6373-
6374 HAPLN1 Hyaluronan and proteoglycan link protein 1 SEQ ID NOS: 6375-6381 HAPLN2
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Hyaluronan and proteoglycan link protein 2 SEQ ID NOS: 6382-6383 HAPLN3 Hyaluronan and
proteoglycan link protein 3 SEQ ID NOS: 6384-6387 HAPLN4 Hyaluronan and proteoglycan link protein
4 SEQ ID NO: 6388 HARS2 Histidyl-tRNA synthetase 2, mitochondrial SEQ ID NOS: 6389-6404
HAVCR1 Hepatitis A virus cellular receptor 1 SEQ ID NOS: 6405-6409 HCCS Holocytochrome c
synthase SEQ ID NOS: 6410-6412 HCRT Hypocretin (orexin) neuropeptide precursor SEQ ID NO: 6413
CECR5 Cat eye syndrome chromosome region, candidate 5 SEQ ID NOS: 6414-6416 HEATR5A HEAT
repeat containing 5A SEQ ID NOS: 6417-6423 HEPH Hephaestin SEQ ID NOS: 6424-6431 HEXA
Hexosaminidase A (alpha polypeptide) SEQ ID NOS: 6432-6441 HEXB Hexosaminidase B (beta
polypeptide) SEQ ID NOS: 6442-6447 HFE2 Hemochromatosis type 2 (juvenile) SEQ ID NOS: 6448-
6454 HGF Hepatocyte growth factor (hepapoietin A; scatter SEQ ID NOS: 6455-6465 factor) HGFAC
HGF activator SEQ ID NOS: 6466-6467 HHIP Hedgehog interacting protein SEQ ID NOS: 6468-6469
HHIPL1 HHIP-like 1 SEQ ID NOS: 6470-6471 HHIPL2 HHIP-like 2 SEQ ID NO: 6472 HHLA1 HERV-
H LTR-associating 1 SEQ ID NOS: 6473-6474 HHLA2 HERV-H LTR-associating 2 SEQ ID NOS: 6475-
6485 HIBADH 3-hydroxyisobutyrate dehydrogenase SEQ ID NOS: 6486-6488 HINT2 Histidine triad
nucleotide binding protein 2 SEQ ID NO: 6489 HLA-A Major histocompatibility complex, class I, A SEQ
ID NOS: 6490-6494 HLA-C Major histocompatibility complex, class I, C SEQ ID NOS: 6495-6499 HLA-
Major histocompatibility complex, class II, DO SEQ ID NOS: 6500-6501 DOA alpha HLA- Major
histocompatibility complex, class II, DP SEQ ID NOS: 6502-6505 DPA1 alpha 1 HLA- Major
histocompatibility complex, class II, DQ SEQ ID NOS: 6506-6511 DQA1 alpha 1 HLA- Major
histocompatibility complex, class II, DQ SEQ ID NOS: 6512-6517 DQB1 beta 1 HLA- Major
histocompatibility complex, class II, DQ SEQ ID NOS: 6518-6521 DQB2 beta 2 HMCN1 Hemicentin 1
SEQ ID NOS: 6522-6523 HMCN2 Hemicentin 2 SEQ ID NOS: 6524-6527 HMGCL 3-hydroxymethyl-3-
methylglutaryl-CoA lyase SEQ ID NOS: 6528-6531 HMSD Histocompatibility (minor) serpin domain
SEQ ID NOS: 6532-6533 containing HP Haptoglobin SEQ ID NOS: 6534-6547 HPR Haptoglobin-related
protein SEQ ID NOS: 6548-6550 HPSE Heparanase SEQ ID NOS: 6551-6557 HPSE2 Heparanase 2
(inactive) SEQ ID NOS: 6558-6563 HPX Hemopexin SEQ ID NOS: 6564-6565 HRC Histidine rich
calcium binding protein SEQ ID NOS: 6566-6568 HRG Histidine-rich glycoprotein SEQ ID NO: 6569
HS2ST1 Heparan sulfate 2-O-sulfotransferase 1 SEQ ID NOS: 6570-6572 HS3ST1 Heparan sulfate
(glucosamine) 3-O- SEQ ID NOS: 6573-6575 sulfotransferase 1 HS6ST1 Heparan sulfate 6-O-
sulfotransferase 1 SEQ ID NO: 6576 HS6ST3 Heparan sulfate 6-O-sulfotransferase 3 SEQ ID NOS: 6577-
6578 HSD11B1L Hydroxysteroid (11-beta) dehydrogenase 1-like SEQ ID NOS: 6579-6597 HSD17B11
Hydroxysteroid (17-beta) dehydrogenase 11 SEQ ID NOS: 6598-6599 HSD17B7 Hydroxysteroid (17-
beta) dehydrogenase 7 SEQ ID NOS: 6600-6604 HSP90B1 Heat shock protein 90 kDa beta (Grp94),
member 1 SEQ ID NOS: 6605-6610 HSPAI3 Heat shock protein 70 kDa family, member 13 SEQ ID NO:
6611 HSPA5 Heat shock 70 kDa protein 5 (glucose-regulated SEQ ID NO: 6612 protein, 78 kDa) HSPG2
Heparan sulfate proteoglycan 2 SEQ ID NOS: 6613-6617 HTATIP2 HIV-1 Tat interactive protein 2, 30
kDa SEQ ID NOS: 6618-6625 HTN1 Histatin 1 SEQ ID NOS: 6626-6628 HTN3 Histatin 3 SEQ ID NOS:
6629-6631 HTRA1 HtrA serine peptidase 1 SEQ ID NOS: 6632-6633 HTRA3 HtrA serine peptidase 3
SEQ ID NOS: 6634-6635 HTRA4 HtrA serine peptidase 4 SEQ ID NO: 6636 HYAL1
Hyaluronoglucosaminidase 1 SEQ ID NOS: 6637-6645 IIYAL2 Hyaluronoglucosaminidase 2 SEQ ID
NOS: 6646-6654 HYAL3 Hyaluronoglucosaminidase 3 SEQ ID NOS: 6655-6661 HYOU1 Hypoxia up-
regulated 1 SEQ ID NOS: 6662-6676 IAPP Islet amyloid polypeptide SEQ ID NOS: 6677-6681 IBSP
Integrin-binding sialoprotein SEQ ID NO: 6682 ICAM1 Intercellular adhesion molecule 1 SEQ ID NOS:
6683-6685 ICAM2 Intercellular adhesion molecule 2 SEQ ID NOS: 6686-6696 ICAM4 Intercellular
adhesion molecule 4 (Landsteiner- SEQ ID NOS: 6697-6699 Wiener blood group) ID1 Inhibitor of DNA
binding 1, dominant negative SEQ ID NOS: 6700-6701 helix-loop-helix protein IDE Insulin-degrading
enzyme SEQ ID NOS: 6702-6705 IDNK IdnK, gluconokinase homolog (E. coli) SEQ ID NOS: 6706-
6711 IDS Iduronate 2-sulfatase SEQ ID NOS: 6712-6717 IDUA Iduronidase, alpha-L- SEQ ID NOS:
6718-6723 IFI27L2 Interferon, alpha-inducible protein 27-like 2 SEQ ID NOS: 6724-6725 IFI30
Interferon, gamma-inducible protein 30 SEQ ID NOS: 6726-6727 IFNA1 Interferon, alpha 1 SEQ ID NO:
6728 IFNA10 Interferon, alpha 10 SEQ ID NO: 6729 IFNA13 Interferon, alpha 13 SEQ ID NOS: 6730-
6731 IFNA14 Interferon, alpha 14 SEQ ID NO: 6732 IFNA16 Interferon, alpha 16 SEQ ID NO: 6733
IFNA17 Interferon, alpha 17 SEQ ID NO: 6734 IFNA2 Interferon, alpha 2 SEQ ID NO: 6735 IFNA21
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Interferon, alpha 21 SEQ ID NO: 6736 IFNA4 Interferon, alpha 4 SEQ ID NO: 6737 IFNA5 Interferon,
alpha 5 SEQ ID NO: 6738 IFNA6 Interferon, alpha 6 SEQ ID NOS: 6739-6740 IFNA7 Interferon, alpha 7
SEQ ID NO: 6741 IFNA8 Interferon, alpha 8 SEQ ID NO: 6742 IFNAR1 Interferon (alpha, beta and
omega) receptor 1 SEQ ID NOS: 6743-6744 IFNB1 Interferon, beta 1, fibroblast SEQ ID NO: 6745 IFNE
Interferon, epsilon SEQ ID NO: 6746 IFXG Interferon, gamma SEQ ID NO: 6747 IFNGR1 Interferon
gamma receptor 1 SEQ ID NOS: 6748-6758 IFNL1 Interferon, lambda 1 SEQ ID NO: 6759 IFNL2
Interferon, lambda 2 SEQ ID NO: 6760 IFNL3 Interferon, lambda 3 SEQ ID NOS: 6761-6762 IFNLR1
Interferon, lambda receptor 1 SEQ ID NOS: 6763-6767 IFNW1 Interferon, omega 1 SEQ ID NO: 6768
IGF1 Insulin-like growth factor I (somatomedin C) SEQ ID NOS: 6769-6774 IGF2 Insulin-like growth
factor 2 SEQ ID NOS: 6775-6782 IGFALS Insulin-like growth factor binding protein, acid SEQ ID NOS:
6783-6785 labile subunit IGFBP1 Insulin-like growth factor binding protein 1 SEQ ID NOS: 6786-6788
IGFBP2 Insulin-like growth factor binding protein 2, 36 kDa SEQ ID NOS: 6789-6792 IGFBP3 Insulin-
like growth factor binding protein 3 SEQ ID NOS: 6793-6800 IGFBP4 Insulin-like growth factor binding
protein 4 SEQ ID NO: 6801 IGFBP5 Insulin-like growth factor binding protein 5 SEQ ID NOS: 6802-
6803 IGFBP6 Insulin-like growth factor binding protein 6 SEQ ID NOS: 6804-6806 IGFBP7 Insulin-like
growth factor binding protein 7 SEQ ID NOS: 6807-6808 IGFBPL1 Insulin-like growth factor binding
protein-like 1 SEQ ID NO: 6809 IGFL1 IGF-like family member 1 SEQ ID NO: 6810 IGFL2 IGF-like
family member 2 SEQ ID NOS: 6811-6813 IGFL3 IGF-like family member 3 SEQ ID NO: 6814 IGFLR1
IGF-like family receptor 1 SEQ ID NOS: 6815-6823 IGIP IgA-inducing protein SEQ ID NO: 6824
IGLON5 IgLON family member 5 SEQ ID NO: 6825 IGSF1 Immunoglobulin superfamily, member 1
SEQ ID NOS: 6826-6831 IGSF10 Immunoglobulin superfamily, member 10 SEQ ID NOS: 6832-6833
IGSF11 Immunoglobulin superfamily, member 11 SEQ ID NOS: 6834-6841 IGSF21 Immunoglobulin
superfamily, member 21 SEQ ID NO: 6842 IGSF8 Immunoglobulin superfamily, member 8 SEQ ID NOS:
6843-6846 IGSF9 Immunoglobulin superfamily, member 9 SEQ ID NOS: 6847-6849 IHH Indian
hedgehog SEQ ID NO: 6850 IL10 Interleukin 10 SEQ ID NOS: 6851-6852 IL11 Interleukin 11 SEQ ID
NOS: 6853-6856 IL11RA Interleukin 11 receptor, alpha SEQ ID NOS: 6857-6867 IL12B Interleukin 12B
SEQ ID NO: 6868 IL12RB1 Interleukin 12 receptor, beta 1 SEQ ID NOS: 6869-6874 IL12RB2
Interleukin 12 receptor, beta 2 SEQ ID NOS: 6875-6879 IL13 Interleukin 13 SEQ ID NOS: 6880-6881
IL13RA1 Interleukin 13 receptor, alpha 1 SEQ ID NOS: 6882-6883 IL15RA Interleukin 15 receptor, alpha
SEQ ID NOS: 6884-6901 IL17A Interleukin 17A SEQ ID NO: 6902 IL17B Interleukin 17B SEQ ID NO:
6903 IL17C Interleukin 17C SEQ ID NO: 6904 IL17D Interleukin 17D SEQ ID NOS: 6905-6907 IL17F
Interleukin 17F SEQ ID NO: 6908 IL17RA Interleukin 17 receptor A SEQ ID NOS: 6909-6910 IL17RC
Interleukin 17 receptor C SEQ ID NOS: 6911-6926 IL17RE Interleukin 17 receptor E SEQ ID NOS:
6927-6933 1L18BP Interleukin 18 binding protein SEQ ID NOS: 6934-6944 IL18R1 Interleukin 18
receptor 1 SEQ ID NOS: 6945-6948 IL18RAP Interleukin 18 receptor accessory protein SEQ ID NOS:
6949-6951 IL19 Interleukin 19 SEQ ID NOS: 6952-6954 IL1R1 Interleukin 1 receptor, type I SEQ ID
NOS: 6955-6967 IL1R2 Interleukin 1 receptor, type II SEQ ID NOS: 6968-6971 IL1RAP Interleukin 1
receptor accessory protein SEQ ID NOS: 6972-6985 IL1RL1 Interleukin 1 receptor-like 1 SEQ ID NOS:
6986-6991 IL1RL2 Interleukin 1 receptor-like 2 SEQ ID NOS: 6992-6994 IL1RN Interleukin 1 receptor
antagonist SEQ ID NOS: 6995-6999 IL2 Interleukin 2 SEQ ID NO: 7000 IL20 Interleukin 20 SEQ ID
NOS: 7001-7003 IL20RA Interleukin 20 receptor, alpha SEQ ID NOS: 7004-7010 IL21 Interleukin 21
SEQ ID NOS: 7011-7012 IL22 Interleukin 22 SEQ ID NOS: 7013-7014 IL22RA2 Interleukin 22 receptor,
alpha 2 SEQ ID NOS: 7015-7017 IL23A Interleukin 23, alpha subunit p19 SEQ ID NO: 7018 IL24
Interleukin 24 SEQ ID NOS: 7019-7024 IL25 Interleukin 25 SEQ ID NOS: 7025-7026 1L26 Interleukin
26 SEQ ID NO: 7027 IL27 Interleukin 27 SEQ ID NOS: 7028-7029 IL2RB Interleukin 2 receptor, beta
SEQ ID NOS: 7030-7034 IL3 Interleukin 3 SEQ ID NO: 7035 IL31 Interleukin 31 SEQ ID NO: 7036
IL31RA Interleukin 31 receptor A SEQ ID NOS: 7037-7044 IL32 Interleukin 32 SEQ ID NOS: 7045-7074
IL34 Interleukin 34 SEQ ID NOS: 7075-7078 IL3RA Interleukin 3 receptor, alpha (low affinity) SEQ ID
NOS: 7079-7081 IL4 Interleukin 4 SEQ ID NOS: 7082-7084 IL4I1 Interleukin 4 induced 1 SEQ ID NOS:
7085-7092 IL4R Interleukin 4 receptor SEQ ID NOS: 7093-7106 IL5 Interleukin 5 SEQ ID NOS: 7107-
7108 IL5RA Interleukin 5 receptor, alpha SEQ ID NOS: 7109-7118 IL6 Interleukin 6 SEQ ID NOS: 7119-
7125 IL6R Interleukin 6 receptor SEQ ID NOS: 7126-7131 IL6ST Interleukin 6 signal transducer SEQ ID
NOS: 7132-7141 IL7 Interleukin 7 SEQ ID NOS: 7142-7149 IL7R Interleukin 7 receptor SEQ ID NOS:
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7150-7156 IL9 Interleukin 9 SEQ ID NO: 7157 ILDR1 Immunoglobulin-like domain containing receptor
1 SEQ ID NOS: 7158-7162 ILDR2 Immunoglobulin-like domain containing receptor 2 SEQ ID NOS:
7163-7169 IMP4 IMP4, U3 small nucleolar ribonucleoprotein SEQ ID NOS: 7170-7175 IMPG1
Interphotoreceptor matrix proteoglycan 1 SEQ ID NOS: 7176-7179 INHA Inhibin, alpha SEQ ID NO:
7180 INHBA Inhibin, beta A SEQ ID NOS: 7181-7183 INHBB Inhibin, beta B SEQ ID NO: 7184 INHBC
Inhibin, beta C SEQ ID NO: 7185 INHBE Inhibin, beta E SEQ ID NOS: 7186-7187 INPP5A Inositol
polyphosphate-5-phosphatase A SEQ ID NOS: 7188-7192 INS Insulin SEQ ID NOS: 7193-7197 INS-
INS-IGF2 readthrough SEQ ID NOS: 7198-7199 IGF2 INSL3 Insulin-like 3 (Leydig cell) SEQ ID NOS:
7200-7202 INSL4 Insulin-like 4 (placenta) SEQ ID NO: 7203 INSL5 Insulin-like 5 SEQ ID NO: 7204
INSL6 Insulin-like 6 SEQ ID NO: 7205 INTS3 Integrator complex subunit 3 SEQ ID NOS: 7206-7211
IPO11 Importin 11 SEQ ID NOS: 7212-7220 IPO9 Importin 9 SEQ ID NOS: 7221-7222 IQCF6 IQ motif
containing F6 SEQ ID NOS: 7223-7224 IRAK3 Interleukin-1 receptor-associated kinase 3 SEQ ID NOS:
7225-7227 IRS4 Insulin receptor substrate 4 SEQ ID NO: 7228 ISLR Immunoglobulin superfamily
containing leucine- SEQ ID NOS: 7229-7232 rich repeat ISLR2 Immunoglobulin superfamily containing
leucine- SEQ ID NOS: 7233-7242 rich repeat 2 ISM1 Isthmin 1, angiogenesis inhibitor SEQ ID NO: 7243
ISM2 Isthmin 2 SEQ ID NOS: 7244-7249 ITGA4 Integrin, alpha 4 (antigen CD49D, alpha 4 subunit SEQ
ID NOS: 7250-7252 of VLA-4 receptor) ITGA9 Integrin, alpha 9 SEQ ID NOS: 7253-7255 ITGAL
Integrity alpha L (antigen CD11A (p180), SEQ ID NOS: 7256-7265 lymphocyte function-associated
antigen 1; alpha polypeptide) ITGAX Integrin, alpha X (complement component 3 SEQ ID NOS: 7266-
7268 receptor 4 subunit) ITGB1 Integrin, beta 1 (fibronectin receptor, beta SEQ ID NOS: 7269-7284
polypeptide, antigen CD29 includes MDF2, MSK12) ITGB2 Integrin, beta 2 (complement component 3
receptor SEQ ID NOS: 7285-7301 3 and 4 subunit) ITGB3 Integrin, beta 3 (platelet glycoprotein IIIa,
antigen SEQ ID NOS: 7302-7304 CD61) ITGB7 Integrin, beta 7 SEQ ID NOS: 7305-7312 ITGBL1
Integrin, beta-like 1 (with EGF-like repeat SEQ ID NOS: 7313-7318 domains) ITIH1 Inter-alpha-trypsin
inhibitor heavy chain 1 SEQ ID NOS: 7319-7324 ITIH2 Inter-alpha-trypsin inhibitor heavy chain 2 SEQ
ID NOS: 7325-7327 ITIH3 Inter-alpha-trypsin inhibitor heavy chain 3 SEQ ID NOS: 7328-7330 ITIH4
Inter-alpha-trypsin inhibitor heavy chain family, SEQ ID NOS: 7331-7334 member 4 ITIH5 Inter-alpha-
trypsin inhibitor heavy chain family, SEQ ID NOS: 7335-7338 member 5 ITIH6 Inter-alpha-trypsin
inhibitor heavy chain family, SEQ ID NO: 7339 member 6 ITLN1 Intelectin 1 (galactofuranose binding)
SEQ ID NO: 7340 ITLN2 Intelectin 2 SEQ ID NO: 7341 IZUMO1R IZUMO1 receptor, JUNO SEQ ID
NOS: 7342-7343 IZUMO4 IZUMO family member 4 SEQ ID NOS: 7344-7350 AMICA1 Adhesion
molecule, interacts with CXADR antigen SEQ ID NOS: 7351-7359 1 JCHAIN Joining chain of
multimeric IgA and IgM SEQ ID NOS: 7360-7365 JMJD8 Jumonji domain containing 8 SEQ ID NOS:
7366-7370 JSRP1 Junctional sarcoplasmic reticulum protein 1 SEQ ID NO: 7371 KANSL2 KAT8
regulatory NSL complex subunit 2 SEQ ID NOS: 7372-7382 KAZALD1 Kazal-type serine peptidase
inhibitor domain 1 SEQ ID NO: 7383 KCNIP3 Kv channel interacting protein 3, calsenilin SEQ ID NOS:
7384-7386 KCNK7 Potassium channel, two pore domain subfamily K, SEQ ID NOS: 7387-7392 member
7 KCNN4 Potassium channel, calcium activated SEQ ID NOS: 7393-7398 intermediate/small
conductance subfamily N alpha, member 4 KCNU1 Potassium channel, subfamily U, member 1 SEQ ID
NOS: 7399-7403 KCP Kielin/chordin-like protein SEQ ID NOS: 7404-7407 KDELC1 KDEL (Lys-Asp-
Glu-Leu) containing 1 SEQ ID NO: 7408 KDELC2 KDEL (Lys-Asp-Glu-Leu) containing 2 SEQ ID NOS:
7409-7412 KDM1A Lysine (K)-specific demethylase 1A SEQ ID NOS: 7413-7416 KDM3B Lysine (K)-
specific demethylase 3B SEQ ID NOS: 7417-7420 KDM6A Lysine (K)-specific demethylase 6A SEQ ID
NOS: 7421-7430 KDM7A Lysine (K)-specific demethylase 7A SEQ ID NOS: 7431-7432 KDSR 3-
ketodihydrosphingosine reductase SEQ ID NOS: 7433-7439 KERA Keratocan SEQ ID NO: 7440
KIAA0100 KIAA0100 SEQ ID NOS: 7441-7446 KIAA0319 KIAA0319 SEQ ID NOS: 7447-7452
KIAA1324 KIAA1324 SEQ ID NOS: 7453-7461 KIFC2 Kinesin family member C2 SEQ ID NOS: 7462-
7464 KIR2DL4 Killer cell immunoglobulin-like receptor, two SEQ ID NOS: 7465-7471 domains, long
cytoplasmic tail, 4 KIR3DX1 Killer cell immunoglobulin-like receptor, three SEQ ID NOS: 7472-7476
domains, X1 KIRREL2 Kin of IRRE like 2 (Drosophila) SEQ ID NOS: 7477-7481 KISS1 KiSS-1
metastasis-suppressor SEQ ID NOS: 7482-7483 KLHL11 Kelch-like family member 11 SEQ ID NO:
7484 KLHL22 Kelch-like family member 22 SEQ ID NOS: 7485-7491 KLK1 Kallikrein 1 SEQ ID NOS:
7492-7493 KLK10 Kallikrein-related peptidase 10 SEQ ID NOS: 7494-7498 KLK11 Kallikrein-related
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peptidase 11 SEQ ID NOS: 7499-7507 KLK12 Kallikrein-related peptidase 12 SEQ ID NOS: 7508-7514
KLK13 Kallikrein-related peptidase 13 SEQ ID NOS: 7515-7523 KLK14 Kallikrein-related peptidase 14
SEQ ID NOS: 7524-7525 KLK15 Kallikrein-related peptidase 15 SEQ ID NOS: 7526-7530 KLK2
Kallikrein-related peptidase 2 SEQ ID NOS: 7531-7543 KLK3 Kallikrein-related peptidase 3 SEQ ID
NOS: 7544-7555 KLK4 Kallikrein-related peptidase 4 SEQ ID NOS: 7556-7560 KLK5 Kallikrein-related
peptidase 5 SEQ ID NOS: 7561-7564 KLK6 Kallikrein-related peptidase 6 SEQ ID NOS: 7565-7571
KLK7 Kallikrein-related peptidase 7 SEQ ID NOS: 7572-7576 KLK8 Kallikrein-related peptidase 8 SEQ
ID NOS: 7577-7584 KLK9 Kallikrein-related peptidase 9 SEQ ID NOS: 7585-7586 KLKB1 Kallikrein B,
plasma (Fletcher factor) 1 SEQ ID NOS: 7587-7591 SETD8 SET domain containing (lysine
methyltransferase) SEQ ID NOS: 7592-7595 8 KNDC1 Kinase non-catalytic C-lobe domain (KIND) SEQ
ID NOS: 7596-7597 containing 1 KNG1 Kininogen 1 SEQ ID NOS: 7598-7602 KRBA2 KRAB-A
domain containing 2 SEQ ID NOS: 7603-7606 KREMEN2 Kringle containing transmembrane protein 2
SEQ ID NOS: 7607-7612 KRTDAP Keratinocyte differentiation-associated protein SEQ ID NOS: 7613-
7614 L1CAM L1 cell adhesion molecule SEQ ID NOS: 7615-7624 L3MBTL2 L(3)mbt-like 2
(Drosophila) SEQ ID NOS: 7625-7629 LACRT Lacritin SEQ ID NOS: 7630-7632 LACTB Lactamase,
beta SEQ ID NOS: 7633-7635 LAG3 Lymphocyte-activation gene 3 SEQ ID NOS: 7636-7637 LAIR2
Leukocyte-associated immunoglobulin-like SEQ ID NOS: 7638-7641 receptor 2 LALBA Lactalbumin,
alpha- SEQ ID NOS: 7642-7643 LAMA1 Laminin, alpha 1 SEQ ID NOS: 7644-7645 LAMA2 Laminin,
alpha 2 SEQ ID NOS: 7646-7649 LAMA3 Laminin, alpha 3 SEQ ID NOS: 7650-7659 LAMA4 Laminin,
alpha 4 SEQ ID NOS: 7660-7674 LAMA5 Laminin, alpha 5 SEQ ID NOS: 7675-7677 LAMB1 Laminin,
beta 1 SEQ ID NOS: 7678-7682 LAMB2 Laminin, beta 2 (laminin S) SEQ ID NOS: 7683-7685 LAMB3
Laminin, beta 3 SEQ ID NOS: 7686-7690 LAMB4 Laminin, beta 4 SEQ ID NOS: 7691-7694 LAMC1
Laminin, gamma 1 (formerly LAMB2) SEQ ID NOS: 7695-7696 LAMC2 Laminin, gamma 2 SEQ ID
NOS: 7697-7698 LAMC3 Laminin, gamma 3 SEQ ID NOS: 7699-7700 LAMP3 Lysosomal-associated
membrane protein 3 SEQ ID NOS: 7701-7704 GYLTL1B Glycosyltransferase-like 1B SEQ ID NOS:
7705-7710 LAT Linker for activation of T cells SEQ ID NOS: 7711-7720 LAT2 Linker for activation of T
cells family, member 2 SEQ ID NOS: 7721-7729 LBP Lipopolysaccharide binding protein SEQ ID NO:
7730 LCAT Lecithin-cholesterol acyltransferase SEQ ID NOS: 7731-7737 LCN1 Lipocalin 1 SEQ ID
NOS: 7738-7739 LCN10 Lipocalin 10 SEQ ID NOS: 7740-7745 LCN12 Lipocalin 12 SEQ ID NOS:
7746-7748 LCN15 Lipocalin 15 SEQ ID NO: 7749 LCN2 Lipocalin 2 SEQ ID NOS: 7750-7752 LCN6
Lipocalin 6 SEQ ID NOS: 7753-7754 LCN8 Lipocalin 8 SEQ ID NOS: 7755-7756 LCN9 Lipocalin 9
SEQ ID NOS: 7757-7758 LCORL Ligand dependent nuclear receptor corepressor-like SEQ ID NOS:
7759-7764 LDLR Low density lipoprotein receptor SEQ ID NOS: 7765-7773 LDLRAD2 Low density
lipoprotein receptor class A domain SEQ ID NOS: 7774-7775 containing 2 LEAP2 Liver expressed
antimicrobial peptide 2 SEQ ID NO: 7776 LECT2 Leukocyte cell-derived chemotaxin 2 SEQ ID NOS:
7777-7780 LEFTY1 Left-right determination factor 1 SEQ ID NOS: 7781-7782 LEFTY2 Left-right
determination factor 2 SEQ ID NOS: 7783-7784 LEP Leptin SEQ ID NO: 7785 LFNG LFNG O-
fucosylpeptide 3-beta-N- SEQ ID NOS: 7786-7791 acetylglucosaminyltransferase LGALS3BP Lectin,
galactoside-binding, soluble, 3 binding SEQ ID NOS: 7792-7806 protein LGI1 Leucine-rich, glioma
inactivated 1 SEQ ID NOS: 7807-7825 LGI2 Leucine-rich repeat LGI family, member 2 SEQ ID NOS:
7826-7827 LGI3 Leucine-rich repeat LGI family, member 3 SEQ ID NOS: 7828-7831 LGI4 Leucine-rich
repeat LGI family, member 4 SEQ ID NOS: 7832-7835 LGMN Legumain SEQ ID NOS: 7836-7849
LGR4 Leucine-rich repeat containing G protein-coupled SEQ ID NOS: 7850-7852 receptor 4 LHB
Luteinizing hormone beta polypeptide SEQ ID NO: 7853 LHCGR Luteinizing
hormone/choriogonadotropin receptor SEQ ID NOS: 7854-7858 LIF Leukemia inhibitory factor SEQ ID
NOS: 7859-7860 LIFR Leukemia inhibitory factor receptor alpha SEQ ID NOS: 7861-7865 LILRA1
Leukocyte immunoglobulin-like receptor, SEQ ID NOS: 7866-7867 subfamily A (with TM domain),
member 1 LILRA2 Leukocyte immunoglobulin-like receptor, SEQ ID NOS: 7868-7874 subfamily A (with
TM domain), member 2 LILRB3 Leukocyte immunoglobulin-like receptor, SEQ ID NOS: 7875-7879
subfamily B (with TM and ITIM domains), member 3 LIME1 Lck interacting transmembrane adaptor 1
SEQ ID NOS: 7880-7885 LINGO1 Leucine rich repeat and Ig domain containing 1 SEQ ID NOS: 7886-
7896 LIPA Lipase A, lysosomal acid, cholesterol esterase SEQ ID NOS: 7897-7901 LIPC Lipase, hepatic
SEQ ID NOS: 7902-7905 LIPF Lipase, gastric SEQ ID NOS: 7906-7909 LIPG Lipase, endothelial SEQ
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ID NOS: 7910-7915 LIPH Lipase, member H SEQ ID NOS: 7916-7920 LIPK Lipase, family member K
SEQ ID NO: 7921 LIPM Lipase, family member M SEQ ID NOS: 7922-7923 LIPN Lipase, family
member N SEQ ID NO: 7924 LMAN2 Lectin, mannose-binding 2 SEQ ID NOS: 7925-7929 LMNTD1
Lamin tail domain containing 1 SEQ ID NOS: 7930-7940 LNX1 Ligand of numb-protein X 1, E3
ubiquitin protein SEQ ID NOS: 7941-7947 ligase LOX Lysyl oxidase SEQ ID NOS: 7948-7950 LOXL1
Lysyl oxidase-like 1 SEQ ID NOS: 7951-7952 LOXL2 Lysyl oxidase-like 2 SEQ ID NOS: 7953-7961
LOXL3 Lysyl oxidase-like 3 SEQ ID NOS: 7962-7968 LOXL4 Lysyl oxidase-like 4 SEQ ID NO: 7969
LPA Lipoprotein, Lp(a) SEQ ID NOS: 7970-7972 LPL Lipoprotein lipase SEQ ID NOS: 7973-7977 LPO
Lactoperoxidase SEQ ID NOS: 7978-7984 LRAT Lecithin retinol acyltransferase SEQ ID NOS: 7985-
7987 (phosphatidylcholine--retinol O-acyltransferase) LRCH3 Leucine-rich repeats and calponin
homology (CH) SEQ ID NOS: 7988-7996 domain containing 3 LRCOL1 Leucine rich colipase-like 1
SEQ ID NOS: 7997-8000 LRFN4 Leucine rich repeat and fibronectin type III domain SEQ ID NOS:
8001-8002 containing 4 LRFN5 Leucine rich repeat and fibronectin type III domain SEQ ID NOS: 8003-
8005 containing 5 LRG1 Lcucinc-rich alpha-2-glycoprotcin 1 SEQ ID NO: 8006 LRP1 Low density
lipoprotein receptor-related protein 1 SEQ ID NOS: 8007-8012 LRP11 Low density lipoprotein receptor-
related protein 11 SEQ ID NOS: 8013-8014 LRP1B Low density lipoprotein receptor-related protein SEQ
ID NOS: 8015-8018 1B LRP2 Low density lipoprotein receptor-related protein 2 SEQ ID NOS: 8019-
8020 LRP4 Low density lipoprotein receptor-related protein 4 SEQ ID NOS: 8021-8022 LRPAP1 Low
density lipoprotein receptor-related protein SEQ ID NOS: 8023-8024 associated protein 1 LRRC17
Leucine rich repeat containing 17 SEO ID NOS: 8025-8027 LRRC32 Leucine rich repeat containing 32
SEQ ID NOS: 8028-8031 LRRC3B Leucine rich repeat containing 3B SEQ ID NOS: 8032-8036 LRRC4B
Leucine rich repeat containing 4B SEQ ID NOS: 8037-8039 LRRC70 Leucine rich repeat containing 70
SEQ ID NOS: 8040-8041 LRRN3 Leucine rich repeat neuronal 3 SEQ ID NOS: 8042-8045 LRRTM1
Leucine rich repeat transmembrane neuronal 1 SEQ ID NOS: 8046-8052 LRRTM2 Leucine rich repeat
transmembrane neuronal 2 SEQ ID NOS: 8053-8055 LRRTM4 Leucine rich repeat transmembrane
neuronal 4 SEQ ID NOS: 8056-8061 LRTM2 Leucine-rich repeats and transmembrane domains SEQ ID
NOS: 8062-8066 2 LSR Lipolysis stimulated lipoprotein receptor SEQ ID NOS: 8067-8077 LST1
Leukocyte specific transcript 1 SEQ ID NOS: 8078-8095 LTA Lymphotoxin alpha SEQ ID NOS: 8096-
8097 LTBP1 Latent transforming growth factor beta binding SEQ ID NOS: 8098-8107 protein 1 LTBP2
Latent transforming growth factor beta binding SEQ ID NOS: 8108-8111 protein 2 LTBP3 Latent
transforming growth factor beta binding SEQ ID NOS: 8112-8124 protein 3 LTBP4 Latent transforming
growth factor beta binding SEQ ID NOS: 8125-8140 protein 4 LTBR Lymphotoxin beta receptor (TNFR
superfamily, SEQ ID NOS: 8141-8146 member 3) LTF Lactotransferrin SEQ ID NOS: 8147-8151 LTK
Leukocyte receptor tyrosine kinase SEQ ID NOS: 8152-8155 LUM Lumican SEQ ID NO: 8156 LUZP2
Leucine zipper protein 2 SEQ ID NOS: 8157-8160 LVRN Laeverin SEQ ID NOS: 8161-8166 LY6E
Lymphocyte antigen 6 complex, locus E SEQ ID NOS: 8167-8180 LY6G5B Lymphocyte antigen 6
complex, locus G5B SEQ ID NOS: 8181-8182 LY6G6D Lymphocyte antigen 6 complex, locus G6D SEQ
ID NOS: 8183-8184 LY6G6E Lymphocyte antigen 6 complex, locus G6E SEQ ID NOS: 8185-8188
(pseudogene) LY6H Lymphocyte antigen 6 complex, locus H SEQ ID NOS: 8189-8192 LY6K lymphocyte
antigen 6 complex, locus K SEQ ID NOS: 8193-8196 RP11- SEQ ID NO: 8197 520P18.5 LY86
Lymphocyte antigen 86 SEQ ID NOS: 8198-8199 LY96 Lymphocyte antigen 96 SEQ ID NOS: 8200-8201
LYG1 Lysozyme G-like 1 SEQ ID NOS: 8202-8203 LYG2 Lysozyme G-like 2 SEQ ID NOS: 8204-8209
LYNX1 Ly6/neurotoxin 1 SEQ ID NOS: 8210-8214 LYPD1 LY6/PLAUR domain containing 1 SEQ ID
NOS: 8215-8217 LYPD2 LY6/PLAUR domain containing 2 SEQ ID NO: 8218 LYPD4 LY6/PLAUR
domain containing 4 SEQ ID NOS: 8219-8221 LYPD6 LY6/PLAUR domain containing 6 SEQ ID NOS:
8222-8226 LYPD6B LY6/PLAUR domain containing 6B SEQ ID NOS: 8227-8233 LYPD8 LY6/PLAUR
domain containing 8 SEQ ID NOS: 8234-8235 LYZ Lysozyme SEQ ID NOS: 8236-8238 LYZL4
Lysozyme-like 4 SEQ ID NOS: 8239-8240 LYZL6 Lysozyme-like 6 SEQ ID NOS: 8241-8243 M6PR
Mannose-6-phosphate receptor (cation dependent) SEQ ID NOS: 8244-8254 MAD1L1 MAD1 mitotic
arrest deficient-like 1 (yeast) SEQ ID NOS: 8255-8267 MAG Myelin associated glycoprotein SEQ ID
NOS: 8268-8273 MAGT1 Magnesium transporter 1 SEQ ID NOS: 8274-8277 MALSU1 Mitochondrial
assembly of ribosomal large subunit SEQ ID NO: 8278 1 MAMDC2 MAM domain containing 2 SEQ ID
NO: 8279 MAN2B1 Mannosidase, alpha, class 2B, member 1 SEQ ID NOS: 8280-8285 MAN2B2
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Mannosidase, alpha, class 2B, member 2 SEQ ID NOS: 8286-8288 MANBA Mannosidase, beta A,
lysosomal SEQ ID NOS: 8289-8302 MANEAL Mannosidase, endo-alpha-like SEQ ID NOS: 8303-8307
MANF Mesencephalic astrocyte-derived neurotrophic SEQ ID NOS: 8308-8309 factor MANSC1
MANSC domain containing 1 SEQ ID NOS: 8310-8313 MAP3K9 Mitogen-activated protein kinase 9
SEQ ID NOS: 8314-8319 MASP1 Mannan-binding lectin serine peptidase 1 (C4/C2 SEQ ID NOS: 8320-
8327 activating component of Ra-reactive factor) MASP2 Mannan-binding lectin serine peptidase 2 SEQ
ID NOS: 8328-8329 MATN1 Matrilin 1, cartilage matrix protein SEQ ID NO: 8330 MATN2 Matrilin 2
SEQ ID NOS: 8331-8343 MATN3 Matrilin 3 SEQ ID NOS: 8344-8345 MATN4 Matrilin 4 SEQ ID NOS:
8346-8350 MATR3 Matrin 3 SEQ ID NOS: 8351-8378 MAU2 MAU2 sister chromatid cohesion factor
SEQ ID NOS: 8379-8381 MAZ MYC-associated zinc finger protein (purine- SEQ ID NOS: 8382-8396
binding transcription factor) MBD6 Methyl-CpG binding domain protein 6 SEQ ID NOS: 8397-8408
MBL2 Mannose-binding lectin (protein C) 2, soluble SEQ ID NO: 8409 MBNL1 Muscleblind-like
splicing regulator 1 SEQ ID NOS: 8410-8428 MCCC1 Methylcrotonoyl-CoA carboxylase 1 (alpha) SEQ
ID NOS: 8429-8440 MCCD1 Mitochondrial coiled-coil domain 1 SEQ ID NO: 8441 MCEE
Methylmalonyl CoA epimerase SEQ ID NOS: 8442-8445 MCF2L MCF.2 cell line derived transforming
sequence- SEQ ID NOS: 8446-8467 like MCFD2 Multiple coagulation factor deficiency 2 SEQ ID NOS:
8468-8479 MDFIC MyoD family inhibitor domain containing SEQ ID NOS: 8480-8487 MDGA1 MAM
domain containing SEQ ID NOS: 8488-8493 glycosylphosphatidylinositol anchor 1 MDK Midkine
(neurite growth-promoting factor 2) SEQ ID NOS: 8494-8503 MED20 Mediator complex subunit 20 SEQ
ID NOS: 8504-8508 MEGF10 Multiple EGF-like-domains 10 SEQ ID NOS: 8509-8512 MEGF6 Multiple
EGF-like-domains 6 SEQ ID NOS: 8513-8516 MEI1 Meiotic double-stranded break formation protein 1
SEQ ID NOS: 8517-8520 MEI4 Meiotic double-stranded break formation protein 4 SEQ ID NO: 8521
MEIS1 Meis homeobox 1 SEQ ID NOS: 8522-8527 MEIS3 Meis homeobox 3 SEQ ID NOS: 8528-8537
MFI2 Antigen p97 (melanoma associated) identified by SEQ ID NOS: 8538-8540 monoclonal antibodies
133.2 and 96.5 MEPE Matrix extracellular phosphoglycoprotein SEQ ID NOS: 8541-8547 MESDC2
Mesoderm development candidate 2 SEQ ID NOS: 8548-8552 MEST Mesoderm specific transcript SEQ
ID NOS: 8553-8566 MET MET proto-oncogene, receptor tyrosine kinase SEQ ID NOS: 8567-8572
METRN Meteorin, glial cell differentiation regulator SEQ ID NOS: 8573-8577 METRNL Meteorin, glial
cell differentiation regulator-like SEQ ID NOS: 8578-8581 METTL17 Methyltransferase like 17 SEQ ID
NOS: 8582-8592 METTL24 Methyltransferase like 24 SEQ ID NO: 8593 METTL7B Methyltransferase
like 7B SEQ ID NOS: 8594-8595 METTL9 Methyltransferase like 9 SEQ ID NOS: 8596-8604 MEX3C
Mex-3 RNA binding family member C SEQ ID NOS: 8605-8607 MFAP2 Microfibrillar-associated protein
2 SEQ ID NOS: 8608-8609 MFAP3 Microfibrillar-associated protein 3 SEQ ID NOS: 8610-8614
MFAP3L Microfibrillar-associated protein 3-like SEQ ID NOS: 8615-8624 MFAP4 Microfibrillar-
associated protein 4 SEQ ID NOS: 8625-8627 MFAP5 Microfibrillar associated protein 5 SEQ ID NOS:
8628-8638 MFGE8 Milk fat globule-EGF factor 8 protein SEQ ID NOS: 8639-8645 MFNG MFNG O-
fucosylpeptide 3-beta-N- SEQ ID NOS: 8646-8653 acetylglucosaminyltransferase MGA MGA, MAX
dimerization protein SEQ ID NOS: 8654-8662 MGAT2 Mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-
SEQ ID NO: 8663 acetylglucosaminyltransferase MGAT3 Mannosyl (beta-1,4-)-glycoprotein beta-1,4-N-
SEQ ID NOS: 8664-8666 acetylglucosaminyltransferase MGAT4A Mannosyl (alpha-1,3-)-glycoprotein
beta-1,4-N- SEQ ID NOS: 8667-8671 acetylglucosaminyltransferase, isozyme A MGAT4B Mannosyl
(alpha-1,3-)-glycoprotein beta-1,4-N- SEQ ID NOS: 8672-8682 acetylglucosaminyltransferase, isozyme B
MGAT4D MGAT4 family, member D SEQ ID NOS: 8683-8688 MGLL Monoglyceride lipase SEQ ID
NOS: 8689-8698 MGP Matrix Gla protein SEQ ID NOS: 8699-8701 MGST2 Microsomal glutathione S-
transferase 2 SEQ ID NOS: 8702-8705 MIA Melanoma inhibitory activity SEQ ID NOS: 8706-8711
MIA2 Melanoma inhibitory activity 2 SEQ ID NO: 8712 MIA3 Melanoma inhibitory activity family,
member 3 SEQ ID NOS: 8713-8717 MICU1 Mitochondrial calcium uptake 1 SEQ ID NOS: 8718-8727
M1ER1 Mesoderm induction early response 1, SEQ ID NOS: 8728-8736 transcriptional regulator
MINOS1- MINOS1-NBL1 readthrough SEQ ID NOS: 8737-8739 NBL1 MINPP1 Multiple inositol-
polyphosphate phosphatase 1 SEO ID NOS: 8740-8742 MLEC Malectin SEO ID NOS: 8743-8746 MLN
Motilin SEQ ID NOS: 8747-8749 MLXIP MLX interacting protein SEQ ID NOS: 8750-8755 MLXIPL
MLX interacting protein-like SEQ ID NOS: 8756-8763 MMP1 Matrix metallopeptidase 1 SEQ ID NO:
8764 MMP10 Matrix metallopeptidase 10 SEQ ID NOS: 8765-8766 MMP11 Matrix metallopeptidase 11
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SEQ ID NOS: 8767-8770 MMP12 Matrix metallopeptidase 12 SEQ ID NO: 8771 MMP13 Matrix
metallopeptidase 13 SEQ ID NOS: 8772-8774 MMP14 Matrix metallopeptidase 14 (membrane-inserted)
SEQ ID NOS: 8775-8777 MMP17 Matrix metallopeptidase 17 (membrane-inserted) SEQ ID NOS: 8778-
8785 MMP19 Matrix metallopeptidase 19 SEQ ID NOS: 8786-8791 MMP2 Matrix metallopeptidase 2
SEQ ID NOS: 8792-8799 MMP20 Matrix metallopeptidase 20 SEQ ID NO: 8800 MMP21 Matrix
metallopeptidase 21 SEQ ID NO: 8801 MMP25 Matrix metallopeptidase 25 SEQ ID NOS: 8802-8803
MMP26 Matrix metallopeptidase 26 SEQ ID NOS: 8804-8805 MMP27 Matrix metallopeptidase 27 SEQ
ID NO: 8806 MMP28 Matrix metallopeptidase 28 SEQ ID NOS: 8807-8812 MMP3 Matrix
metallopeptidase 3 SEQ ID NOS: 8813-8815 MMP7 Matrix metallopeptidase 7 SEQ ID NO: 8816 MMP8
Matrix metallopeptidase 8 SEQ ID NOS: 8817-8822 MMP9 Matrix metallopeptidase 9 SEQ ID NO: 8823
MMRN1 Multimerin 1 SEQ ID NOS: 8824-8826 MMRN2 Multimerin 2 SEQ ID NOS: 8827-8831
MOXD1 Monooxygenase, DBH-like 1 SEQ ID NOS: 8832-8834 C6orf25 Chromosome 6 open reading
frame 25 SEQ ID NOS: 8835-8842 MPO Myeloperoxidase SEQ ID NOS: 8843-8844 MPPED1
Metallophosphoesterase domain containing 1 SEQ ID NOS: 8845-8848 MPZL1 Myelin protein zero-like
1 SEQ ID NOS: 8849-8853 MR1 Major histocompatibility complex, class I-related SEQ ID NOS: 8854-
8859 MRPL2 Mitochondrial ribosomal protein L2 SEQ ID NOS: 8860-8864 MRPL21 Mitochondrial
ribosomal protein L21 SEQ ID NOS: 8865-8871 MRPL22 Mitochondrial ribosomal protein L22 SEQ ID
NOS: 8872-8876 MRPL24 Mitochondrial ribosomal protein L24 SEQ ID NOS: 8877-8881 MRPL27
Mitochondrial ribosomal protein L27 SEQ ID NOS: 8882-8887 MRPL32 Mitochondrial ribosomal protein
L32 SEO ID NOS: 8888-8890 MRPL34 Mitochondrial ribosomal protein L34 SEO ID NOS: 8891-8895
MRPL35 Mitochondrial ribosomal protein L35 SEQ ID NOS: 8896-8899 MRPL52 Mitochondrial
ribosomal protein L52 SEQ ID NOS: 8900-8910 MRPL55 Mitochondrial ribosomal protein L55 SEQ ID
NOS: 8911-8936 MRPS14 Mitochondrial ribosomal protein S14 SEQ ID NOS: 8937-8938 MRPS22
Mitochondrial ribosomal protein S22 SEQ ID NOS: 8939-8947 MRPS28 Mitochondrial ribosomal protein
S28 SEQ ID NOS: 8948-8955 MS4A14 Membrane-spanning 4-domains, subfamily A, SEQ ID NOS:
8956-8966 member 14 MS4A3 Membrane-spanning 4-domains, subfamily A, SEQ ID NOS: 8967-8971
member 3 (hematopoietic cell-specific) MSH3 MutS homolog 3 SEQ ID NO: 8972 MSH5 MutS homolog
5 SEQ ID NOS: 8973-8984 MSLN Mesothelin SEQ ID NOS: 8985-8992 MSMB Microseminoprotein,
beta- SEQ ID NOS: 8993-8994 MSRA Methionine sulfoxide reductase A SEQ ID NOS: 8995-9002
MSRB2 Methionine sulfoxide reductase B2 SEQ ID NOS: 9003-9004 MSRB3 Methionine sulfoxide
reductase B3 SEQ ID NOS: 9005-9018 MST1 Macrophage stimulating 1 SEQ ID NOS: 9019-9020
MSTN Myostatin SEQ ID NO: 9021 MT1G Metallothionein 1G SEQ ID NOS: 9022-9025 MTHFD2
Methylenetettahydrofolate dehydrogenase SEQ ID NOS: 9026-9030 (NADP+ dependent) 2,
methenyltetrahydrofolate cyclohydrolase MTMR14 Myotubularin related protein 14 SEQ ID NOS: 9031-
9041 MTRNR2L11 MT-RNR2-like 11 (pseudogene) SEQ ID NO: 9042 MTRR 5-methyltetrahydrofolate-
homocysteine SEQ ID NOS: 9043-9055 methyltransferase reductase MTTP Microsomal triglyceride
transfer protein SEQ ID NOS: 9056-9066 MTX2 Metaxin 2 SEQ ID NOS: 9067-9071 MUC1 Mucin 1,
cell surface associated SEQ ID NOS: 9072-9097 MUC13 Mucin 13, cell surface associated SEQ ID NOS:
9098-9099 MUC20 Mucin 20, cell surface associated SEQ ID NOS: 9100-9104 MUC3A Mucin 3A, cell
surface associated SEQ ID NOS: 9105-9107 MUC5AC Mucin 5AC, oligomeric mucus/gel-forming SEQ
ID NO: 9108 MUC5B Mucin 5B, oligomeric mucus/gel-forming SEQ ID NOS: 9109-9110 MUC6 Mucin
6, oligomeric mucus/gel-forming SEQ ID NOS: 9111-9114 MUC7 Mucin 7, secreted SEQ ID NOS: 9115-
9118 MUCL1 Mucin-like 1 SEQ ID NOS: 9119-9121 MXRA5 Matrix-remodelling associated 5 SEQ ID
NO: 9122 MXRA7 Matrix-remodelling associated 7 SEQ ID NOS: 9123-9129 MYDGF Myeloid-derived
growth factor SEQ ID NOS: 9130-9132 MYL1 Myosin, light chain 1, alkali; skeletal, fast SEQ ID NOS:
9133-9134 MYOC Myocilin, trabecular meshwork inducible SEQ ID NOS: 9135-9136 glucocorticoid
response MYRFL Myclin regulatory factor-like SEQ ID NOS: 9137-9141 MZB1 Marginal zone B and B1
cell-specific protein SEQ ID NOS: 9142-9146 N4BP2L2 NEDD4 binding protein 2-like 2 SEQ ID NOS:
9147-9152 NAA38 N(alpha)-acetyltransferase 38, NatC auxiliary SEQ ID NOS: 9153-9158 subunit
NAAA N-acylethanolamine acid amidase SEQ ID NOS: 9159-9164 NAGA N-acetylgalactosaminidase,
alpha- SEQ ID NOS: 9165-9167 NAGLU N-acetylglucosarninidase, alpha SEQ ID NOS: 9168-9172
NAGS N-acetylglutamate synthase SEQ ID NOS: 9173-9174 NAPSA Napsin A aspartic peptidase SEQ
ID NOS: 9175-9177 CARKD Carbohydrate kinase domain containing SEQ ID NOS: 9178-9179
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APOA1BP Apolipoprotein A-I binding protein SEQ ID NOS: 9180-9182 NBL1 Neuroblastoma 1, DAN
family BMP antagonist SEQ ID NOS: 9183-9196 NCAM1 Neural cell adhesion molecule 1 SEQ ID NOS:
9197-9216 NCAN Neurocan SEQ ID NOS: 9217-9218 NCBP2-AS2 NCBP2 antisense RNA 2 (head to
head) SEQ ID NO: 9219 NCSTN Nicastrin SEQ ID NOS: 9220-9229 NDNF Neuron-derived neurotrophic
factor SEQ ID NOS: 9230-9232 NDP Norrie disease (pseudoglioma) SEQ ID NOS: 9233-9235 NDUFA10
NADH dehydrogenase (ubiquinone) 1 alpha SEQ ID NOS: 9236-9245 subcomplex, 10, 42 kDa NDUFB5
NADH dehydrogenase (ubiquinone) 1 beta SEQ ID NOS: 9246-9254 subcomplex, 5, 16 kDa NDUFS8
NADH dehydrogenase (ubiquinone) Fe—S protein SEQ ID NOS: 9255-9264 8, 23 kDa (NADH-
coenzyme Q reductase) NDUFV1 NADH dehydrogenase (ubiquinone) flavoprotein SEQ ID NOS: 9265-
9278 1, 51 kDa NECAB3 N-terminal EF-hand calcium binding protein 3 SEQ ID NOS: 9279-9288
PVRL1 Poliovirus receptor-related 1 (herpesvirus entry SEQ ID NOS: 9289-92.91 mediator C) NELL1
Neural EGFL like 1 SEQ ID NOS: 9292-9295 NELL2 Neural EGFL like 2 SEQ ID NOS: 9296-9310
NENF Neudesin neurotrophic factor SEQ ID NO: 9311 NETO1 Neuropilin (NRP) and tolloid (TLL)-like
1 SEQ ID NOS: 9312-9316 NFASC Neurofascin SEQ ID NOS: 9317-9331 NFE2L1 Nuclear factor,
erythroid 2-like 1 SEQ ID NOS: 9332-9350 NFE2L3 Nuclear factor, erythroid 2-like 3 SEQ ID NOS:
9351-9352 NGEF Neuronal guaninc nucleotide exchange factor SEQ ID NOS: 9353-9358 NGF Nerve
growth factor (beta polypeptide) SEQ ID NO: 9359 NGLY1 N-glycanase 1 SEQ ID NOS: 9360-9366
NGRN Neugrin, neurite outgrowth associated SEQ ID NOS: 9367-9368 NHLRC3 NHL repeat containing
3 SEQ ID NOS: 9369-9371 NID1 Nidogen 1 SEQ ID NOS: 9372-9373 NID2 Nidogen 2 (osteonidogen)
SEO ID NOS: 9374-9376 NKG7 Natural killer cell granule protein 7 SEO ID NOS: 9377-9381 NLGN3
Neuroligin 3 SEQ ID NOS: 9382-9386 NLGN4Y Neuroligin 4, Y-linked SEQ ID NOS: 9387-9393
NLRP5 NLR family, pyrin domain containing 5 SEQ ID NOS: 9394-9396 NMB Neuromedin B SEQ ID
NOS: 9397-9398 NME1 NME/NM23 nucleoside diphosphate kinase 1 SEQ ID NOS: 9399-9405 NME1-
NME1-NME2 readthrough SEQ ID NOS: 9406-9408 NME2 NME3 NME/NM23 nucleoside diphosphate
kinase 3 SEQ ID NOS: 9409-9413 NMS Neuromedin S SEQ ID NO: 9414 NMU Neuromedin U SEQ ID
NOS: 9415-9418 NOA1 Nitric oxide associated 1 SEQ ID NO: 9419 NODAL Nodal growth
differentiation factor SEQ ID NOS: 9420-9421 NOG Noggin SEQ ID NO: 9422 NOMO3 NODAL
modulator 3 SEQ ID NOS: 9423-9429 NOS1AP Nitric oxide synthase 1 (neuronal) adaptor protein SEQ
ID NOS: 9430-9434 NOTCH3 Notch 3 SEQ ID NOS: 9435-9438 NOTUM Notum pectinacetylesterase
homolog (Drosophila) SEQ ID NOS: 9439-9441 NOV Nephroblastoma overexpressed SEQ ID NO: 9442
NPB Neuropeptide B SEQ ID NOS: 9443-9444 NPC2 Niemann-Pick disease, type C2 SEQ ID NOS:
9445-9453 NPFF Neuropeptide FF-amide peptide precursor SEQ ID NO: 9454 NPFFR2 Neuropeptide FF
receptor 2 SEQ ID NOS: 9455-9458 NPHS1 Nephrosis I, congenital, Finnish type (nephrin) SEQ ID
NOS: 9459-9460 NPNT Nephronectin SEQ ID NOS: 9461-9471 NPPA Natriuretic peptide A SEQ ID
NOS: 9472-9474 NPPB Natriuretic peptide B SEQ ID NO: 9475 NPPC Natriuretic peptide C SEQ ID
NOS: 9476-9477 NPS Neuropeptide S SEQ ID NO: 9478 NPTX1 Neuronal pentraxin I SEQ ID NO: 9479
NPTX2 Neuronal pentraxin II SEQ ID NO: 9480 NPTXR Neuronal pentraxin receptor SEQ ID NOS:
9481-9482 NPVF Neuropeptide VF precursor SEQ ID NO: 9483 NPW Neuropeptide W SEQ ID NOS:
9484-9486 NPY Neuropeptide Y SEQ ID NOS: 9487-9489 NQO2 NAD(P)H dehydrogenase, quinone 2
SEQ ID NOS: 9490-9498 NRCAM Neuronal cell adhesion molecule SEQ ID NOS: 9499-9511 NRG1
Neuregulin 1 SEQ ID NOS: 9512-9529 NRN1L Neuritin 1-like SEQ ID NOS: 9530-9532 NRP1
Neuropilin 1 SEQ ID NOS: 9533-9546 NRP2 Neuropilin 2 SEQ ID NOS: 9547-9553 NRTN Neurturin
SEQ ID NO: 9554 NRXN1 Neurexin 1 SEQ ID NOS: 9555-9585 NRXN2 Neurexin 2 SEQ ID NOS:
9586-9594 NT5C3A 5'-nucleotidase, cytosolic IIIA SEQ ID NOS: 9595-9605 NT5DC3 5'-nucleotidase
domain containing 3 SEQ ID NOS: 9606-9608 NT5E 5'-nucleotidase, ecto (CD73) SEQ ID NOS: 9609-
9613 NTF3 Neurotrophin 3 SEQ ID NOS: 9614-9615 NTF4 Neurotrophin 4 SEQ ID NOS: 9616-9617
NTM Neurotrimin SEQ ID NOS: 9618-9627 NTN1 Netrin 1 SEQ ID NOS: 9628-9629 NTN3 Netrin 3
SEQ ID NO; 9630 NTN4 Netrin 4 SEQ ID NOS: 9631-9635 NTN5 Netrin 5 SEQ ID NOS: 9636-9637
NTNG1 Netrin G1 SEQ ID NOS: 9638-9644 NTNG2 Netrin G2 SEQ ID NOS: 9645-9646 NTS
Neurotensin SEQ ID NOS: 9647-9648 NUBPL Nucleotide binding protein-like SEQ ID NOS: 9649-9655
NUCB1 Nucleobindin 1 SEQ ID NOS: 9656-9662 NUCB2 Nucleobindin 2 SEQ ID NOS: 9663-9678
NUDT19 Nudix (nucleoside diphosphate linked moiety X)- SEQ ID NO: 9679 type motif 19 NUDT9
Nudix (nucleoside diphosphate linked moiety X)- SEQ ID NOS: 9680-9684 type motif 9 NUP155
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Nucleoporin 155 kDa SEQ ID NOS: 9685-9688 NUP214 Nucleoporin 214 kDa SEQ ID NOS: 9689-9700
NUP85 Nucleoporin 85 kDa SEQ ID NOS: 9701-9715 NXPE3 Neurexopbilin and PC-esterase domain
family, SEQ ID NOS: 9716-9721 member 3 NXPE4 Neurexopbilin and PC-esterase domain family, SEQ
ID NOS: 9722-9723 member 4 NXPH1 Neurexopbilin 1 SEQ ID NOS: 9724-9727 NXPH2 Neurexophilin
2 SEQ ID NO: 9728 NXPH3 Neurexophilin 3 SEQ ID NOS: 9729-9730 NXPH4 Neurexophilin 4 SEQ ID
NOS: 9731-9732 NYX Nyctalopin SEQ ID NOS: 9733-9734 OAF Out at first homolog SEQ ID NOS:
9735-9736 OBP2A Odorant binding protein 2A SEQ ID NOS: 9737-9743 OBP2B Odorant binding
protein 2B SEQ ID NOS: 9744-9747 OC90 Otoconin 90 SEQ ID NO: 9748 OCLN Occludin SEQ ID
NOS: 9749-9751 ODAM Odontogenic, ameloblast asssociated SEQ ID NOS: 9752-9755 C4orf26
Chromosome 4 open reading frame 26 SEQ ID NOS: 9756-9759 OGG1 8-oxoguanine DMA glycosylase
SEQ ID NOS: 9760-9773 OGN Osteoglycin SEQ ID NOS: 9774-9776 OIT3 Oncoprotein induced
transcript 3 SEQ ID NOS: 9777-9778 OLFM1 Olfactomedin 1 SEQ ID NOS: 9779-9789 OLFM2
Olfactomedin 2 SEQ ID NOS: 9790-9793 OLFM3 Olfactomedin 3 SEQ ID NOS: 9794-9796 OLFM4
Olfactomedin 4 SEQ ID NO: 9797 OLFML1 Olfactomedin-like 1 SEQ ID NOS: 9798-9801 OLFML2A
Olfactomedin-like 2A SEQ ID NOS: 9802-9804 OLFML2B Olfactomedin-like 2B SEQ ID NOS: 9805-
9809 OLFML3 Olfactomedin-like 3 SEQ ID NOS: 9810-9812 OMD Osteomodulin SEQ ID NO: 9813
OMG Oligodendrocyte myelin glycoprotein SEQ ID NO: 9814 OOSP2 Oocyte secreted protein 2 SEQ ID
NOS: 9815-9816 OPCML Opioid binding protein/cell adhesion molecule-like SEQ ID NOS: 9817-9821
PROL1 Proline rich, lacrimal 1 SEQ ID NO: 9822 OPTC Opticin SEQ ID NOS: 9823-9824 ORAI1 ORAI
calcium release-activated calcium modulator SEQ ID NO: 9825 1 ORM1 Orosomucoid 1 SEQ ID NO:
9826 ORM2 Orosomucoid 2 SEQ ID NO: 9827 ORMDL2 ORMDL sphingolipid biosynthesis regulator 2
SEQ ID NOS: 9828-9831 OS9 Osteosarcoma amplified 9, endoplasmic reticulum SEQ ID NOS: 9832-
9846 lectin OSCAR Osteoclast associated, immunoglobulin-like SEQ ID NOS: 9847-9857 receptor OSM
Oncostatin M SEQ ID NOS: 9858-9860 OSMR Oncostatin M receptor SEQ ID NOS: 9861-9865 OSTN
Osteocrin SEQ ID NOS: 9866-9867 OTOA Otoancorin SEQ ID NOS: 9868-9873 OTOG Otogelin SEQ
ID NOS: 9874-9876 OTOGL Otogelin-like SEQ ID NOS: 9877-9883 OTOL1 Otolin 1 SEQ ID NO: 9884
OTOR Otoraplin SEQ ID NO: 9885 OTOS Otospiralin SEQ ID NOS: 9886-9887 OVCH1 Ovochymase 1
SEQ ID NOS: 9888-9890 OVCH2 Ovochymase 2 (gene/pseudogene) SEQ ID NOS: 9891-9892 OVGP1
Oviductal glycoprotein 1, 120 kDa SEQ ID NO: 9893 OXCT1 3-oxoacid CoA transferase 1 SEQ ID NOS:
9894-9897 OXCT2 3-oxoacid CoA transferase 2 SEQ ID NO: 9898 OXNAD1 Oxidoreductase NAD-
binding domain containing 1 SEQ ID NOS: 9899-9905 OXT Oxytocin/neurophysin I prepropeptide SEQ
ID NO: 9906 P3H1 Prolyl 3-hydroxylase 1 SEQ ID NOS: 9907-9911 P3H2 Prolyl 3-hydroxylase 2 SEQ
ID NOS: 9912-9915 P3H3 Prolyl 3-hydroxylase 3 SEQ ID NO: 9916 P3H4 Prolyl 3-hydroxylase family
member 4 (non- SEQ ID NOS: 9917-9921 enzymatic) P4HA1 Prolyl 4-hydroxylase, alpha polypeptide I
SEQ ID NOS: 9922-9926 P4HA2 Prolyl 4-hydroxylase, alpha polypeptide II SEQ ID NOS: 9927-9941
P4HA3 Prolyl 4-hydroxylase, alpha polypeptide III SEQ ID NOS: 9942-9946 P4HB Prolyl 4-hydroxylase,
beta polypeptide SEQ ID NOS: 9947-9958 PAEP Progestagen-associated endometrial protein SEQ ID
NOS: 9959-9967 PAM Peplidylglycine alpha-amidating monooxygenase SEQ ID NOS: 9968-9981
PAMR1 Peptidase domain containing associated with SEQ ID NOS: 9982-9988 muscle regeneration 1
PAPLN Papilin, proteoglycan-like sulfated glycoprotein SEQ ID NOS: 9989-9996 PAPPA Pregnancy-
associated plasma protein A, pappalysin SEQ ID NO: 9997 1 PAPPA2 Pappalysin 2 SEQ ID NOS: 9998-
9999 PARP15 Poly (ADP-ribose) polymerase family, member 15 SEQ ID NOS: 10000-10003 PARVB
Parvin, beta SEQ ID NOS: 10004-10008 PATE1 Prostate and testis expressed 1 SEQ ID NOS: 10009-
10010 PATE2 Prostate and testis expressed 2 SEQ ID NOS: 10011-10012 PATE3 Prostate and testis
expressed 3 SEQ ID NO: 10013 PATE4 Prostate and testis expressed 4 SEQ ID NOS: 10014-10015
PATL2 Protein associated with topoisomerase II homolog SEQ ID NOS: 10016-10021 2 (yeast) PAX2
Paired box 2 SEQ ID NOS: 10022-10027 PAX4 Paired box 4 SEQ ID NOS: 10028-10034 PCCB
Propionyl CoA carboxylase, beta polypeptide SEQ ID NOS: 10035-10049 PCDH1 Protocadherin 1 SEQ
ID NOS: 10050-10055 PCDH12 Protocadherin 12 SEQ ID NOS: 10056-10057 PCDH15 Protocadherin-
related 15 SEO ID NOS: 10058-10091 PCDHA1 Protocadherin alpha 1 SEO ID NOS: 10092-10094
PCDHA10 Protocadherin alpha 10 SEQ ID NOS: 10095-10097 PCDHA11 Protocadherin alpha 11 SEQ
ID NOS: 10098-10100 PCDHA6 Protocadherin alpha 6 SEQ ID NOS: 10101-10103 PCDHB12
Protocadherin beta 12 SEQ ID NOS: 10104-10106 PCDHGA11 Protocadherin gamma subfamily A, 11
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SEQ ID NOS: 10107-10109 PCF11 PCF11 cleavage and polyadenylation factor SEQ ID NOS: 10110-
10114 subunit PCOLCE Procollagen C-endopeptidase enhancer SEQ ID NO: 10115 PCOLCE2
Procollagen C-endopeptidase enhancer 2 SEQ ID NOS: 10116-10119 PCSK1 Proprotein convertase
subtilisin/kexin type 1 SEQ ID NOS: 10120-10122 PCSK1N Proprotein convertase subtilisin/kexin type 1
SEQ ID NO: 10123 inhibitor PCSK2 Proprotein convertase subtilisin/kexin type 2 SEQ ID NOS: 10124-
10126 PCSK4 Proprotein convertase subtilisin/kexin type 4 SEQ ID NOS: 10127-10129 PCSK5
Proprotein convertase subtilisin/kexin type 5 SEQ ID NOS: 10130-10134 PCSK9 Proprotein convertase
subtilisin/kexin type 9 SEQ ID NO: 10135 PCYOX1 Prenylcysteine oxidase 1 SEQ ID NOS: 10136-
10140 PCYOX1L Prenylcysteine oxidase 1 like SEQ ID NOS: 10141-10145 PDE11A Phosphodicsterase
11A SEO ID NOS: 10146-10151 PDE2A Phosphodiesterase 2A, cGMP-stimulated SEO ID NOS: 10152-
10173 PDE7A Phosphodiesterase 7A SEQ ID NOS: 10174-10177 PDF Peptide deformylase
(mitochondrial) SEQ ID NO: 10178 PDGFA Platelet-derived growth factor alpha polypeptide SEQ ID
NOS: 10179-10182 PDGFB Platelet-derived growth factor beta polypeptide SEQ ID NOS: 10183-10186
PDGFC Platelet derived growth factor C SEQ ID NOS: 10187-10190 PDGFD Platelet derived growth
factor D SEQ ID NOS: 10191-10193 PDGFRA Platelet-derived growth factor receptor, alpha SEQ ID
NOS: 10194-10200 polypeptide PDGFRB Platelet-derived growth factor receptor, beta SEQ ID NOS:
10201-10204 polypeptide PDGFRL Platelet-derived growth factor receptor-like SEO ID NOS: 10205-
10206 PDHA1 Pyruvate dehydrogenase (lipoamide) alpha 1 SEQ ID NOS: 10207-10215 PDIA2 Protein
disulfide isomerase family A, member 2 SEQ ID NOS: 10216-10219 PDIA3 Protein disulfide isomerase
family A, member 3 SEO ID NOS: 10220-10223 PDIA4 Protein disulfide isomerase family A, member 4
SEQ ID NOS: 10224-10225 PDIA5 Protein disulfide isomerase family A, member 5 SEQ ID NOS:
10226-10229 PDIA6 Protein disulfide isomerase family A, member 6 SEQ ID NOS: 10230-10236 PDILT
Protein disulfide isomerase-like, testis expressed SEQ ID NOS: 10237-10238 PDYN Prodynorphin SEQ
ID NOS: 10239-10241 PDZD8 PDZ domain containing 8 SEQ ID NO: 10242 PDZRN4 PDZ domain
containing ring finger 4 SEQ ID NOS: 10243-10245 PEAR1 Platelet endothelial aggregation receptor 1
SEQ ID NOS: 10246-10249 PEBP4 Phosphatidylethanolamine-binding protein 4 SEQ ID NOS: 10250-
10251 PECAM1 Platelet/endothelial cell adhesion molecule 1 SEQ ID NOS: 10252-10255 PENK
Proenkephalin SEQ ID NOS: 10256-10261 PET117 PET117 homolog SEQ ID NO: 10262 PF4 Platelet
factor 4 SEQ ID NO: 10263 PF4V1 Platelet factor 4 variant 1 SEQ ID NO: 10264 PFKP
Phosphofructokinase, platelet SEQ ID NOS: 10265-10273 PFN1 Profilin 1 SEQ ID NOS: 10274-10276
PGA3 Pepsinogen 3, group I (pepsinogen A) SEQ ID NOS: 10277-10280 PGA4 Pepsinogen 4, group I
(pepsinogen A) SEQ ID NOS: 10281-10283 PGA5 Pepsinogen 5, group I (pepsinogen A) SEQ ID NOS:
10284-10286 PGAM5 PGAM family member 5, serine/threonine protein SEQ ID NOS: 10287-10290
phosphatase, mitochondrial PGAP3 Post-GPI attachment to proteins 3 SEQ ID NOS: 10291-10298 PGC
Progastricsin (pepsinogen C) SEQ ID NOS: 10299-10302 PGF Placental growth factor SEQ ID NOS:
10303-10306 PGLYRP1 Peptidoglycan recognition protein 1 SEQ ID NO: 10307 PGLYRP2
Peptidoglycan recognition protein 2 SEQ ID NOS: 10308-10311 PGLYRP3 Peptidoglycan recognition
protein 3 SEQ ID NO: 10312 PGLYRP4 Peptidoglycan recognition protein 4 SEQ ID NOS: 10313-10314
PHACTR1 Phosphatase and actin regulator 1 SEQ ID NOS: 10315-10321 PHB Prohibitin SEQ ID NOS:
10322-10330 PI15 Peptidase inhibitor 15 SEQ ID NOS: 10331-10332 PI3 Peptidase inhibitor 3, skin-
derived SEQ ID NO: 10333 PIANP PILR alpha associated neural protein SEQ ID NOS: 10334-10339
PICK Phosphatidylinositol glycan anchor biosynthesis, SEQ ID NOS: 10340-10343 class K PIGL
Phosphatidylinositol glycan anchor biosynthesis, SEQ ID NOS: 10344-10351 class L PIGT
Phosphatidylinositol glycan anchor biosynthesis, SEQ ID NOS: 10352-10406 class T PIGZ
Phosphatidylinositol glycan anchor biosynthesis, SEQ ID NOS: 10407-10409 class Z PIK3AP1
Phosphoinositide-3-kinase adaptor protein 1 SEQ ID NOS: 10410-10412 PIK3IP1 Phosphoinositide-3-
kinase interacting protein 1 SEQ ID NOS: 10413-10416 PILRA Paired immunoglobin-like type 2 receptor
alpha SEQ ID NOS: 10417-10421 PILRB Paired immunoglobin-like type 2 receptor beta SEQ ID NOS:
10422-10433 PINLYP Phospholipase A2 inhibitor and LY6/PLAUR SEQ ID NOS: 10434-10438 domain
containing PIP Prolactin-induced protein SEQ ID NO: 10439 PIWIL4 Piwi-like RNA-mediated gene
silencing 4 SEQ ID NOS: 10440-10444 PKDCC Protein kinase domain containing, cytoplasmic SEQ ID
NOS: 10445-10446 PKHD1 Polycystic kidney and hepatic disease 1 (autosomal SEQ ID NOS: 10447-
10448 recessive) PLA1A Phospholipase A1 member A SEQ ID NOS: 10449-10453 PLA2G10
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Phospholipase A2, group X SEQ ID NOS: 10454-10455 PLA2G12A Phospholipase A2, group XIIA SEQ
ID NOS: 10456-10458 PLA2G12B Phospholipase A2, group XIIB SEQ ID NO: 10459 PLA2G15
Phospholipase A2, group XV SEQ ID NOS: 10460-10467 PLA2G1B Phospholipase A2, group IB
(pancreas) SEQ ID NOS: 10468-10470 PLA2G2A Phospholipase A2, group IIA (platelets, synovial SEQ
ID NOS: 10471-10472 fluid) PLA2G2C Phospholipase A2, group IIC SEQ ID NOS: 10473-10474
PLA2G2D Phospholipase A2, group IID SEQ ID NOS: 10475-10476 PLA2G2E Phospholipase A2, group
IIE SEQ ID NO: 10477 PLA2G3 Phospholipase A2, group III SEQ ID NO: 10478 PLA2G5
Phospholipase A2, group V SEQ ID NO: 10479 PLA2G7 Phospholipase A2, group VII (platelet-activating
SEQ ID NOS: 10480-10481 factor acetylhydrolase, plasma) PLA2R1 Phospholipase A2 receptor 1, 180
kDa SEQ ID NOS: 10482-10483 PLAC1 Placenta-specific 1 SEQ ID NO: 10484 PLAC9 Placenta-
specific 9 SEQ ID NOS: 10485-10487 PLAT Plasminogen activator, tissue SEQ ID NOS: 10488-10496
PLAU Plasminogen activator, urokinase SEQ ID NOS: 10497-10499 PLAUR Plasminogen activator,
urokinase receptor SEQ ID NOS: 10500-10511 PLBD1 Phospholipase B domain containing 1 SEQ ID
NOS: 10512-10514 PLBD2 Phospholipase B domain containing 2 SEQ ID NOS: 10515-10517 PLG
Plasminogen SEQ ID NOS: 10518-10520 PLGLB1 Plasminogen-like B1 SEQ ID NOS: 10521-10524
PLGLB2 Plasminogen-like B2 SEQ ID NOS: 10525-10526 PLOD1 Procollagen-lysine, 2-oxoglutarate 5-
dioxygenase 1 SEQ ID NOS: 10527-10529 PLOD2 Procollagen-lysine, 2-oxoglutarate 5-dioxygenase 2
SEQ ID NOS: 10530-10535 PLOD3 Procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 SEQ ID NOS:
10536-10542 PLTP Phospholipid transfer protein SEQ ID NOS: 10543-10547 PLXNA4 Plexin A4 SEQ
ID NOS: 10548-10551 PLXNB2 Plexin B2 SEQ ID NOS: 10552-10560 PM20D1 Peptidase M20 domain
containing 1 SEQ ID NO: 10561 PMCH Pro-melanin-concentrating hormone SEQ ID NO: 10562 PMEL
Premelanosorne protein SEQ ID NOS: 10563-10574 PMEPA1 Prostate transmembrane protein, androgen
induced SEQ ID NOS: 10575-10581 1 PNLIP Pancreatic lipase SEQ ID NO: 10582 PNLIPRP1 Pancreatic
lipase-related protein 1 SEQ ID NOS: 10583-10591 PNLIPRP3 Pancreatic lipase-related protein 3 SEQ
ID NO: 10592 FNOC Prepronociceptin SEQ ID NOS: 10593-10595 PNP Purine nucleoside phosphorylase
SEQ ID NOS: 10596-10599 PNPLA4 Patatin-like phospholipase domain containing 4 SEQ ID NOS:
10600-10603 PODNL1 Podocan-like 1 SEQ ID NOS: 10604-10615 POFUT1 Protein O-
fucosyltransferase 1 SEQ ID NOS: 10616-10617 POFUT2 Protein O-fucosyltransferase 2 SEQ ID NOS:
10618-10623 POGLUT1 Protein O-glucosyltransferase 1 SEQ ID NOS: 10624-10628 POLL Polymerase
(DNA directed), lambda SEQ ID NOS: 10629-10641 POMC Proopiomelanocortin SEQ ED NOS: 10642-
10646 POMGNT2 Protein O-linked mannose N- SEQ ID NOS: 10647-10648
acetylglucosaminyltransferase 2 (beta 1,4-) PON1 Paraoxonase 1 SEQ ID NOS: 10649-10650 PON2
Paraoxonase 2 SEQ ID NOS: 10651-10663 PON3 Paraoxonase 3 SEQ ID NOS: 10664-10669 POSTN
Periostin, osteoblast specific factor SEQ ID NOS: 10670-10675 PPBP Pro-platelet basic protein
(chemokine (C-X-C SEQ ID NO: 10676 motif) ligand 7) PPIB Peptidylprolyl isomerase B (cyclophilin B)
SEQ ID NO: 10677 PPIC Peptidylprolyl isomerase C (cyclophilin C) SEQ ID NO: 10678 PPOX
Protoporphyrinogen oxidase SEQ ID NOS: 10679-10689 PPP1CA Protein phosphatase 1, catalytic
subunit, alpha SEQ ID NOS: 10690-10695 isozyme PPT1 Palmitoyl-protein thioesterase 1 SEQ ID NOS:
10696-10712 PPT2 Palmitoyl-protein thioesterase 2 SEQ ID NOS: 10713-10720 PPY Pancreatic
polypeptide SEQ ID NOS: 10721-10725 PRAC2 Prostate cancer susceptibility candidate 2 SEQ ID NOS:
10726-10727 PRADC1 Protease-associated domain containing 1 SEQ ID NO: 10728 PRAP1 Proline-rich
acidic protein 1 SEQ ID NOS: 10729-10730 PRB1 Proline-rich protein BstNI subfamily 1 SEQ ID NOS:
10731-10734 PRB2 Proline-rich protein BstNI subfamily 2 SEQ ID NOS: 10735-10736 PRB3 Proline-
rich protein BstNI subfamily 3 SEQ ID NOS: 10737-10738 PRB4 Proline-rich protein BstNI subfamily 4
SEQ ID NOS: 10739-10742 PRCD Progressive rod-cone degeneration SEQ ID NOS: 10743-10744 PRCP
Prolylcarboxypeptidase (angiotensinase C) SEQ ID NOS: 10745-10756 PRDM12 PR domain containing
12 SEQ ID NO: 10757 PRDX4 Peroxiredoxin 4 SEQ ID NOS: 10758-10761 PRELP Proline/arginine-rich
end leucine-rich repeat SEQ ID NO: 10762 protein PRF1 Perforin 1 (pore forming protein) SEQ ID NOS:
10763-10765 PRG2 Proteoglycan 2, bone marrow (natural killer cell SEQ ID NOS: 10766-10768
activator, eosinophil granule major basic protein) PRG3 Proteoglycan 3 SEQ ID NO: 10769 PRG4
Proteoglycan 4 SEQ ID NOS: 10770-10775 PRH1 Proline-rich protein HaeIII subfamily 1 SEQ ID NOS:
10776-10778 PRH2 Proline-rich protein HaeIII subfamily 2 SEQ ID NOS: 10779-10780 PRKAG1
Protein kinase, AMP-activated, gamma 1 non- SEQ ID NOS: 10781-10795 catalytic subunit PRKCSH
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Protein kinase C substrate 80K-H SEQ ID NOS: 10796-10805 PRKD1 Protein kinase D1 SEQ ID NOS:
10806-10811 PRL Prolactin SEQ ID NOS: 10812-10814 PRLH Prolactin releasing hormone SEQ ID NO:
10815 PRLR Prolactin receptor SEQ ID NOS: 10816-10834 PRNP Prion protein SEQ ID NOS: 10835-
10838 PRNT Prion protein (testis specific) SEQ ID NO: 10839 PROC Protein C (inactivator of
coagulation factors Va SEQ ID NOS: 10840-10847 and VIIIa) PROK1 Prokineticin 1 SEQ ID NO: 10848
PROK2 Prokineticin 2 SEQ ID NOS: 10849-10850 PROM1 Prominin 1 SEQ ID NOS: 10851-10862
PROS1 Protein S (alpha) SEQ ID NOS: 10863-10866 PROZ Protein Z, vitamin K-dependent plasma SEQ
ID NOS: 10867-10868 glycoprotein PRR27 Proline rich 27 SEQ ID NOS: 10869-10872 PRR4 Proline
rich 4 (lacrimal) SEQ ID NOS: 10873-10875 PRRG2 Proline rich Gla (G-carboxyglutamic acid) 2 SEQ
ID NOS: 10876-10878 PRRT3 Proline-rich transmembrane protein 3 SEQ ID NOS: 10879-10881 PRRT4
Proline-rich transmembrane protein 4 SEQ ID NOS: 10882-10888 PRSS1 Protease, serine, 1 (trypsin 1)
SEQ ID NOS: 10889-10892 PRSS12 Protease, serine, 12 (neurotrypsin, motopsin) SEQ ID NO: 10893
PRSS16 Protease, serine, 16 (thymus) SEQ ID NOS: 10894-10901 PRSS2 Protease, serine, 2 (trypsin 2)
SEQ ID NOS: 10902-10905 PRSS21 Protease, serine, 21 (testisin) SEQ ID NOS: 10906-10911 PRSS22
Protease, serine, 22 SEQ ID NOS: 10912-10914 PRSS23 Protease, serine. 23 SEQ ID NOS: 10915-10918
PRSS27 Protease, serine 27 SEQ ID NOS: 10919-10921 PRSS3 Protease, serine, 3 SEQ ID NOS: 10922-
10926 PRSS33 Protease, serine, 33 SEQ ID NOS: 10927-10930 PRSS35 Protease, serine, 35 SEQ ID NO:
10931 PRSS36 Protease, serine, 36 SEQ ID NOS: 10932-10935 PRSS37 Protease, serine, 37 SEQ ID
NOS: 10936-10939 PRSS38 Protease, serine, 38 SEQ ID NO: 10940 PRSS42 Protease, serine, 42 SEQ ID
NOS: 10941-10942 PRSS48 Protease, serine, 48 SEQ ID NOS: 10943-10944 PRSS50 Protease, serine, 50
SEQ ID NO: 10945 PRSS53 Protease, serine, 53 SEQ ID NO: 10946 PRSS54 Protease, serine, 54 SEQ ID
NOS: 10947-10951 PRSS55 Protease, serine, 55 SEQ ID NOS: 10952-10954 PRSS56 Protease, serine, 56
SEQ ID NOS: 10955-10956 PRSS57 Protease, serine, 57 SEQ ID NOS: 10957-10958 PRSS58 Protease,
serine, 58 SEQ ID NOS: 10959-10960 PRSS8 Protease, serine, 8 SEQ ID NOS: 10961-10964 PRTG
Protogenin SEQ ID NOS: 10965-10968 PRTN3 Proteinase 3 SEQ ID NOS: 10969-10970 PSAP
Prosaposin SEQ ID NOS: 10971-10974 PSAPL1 Prosaposin-like 1 (gene/pseudogene) SEQ ID NO:
10975 PSG1 Pregnancy specific beta-1-glycoprotein 1 SEQ ID NOS: 10976-10983 PSG11 Pregnancy
specific beta-1-glycoprotein 11 SEQ ID NOS: 10984-10988 PSG2 Pregnancy specific beta-1-glycoprotein
2 SEQ ID NOS: 10989-10990 PSG3 Pregnancy specific beta-1-glycoprotein 3 SEQ ID NOS: 10991-
10994 PSG4 Pregnancy specific beta-1-glycoprotein 4 SEQ ID NOS: 10995-11006 PSG5 Pregnancy
specific beta-1-glycoprotein 5 SEQ ID NOS: 11007-11012 PSG6 Pregnancy specific beta-1-glycoprotein 6
SEQ ID NOS: 11013-11018 PSG7 Pregnancy specific beta-1-glycoprotein 7 SEQ ID NOS: 11019-11021
(gene/pseudogene) PSG8 Pregnancy specific beta-1-glycoprotein 8 SEQ ID NOS: 11022-11026 PSG9
Pregnancy specific beta-1-glycoprotein 9 SEQ ID NOS: 11027-11034 PSMD1 Proteasome 26S subunit,
non-ATPase 1 SEQ ID NOS: 11035-11042 PSORS1C2 Psoriasis susceptibility 1 candidate 2 SEQ ID NO:
11043 PSPN Persephin SEQ ID NOS: 11044-11045 PTGDS Prostaglandin D2 synthase 21 kDa (brain)
SEQ ID NOS: 11046-11050 PTGIR Prostaglandin I2 (prostacyclin) receptor (IP) SEQ ID NOS: 11051-
11055 PTGS1 Prostaglandin-endoperoxide synthase 1 SEQ ID NOS: 11056-11064 (prostaglandin G/H
synthase and cyclooxygenase) PTGS2 Prostaglandin-endoperoxide synthase 2 SEQ ID NOS: 11065-11066
(prostaglandin G/H synthase and cyclooxygenase) PTH Parathyroid hormone SEQ ID NOS: 11067-11068
PTH2 Parathyroid hormone 2 SEQ ID NO: 11069 PTHLH Parathyroid hormone-like hormone SEQ ID
NOS: 11070-11078 PTK7 Protein tyrosine kinase 7 (inactive) SEQ ID NOS: 11079-11094 PTN
Pleiotrophin SEQ ID NOS: 11095-11096 PTPRA Protein tyrosine phosphatase, receptor type, A SEQ ID
NOS: 11097-11104 PTPRB Protein tyrosine phosphatase, receptor type, B SEQ ID NOS: 11105-11112
PTPRC Protein tyrosine phosphatase, receptor type, C SEQ ID NOS: 11113-11123 PTPRCAP Protein
tyrosine phosphatase, receptor type, C- SEQ ID NO: 11124 associated protein PTPRD Protein tyrosine
phosphatase, receptor type, D SEQ ID NOS: 11125-11136 PTPRF Protein tyrosine phosphatase, receptor
type, F SEQ ID NOS: 11137-11144 PTPRJ Protein tyrosine phosphatase, receptor type, J SEQ ID NOS:
11145-11150 PTPRO Protein tyrosine phosphatase, receptor type. O SEQ ID NOS: 11151-11159 PTPRS
Protein tyrosine phosphatase, receptor type, S SEQ ID NOS: 11160-11167 PTTG1IP Pituitary tumor-
transforming 1 interacting protein SEQ ID NOS: 11168-11171 PTX3 Pentraxin 3, long SEQ ID NO: 11172
PTX4 Pentraxin 4, long SEQ ID NOS: 11173-11175 PVR Poliovirus receptor SEQ ID NOS: 11176-11181
PXDN Peroxidasin SEQ ID NOS: 11182-11186 PXDNL Peroxidasin-like SEQ ID NOS: 11187-11189
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PXYLP1 2-phosphoxylose phosphatase 1 SEQ ID NOS: 11190-11202 PYY Peptide YY SEQ ID NOS:
11203-11204 PZP Pregnancy-zone protein SEQ ID NOS: 11205-11206 QPCT Glutaminyl-peplide
cyclotransferase SEQ ID NOS: 11207-11209 QPRT Quinolinate phosphoribosyltransferase SEQ ID NOS:
11210-11211 QRFP Pyroglutamylated RFamide peptide SEQ ID NOS: 11212-11213 QSOX1 Quiescin Q6
sulfhydryl oxidase 1 SEQ ID NOS: 11214-11217 R3HDML R3H domain containing-like SEQ ID NO:
11218 RAB26 RAB26, member RAS oncogene family SEQ ID NOS: 11219-11222 RAB36 RAB36,
member RAS oncogene family SEQ ID NOS: 11223-11225 RAB9B RAB9B, member RAS oncogene
family SEQ ID NO: 11226 RAET1E Retinoic acid early transcript 1E SEQ ID NOS: 11227-11232
RAET1G Retinoic acid early transcript 1G SEQ ID NOS: 11233-11235 RAMP2 Receptor (G protein-
coupled) activity modifying SEQ ID NOS: 11236-11240 protein 2 RAPGEF5 Rap guanine nucleotide
exchange factor (GEF) 5 SEQ ID NOS: 11241-11247 RARRES1 Retinoic acid receptor responder
(tazarotene SEQ ID NOS: 11248-11249 induced) 1 RARRES2 Retinoic acid receptor responder
(tazarotene SEQ ID NOS: 11250-11253 induced) 2 RASA2 RAS p21 protein activator 2 SEQ ID NOS:
11254-11256 RBM3 RNA binding motif (RNP1, RRM) protein 3 SEQ ID NOS: 11257-11259 RBP3
Retinol binding protein 3, interstitial SEQ ID NO: 11260 RBP4 Retinol binding protein 4, plasma SEQ ID
NOS: 11261-11264 RCN1 Reticulocalbin 1, EF-hand calcium binding domain SEQ ID NOS: 11265-11268
RCN2 Reticulocalbin 2, EF-hand calcium binding domain SEQ ID NOS: 11269-11272 RCN3
Reticulocalbin 3, EF-hand calcium binding domain SEQ ID NOS: 11273-11276 RCOR1 REST
corepressor 1 SEQ ID NOS: 11277-11278 RDH11 Retinol dehydrogenase 11 (all-trans/9-cis/11-cis) SEQ
ID NOS: 11279-11286 RDH12 Retinol dehydrogenase 12 (all-trans/9-cis/11-cis) SEO ID NOS: 11287-
11288 RDH13 Retinol dehydrogenase 13 (all-trans/9-cis) SEQ ID NOS: 11289-11297 RDH5 Retinol
dehydrogenase 5 (11-cis/9-cis) SEQ ID NOS: 11298-11302 RDH8 Retinol dehydrogenase 8 (all-trans)
SEQ ID NOS: 11303-11304 REGIA Regenerating islet-derived 1 alpha SEQ ID NO: 11305 REG1B
Regenerating islet-derived 1 beta SEQ ID NOS: 11306-11307 REG3A Regenerating islet-derived 3 alpha
SEQ ID NOS: 11308-11310 REG3G Regenerating islet-derived 3 gamma SEQ ID NOS: 11311-11313
REG4 Regenerating islet-derived family, member 4 SEQ ID NOS: 11314-11317 RELN Reelin SEQ ID
NOS: 11318-11321 RELT RELT tumor necrosis factor receptor SEQ ID NOS: 11322-11325 REN Renin
SEQ ID NOS: 11326-11327 REPIN1 Replication initiator 1 SEQ ID NOS: 11328-11341 REPS2 RALBP1
associated Eps domain containing 2 SEQ ID NOS: 11342-11343 RET Ret proto-oncogene SEQ ID NOS:
11344-11349 RETN Resistin SEQ ID NOS: 11350-11352 RETNLB Resistin like beta SEQ ID NO: 11353
RETSAT Retinol saturase (all-trans-retinol 13,14-reductase) SEQ ID NOS: 11354-11358 RFNG RFNG O-
fucosylpeptide 3-beta-N- SEQ ID NOS: 11359-11361 acetylglucosaminyltransferase RGCC Regulator of
cell cycle SEQ ID NO: 11362 RGL4 Ral guanine nucleotide dissociation stimulator-like SEQ ID NOS:
11363-11369 4 RGMA Repulsive guidance molecule family member a SEQ ID NOS: 11370-11379
RGMB Repulsive guidance molecule family member b SEQ ID NOS: 11380-11381 RHOQ Ras homolog
family member Q SEQ ID NOS: 11382-11386 RIC3 RIC3 acetylcholine receptor chaperone SEQ ID
NOS: 11387-11394 HRSP12 Heat-responsive protein 12 SEQ ID NOS: 11395-11398 RIMS1 Regulating
synaptic membrane exocytosis 1 SEQ ID NOS: 11399-11414 RIPPLY1 Ripply transcriptional repressor 1
SEQ ID NOS: 11415-11416 RLN1 Relaxin 1 SEQ ID NO: 11417 RLN2 Relaxin 2 SEQ ID NOS: 11418-
11419 RLN3 Relaxin 3 SEQ ID NOS: 11420-11421 RMDN1 Regulator of microtubule dynamics 1 SEQ
ID NOS: 11422-11435 RNASE1 Ribonuclease, RNase A family, 1 (pancreatic) SEQ ID NOS: 11436-
11440 RNASE10 Ribonuclease, RNase A family, 10 (non-active) SEQ ID NOS: 11441-11442 RNASE11
Ribonuclease, RNase A family, 11 (non-active) SEQ ID NOS: 11443-11453 RNASE12 Ribonuclease,
RNase A family, 12 (non-active) SEQ ID NO: 11454 RNASE13 Ribonuclease, RNase A family, 13 (non-
active) SEQ ID NO: 11455 RNASE2 Ribonuclease, RNase A family, 2 (liver, SEQ ID NO: 11456
eosinophil-derived neurotoxin) RNASE3 Ribonuclease, RNase A family, 3 SEQ ID NO: 11457 RNASE4
Ribonuclease, RNase A family, 4 SEQ ID NOS: 11458-11460 RNASE6 Ribonuclease, RNase A family, k6
SEQ ID NO: 11461 RNASE7 Ribonuclease, RNase A family, 7 SEQ ID NOS: 11462-11463 RNASE8
Ribonuclease, RNase A family, 8 SEQ ID NO: 11464 RNASE9 Ribonuclease, RNase A family, 9 (non-
active) SEQ ID NOS: 11465-11475 RNASEH1 Ribonuclease H1 SEQ ID NOS: 11476-11478 RNASET2
Ribonuclease T2 SEQ ID NOS: 11479-11486 RNF146 Ring finger protein 146 SEQ ID NOS: 11487-
11498 RNF148 Ring finger protein 148 SEQ ID NOS: 11499-11500 RNF150 Ring finger protein 150 SEQ
ID NOS: 11501-11505 RNF167 Ring finger protein 167 SEQ ID NOS: 11506-11516 RNF220 Ring finger
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protein 220 SEQ ID NOS: 11517-11523 RNF34 Ring finger protein 34, E3 ubiquitin protein ligase SEQ
ID NOS: 11524-11531 RNLS Renalase, FAD-dependent amine oxidase SEQ ID NOS: 11532-11534
RNPEP Arginyl aminopeptidase (aminopeptidase B) SEQ ID NOS: 11535-11540 ROR1 Receptor tyrosine
kinase-like orphan receptor 1 SEQ ID NOS: 11541-11543 RPL3 Ribosomal protein L3 SEQ ID NOS:
11544-11549 RPLP2 Ribosomal protein, large, P2 SEQ ID NOS: 11550-11552 RPN2 Ribophorin II SEQ
ID NOS: 11553-11559 RPS27L Ribosomal protein S27-like SEQ ID NOS: 11560-11565 RS1
Retinoschisin 1 SEQ ID NO: 11566 RSF1 Remodeling and spacing factor 1 SEQ ID NOS: 11567-11573
RSPO1 R-spondin 1 SEQ ID NOS: 11574-11577 RSPO2 R-spondin 2 SEQ ID NOS: 11578-11585 RSPO3
R-spondin 3 SEQ ID NOS: 11586-11587 RSPO4 R-spondin 4 SEQ ID NOS: 11588-11589 RSPRY1 Ring
finger and SPRY domain containing 1 SEQ ID NOS: 11590-11596 RTBDN Retbindin SEQ ID NOS:
11597-11609 RTN4RL1 Reticulon 4 receptor-like 1 SEQ ID NO: 11610 RTN4RL2 Reticulon 4 receptor-
like 2 SEQ ID NOS: 11611-11613 SAA1 Seram amyloid A1 SEQ ID NOS: 11614-11616 SAA2 Seram
amyloid A2 SEQ ID NOS: 11617-11622 SAA4 Serum amyloid A4, constitutive SEQ ID NO: 11623
SAP30 Sin3A-associated protein, 30 kDa SEQ ID NO: 11624 SAR1A Secretion associated, Ras related
GTPase 1A SEQ ID NOS: 11625-11631 SARAF Store-operated calcium entry-associated regulatory SEQ
ID NOS: 11632-11642 factor SARM1 Sterile alpha and TIR motif containing 1 SEQ ID NOS: 11643-
11646 SATB1 SATB homeobox 1 SEQ ID NOS: 11647-11659 SAXO2 Stabilizer of axonemal
microtubules 2 SEQ ID NOS: 11660-11664 SBSN Suprabasin SEQ ID NOS: 11665-11667 SBSPON
Somatomedin B and thrombospondin, type 1 SEQ ID NO: 11668 domain containing SCARF1 Scavenger
receptor class F, member 1 SEO ID NOS: 11669-11673 SCG2 Secretogranin II SEO ID NOS: 11674-
11676 SCG3 Secretogranin III SEQ ID NOS: 11677-11679 SCG5 Secretogranin V SEQ ID NOS: 11680-
11684 SCGB1A1 Secretoglobin, family 1A, member 1 (uteroglobin) SEQ ID NOS: 11685-11686
SCGB1C1 Secretoglobin, family 1C, member 1 SEQ ID NO: 11687 SCGB1C2 Secretoglobin, family 1C,
member 2 SEQ ID NO: 11688 SCGB1D1 Secretoglobin, family 1D, member 1 SEQ ID NO: 11689
SCGB1D2 Secretoglobin, family 1D, member 2 SEQ ID NO: 11690 SCGB1D4 Secretoglobin, family 1D,
member 4 SEQ ID NO: 11691 SCGB2A1 Secretoglobin, family 2A, member 1 SEQ ID NO: 11692
SCGB2A2 Secretoglobin, family 2A, member 2 SEQ ID NOS: 11693-11694 SCGB2B2 Secretoglobin,
family 2B, member 2 SEQ ID NOS: 11695-11696 SCGB3A1 Secretoglobin, family 3A, member 1 SEQ
ID NO: 11697 SCGB3A2 Secretoglobin, family 3A, member 2 SEQ ID NOS: 11698-11699 SCN1B
Sodium channel, voltage gated, type I beta subunit SEQ ID NOS: 11700-11705 SCN3B Sodium channel,
voltage gated, type III beta SEQ ID NOS: 11706-11710 subunit SCPEP1 Serine carboxypeptidase 1 SEQ
ID NOS: 11711-11718 SCRG1 Stimulator of chondrogenesis 1 SEQ ID NOS: 11719-11720 SCT Secretin
SEQ ID NO: 11721 SCUBE1 Signal peptide, CUB domain, EGF-like 1 SEQ ID NOS: 11722-11725
SCUBE2 Signal peptide, CUB domain, EGF-like 2 SEQ ID NOS: 11726-11732 SCUBE3 Signal peptide,
CUB domain, EGF-like 3 SEQ ID NO: 11733 SDC1 Syndecan 1 SEQ ID NOS: 11734-11738 SDF2
Stromal cell-derived factor 2 SEQ ID NOS: 11739-11741 SDF2L1 Stromal cell-derived factor 2-like 1
SEQ ID NO: 11742 SDF4 Stromal cell derived factor 4 SEQ ID NOS: 11743-11746 SDHAF2 Succinate
dehydrogenase complex assembly factor SEQ ID NOS: 11747-11754 2 SDHAF4 Succinate
dehydrogenase complex assembly factor SEQ ID NO: 11755 4 SDHB Succinate dehydrogenase complex,
subunit B, iron SEQ ID NOS: 11756-11758 sulfur (Ip) SDHD Succinate dehydrogenase complex, subunit
D, SEQ ID NOS: 11759-11768 integral membrane protein SEC14L3 SEC14-like lipid binding 3 SEQ ID
NOS: 11769-11775 SEC16A SEC16 homolog A, endoplasmic reticulum export SEQ ID NOS: 11776-
11782 factor SEC16B SEC16 homolog B, endoplasmic reticulum export SEQ ID NOS: 11783-11786
factor SEC22C SEC22 homolog C, vesicle trafficking protein SEQ ID NOS: 11787-11799 SEC31A
SEC31 homolog A, COP1I coat complex SEQ ID NOS: 11800-11829 component SECISBP2 SECIS
binding protein 2 SEQ ID NOS: 11830-11834 SECTM1 Secreted and transmembrane 1 SEQ ID NOS:
11835-11842 SEL1L Sel-1 suppressor of lin-12-like (C. elegans) SEQ ID NOS: 11843-11845 SEPT15 15
kDa selenoprotein SEQ ID NOS: 11846-11852 SELM Selenoprotein M SEQ ID NOS: 11853-11855
SEPN1 Selenoprotein N, 1 SEQ ID NOS: 11856-11859 SELO Selenoprotein O SEQ ID NOS: 11860-
11861 SEPP1 Selenoprotein P, plasma, 1 SEQ ID NOS: 11862-11867 SEMA3A Sema domain,
immunoglobulin domain (Ig), short SEQ ID NOS: 11868-11872 basic domain, secreted, (semaphorin) 3A
SEMA3B Sema domain, immunoglobulin domain (Ig), short SEQ ID NOS: 11873-11879 basic domain,
secreted, (semaphorin) 3B SEMA3C Sema domain, immunoglobulin domain (Ig), short SEQ ID NOS:
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11880-11884 basic domain, secreted, (semaphorin) 3C SEMA3E Sema domain, immunoglobulin domain
(Ig), short SEQ ID NOS: 11885-11889 basic domain, secreted, (semaphorin) 3E SEMA3F Sema domain,
immunoglobulin domain (Ig), short SEQ ID NOS: 11890-11896 basic domain, secreted, (semaphorin) 3F
SEMA3G Sema domain, immunoglobulin domain (Ig), short SEQ ID NOS: 11897-11899 basic domain,
secreted, (semaphorin) 3G SEMA4A Sema domain, immunoglobulin domain (Ig), SEQ ID NOS: 11900-
11908 transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 4A SEMA4B Sema
domain, immunoglobulin domain (Ig), SEQ ID NOS: 11909-11919 transmembrane domain (TM) and
short cytoplasmic domain, (semaphorin) 4B SEMA4C Sema domain, immunoglobulin domain (Ig), SEQ
ID NOS: 11920-11922 transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 4C
SEMA4D Sema domain, immunoglobulin domain (Ig), SEQ ID NOS: 11923-11936 transmembrane
domain (TM) and short cytoplasmic domain, (semaphorin) 4D SEMA4F Sema domain, immunoglobulin
domain (Ig), SEQ ID NOS: 11937-11945 transmembrane domain (TM) and short cytoplasmic domain,
(semaphorin) 4F SEMA4G Sema domain, immunoglobulin domain (Ig), SEQ ID NOS: 11946-11953
transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 4G SEMA5A Sema domain,
seven thrombospondin repeats (type SEQ ID NOS: 11954-11955 1 and type 1-like), transmembrane
domain (TM) and short cytoplasmic domain, (semaphorin) 5A SEMA6A Sema domain, transmembrane
domain (TM), and SEQ ID NOS: 11956-11963 cytoplasmic domain, (semaphorin) 6A SEMA6C Sema
domain, transmembrane domain (TM), and SEQ ID NOS: 11964-11969 cytoplasmic domain,
(semaphorin) 6C SEMA6D Sema domain, transmembrane domain (TM), and SEQ ID NOS: 11970-11983
cytoplasmic domain, (semaphorin) 6D SEMG1 Semenogelin I SEO ID NO: 11984 SEMG2 Semenogelin
II SEQ ID NO: 11985 SEPT9 Septin 9 SEQ ID NOS: 11986-12022 SERPINA1 Serpin peptidase inhibitor,
clade A (alpha-1 SEQ ID NOS: 12023-12039 antiproteinase, antitrypsin), member 1 SERPINA10 Setpin
peptidase inhibitor, clade A (alpha-1 SEQ ID NOS: 12040-12043 antiproteinase, antitrypsin), member 10
SERPINA11 Serpin peptidase inhibitor, clade A (alpha-1 SEQ ID NO: 12044 anti proteinase, antitrypsin),
member 11 SERPINA12 Serpin peptidase inhibitor, clade A (alpha-1 SEQ ID NOS: 12045-12046
antiproteinase, antitrypsin), member 12 SERPINA3 Serpin peptidase inhibitor, clade A (alpha-1 SEQ ID
NOS: 12047-12053 antiproteinase, antitrypsin), member 3 SERPINA4 Serpin peptidase inhibitor, clade A
(alpha-1 SEQ ID NOS: 12054-12056 antiproteinase, antitrypsin), member 4 SERPINA5 Serpin peptidase
inhibitor, clade A (alpha-1 SEQ ID NOS: 12057-12068 antiproteinase, antitrypsin), member 5 SERPINA6
Serpin peptidase inhibitor, clade A (alpha-1 SEQ ID NOS: 12069-12071 antiproteinase, antitrypsin),
member 6 SERPINA7 Setpin peptidase inhibitor, clade A (alpha-1 SEQ ID NOS: 12072-12073
antiproteinase, antitrypsin), member 7 SERPINA9 Serpin peptidase inhibitor, clade A (alpha-1 SEQ ID
NOS: 12074-12080 antiproteinase, antitrypsin), member 9 SERPINB2 Serpin peptidase inhibitor, clade B
(ovalbumin), SEQ ID NOS: 12081-12085 member 2 SERPINC1 Serpin peptidase inhibitor, clade C
(antithrombin), SEQ ID NOS: 12086-12087 member 1 SERPIND1 Serpin peptidase inhibitor, clade D
(heparin SEQ ID NOS: 12088-12089 cofactor), member 1 SERPINE1 Serpin peptidase inhibitor, clade E
(nexin, SEQ ID NO: 12090 plasminogen activator inhibitor type 1), member 1 SERPINE2 Serpin
peptidase inhibitor, clade E (nexin, SEQ ID NOS: 12091-12097 plasminogen activator inhibitor type 1),
member 2 SERPINE3 Serpin peptidase inhibitor, clade E (nexin, SEQ ID NOS: 12098-12101 plasminogen
activator inhibitor type 1), member 3 SERPINF1 Serpin peptidase inhibitor, clade F (alpha-2 SEQ ID
NOS: 12102-12110 antiplasmin, pigment epithelium derived factor), member 1 SERPINF2 Serpin
peptidase inhibitor, clade F (alpha-2 SEQ ID NOS: 12111-12115 antiplasmin, pigment epithelium derived
factor), member 2 SERPING1 Serpin peptidase inhibitor, clade G (C1 inhibitor), SEQ ID NOS: 12116-
12126 member 1 SERPINH1 Serpin peptidase inhibitor, clade H (heat shock SEQ ID NOS: 12127-12141
protein 47), member 1, (collagen binding protein 1) SERPINI1 Serpin peptidase inhibitor, clade I
(neuroserpin), SEQ ID NOS: 12142-12146 member 1 SERPINI2 Serpin peptidase inhibitor, clade I
(panepin), SEQ ID NOS: 12147-12153 member 2 SEZ6L2 Seizure related 6 homolog (mouse)-like 2 SEQ
ID NOS: 12154-12160 SFRP1 Secreted frizzled-related protein 1 SEQ ID NOS: 12161-12162 SFRP2
Secreted frizzled-related protein 2 SEQ ID NO: 12163 SFRP4 Secreted frizzled-related protein 4 SEQ ID
NOS: 12164-12165 SFRP5 Secreted frizzled-related protein 5 SEQ ID NO: 12166 SFTA2 Surfactant
associated 2 SEQ ID NOS: 12167-12168 SFTPA1 Surfactant protein A1 SEQ ID NOS: 12169-12173
SFTPA2 Surfactant protein A2 SEQ ID NOS: 12174-12178 SFTPB Surfactant protein B SEQ ID NOS:
12179-12183 SFTPD Surfactant protein D SEQ ID NOS: 12184-12185 SFXN5 Sideroflexin 5 SEQ ID
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NOS: 12186-12190 SGCA Sarcoglycan, alpha (50 kDa dystrophin-associated SEQ ID NOS: 12191-12198
glycoprotein) SGSH N-sulfoglucosamine sulfohydrolase SEQ ID NOS: 12199-12207 SH3RF3 SH3
domain containing ring finger 3 SEQ ID NO: 12208 SHBG Sex hormone-binding globulin SEQ ID NOS:
12209-12227 SHE Src homology 2 domain containing E SEQ ID NOS: 12228-12230 SHH Sonic
hedgehog SEQ ID NOS: 12231-12234 SHKBP1 SH3KBP1 binding protein 1 SEQ ID NOS: 12235-12250
SIAE Sialic acid acetylesterase SEQ ID NOS: 12251-12253 SIDT2 SID1 transmembrane family, member
2 SEQ ID NOS: 12254-12263 SIGLEC10 Sialic acid binding Ig-like lectin 10 SEQ ID NOS: 12264-12272
SIGLEC6 Sialic acid binding Ig-like lectin 6 SEQ ID NOS: 12273-12278 SIGLEC7 Sialic acid binding
Ig-like lectin 7 SEQ ID NOS: 12279-12283 SIGLECL1 SIGLEC family like 1 SEQ ID NOS: 12284-
12289 SIGMAR1 Sigma non-opioid intracellular receptor 1 SEQ ID NOS: 12290-12293 SIL1 SIL1
nucleotide exchange factor SEQ ID NOS: 12294-12302 SIRPB1 Signal-regulatory protein beta 1 SEQ ID
NOS: 12303-12315 SIRPD Signal-regulatory protein delta SEQ ID NOS: 12316-12318 SLAMF1
Signaling lymphocytic activation molecule family SEQ ID NOS: 12319-12321 member 1 SLAMF7
SLAM family member 7 SEQ ID NOS: 12322-12330 SLC10A3 Solute carrier family 10, member 3 SEQ
ID NOS: 12331-12335 SLC15A3 Solute carrier family 15 (oligopeptide transporter), SEQ ID NOS:
12336-12341 member 3 SLC25A14 Solute carrier family 25 (mitochondrial carrier, SEQ ID NOS: 12342-
12348 brain), member 14 SLC25A25 Solute carrier family 25 (mitochondrial carrier; SEQ ID NOS:
12349-12355 phosphate carrier), member 25 SLC2A5 Solute carrier family 2 (facilitated glucose/fructose
SEQ ID NOS: 12356-12364 transporter), member 5 SLC35E3 Solute carrier family 35, member E3 SEQ
ID NOS: 12365-12366 SLC39A10 Solute carrier family 39 (zinc transporter), member SEQ ID NOS:
12367-12373 10 SLC39A14 Solute carrier family 39 (zinc transporter), member SEQ ID NOS: 12374-
12384 14 SLC39A4 Solute carrier family 39 (zinc transporter), member SEQ ID NOS: 12385-12387 4
SLC39A5 Solute carrier family 39 (zinc transporter), member SEQ ID NOS: 12388-12394 5 SLC3A1
Solute carrier family 3 (amino acid transporter SEQ ID NOS: 12395-12404 heavy chain), member 1
SLC51A Solute carrier family 51, alpha subunit SEQ ID NOS: 12405-12409 SLC52A2 Solute carrier
family 52 (riboflavin transporter), SEQ ID NOS: 12410-12420 member 2 SLC5A6 Solute carrier family 5
(sodium/multivitamin and SEQ ID NOS: 12421-12431 iodide cotransporter), member 6 SLC6A9 Solute
carrier family 6 (neurotransmitter SEQ ID NOS: 12432-12439 transporter, glycine), member 9 SLC8A1
Solute carrier family 8 (sodium/calcium SEQ ID NOS: 12440-12451 exchanger), member 1 SLC8B1
Solute carrier family 8 (sodium/lithium/calcium SEQ ID NOS: 12452-12462 exchanger), member B1
SLC9A6 Solute carrier family 9, subfamily A (NHE6, cation SEQ ID NOS: 12463-12474 proton
antiporter 6), member 6 SLCO1A2 Solute carrier organic anion transporter family, SEQ ID NOS: 12475-
12488 member 1A2 SLIT1 Slit guidance ligand 1 SEQ ID NOS: 12489-12492 SLIT2 Slit guidance ligand
2 SEQ ID NOS: 12493-12501 SLIT3 Slit guidance ligand 3 SEQ ID NOS: 12502-12504 SLITRK3 SLIT
and NTRK-like family, member 3 SEQ ID NOS: 12505-12507 SLPI Secretory leukocyte peptidase
inhibitor SEQ ID NO: 12508 SLTM SAFB-like, transcription modulator SEQ ID NOS: 12509-12522
SLURP1 Secreted LY6/PLAUR domain containing 1 SEQ ID NO: 12523 SMARCA2 SWI/SNF related,
matrix associated, actin SEQ ID NOS: 12524-12571 dependent regulator of chromatin, subfamily a,
member 2 SMG6 SMG6 nonsense mediated mRNA decay factor SEQ ID NOS: 12572-12583 SMIM7
Small integral membrane protein 7 SEQ ID NOS: 12584-12600 SMOC1 SPARC related modular calcium
binding 1 SEQ ID NOS: 12601-12602 SMOC2 SPARC related modular calcium binding 2 SEQ ID NOS:
12603-12607 SMPDL3A Sphingomyelin phosphodiesterase, acid-like 3A SEQ ID NOS: 12608-12609
SMPDL3B Sphingomyelin phosphodiesterase, acid-like 3B SEQ ID NOS: 12610-12614 SMR3A
Submaxillary gland androgen regulated protein 3A SEQ ID NO: 12615 SMR3B Submaxillary gland
androgen regulated protein 3B SEQ ID NOS: 12616-12618 SNED1 Sushi, nidogen and EGF-like domains
1 SEQ ID NOS: 12619-12625 SNTB1 Syntrophin, beta 1 (dystrophin-associated protein SEQ ID NOS:
12626-12628 A1, 59 kDa, basic component 1) SNTB2 Syntrophin, beta 2 (dystrophin-associated protein
SEQ ID NOS: 12629-12633 A1, 59 kDa, basic component 2) SNX14 Sorting nexin 14 SEQ ID NOS:
12634-12645 SOD3 Superoxide dismutase 3, extracellular SEQ ID NOS: 12646-12647 SOST Sclerostin
SEQ ID NO: 12648 SOSTDC1 Sclerostin domain containing 1 SEQ ID NOS: 12649-12650 SOWAHA
Sosondowah ankyrin repeat domain family SEQ ID NO: 12651 member A SPACA3 Sperm acrosome
associated 3 SEQ ID NOS: 12652-12654 SPACA4 Sperm acrosome associated 4 SEQ ID NO: 12655
SPACA5 Sperm acrosome associated 5 SEQ ID NOS: 12656-12657 SPACA5B Sperm acrosome
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associated 5B SEQ ID NO: 12658 SPACA7 Sperm acrosome associated 7 SEQ ID NOS: 12659-12662
SPAG11A Sperm associated antigen 11A SEQ ID NOS: 12663-12671 SPAG11B Sperm associated antigen
11B SEQ ID NOS: 12672-12680 SPARC Secreted protein, acidic, cystcinc-rich (osteonectin) SEQ ID
NOS: 12681-12685 SPARCL1 SPARC-like 1 (hevin) SEQ ID NOS: 12686-12695 SPATA20
Spermatogenesis associated 20 SEQ ID NOS: 12696-12709 SPESP1 Sperm equatorial segment protein 1
SEQ ID NO: 12710 SPINK1 Serine peptidase inhibitor, Kazal type 1 SEQ ID NOS: 12711-12712
SPINK13 Serine peptidase inhibitor, Kazal type 13 (putative) SEQ ID NOS: 12713-12715 SPINK14
Serine peptidase inhibitor, Kazal type 14 (putative) SEQ ID NOS: 12716-12717 SPINK2 Serine peptidase
inhibitor, Kazal type 2 (acrosin- SEQ ID NOS: 12718-12723 trypsin inhibitor) SPINK4 Serine peptidase
inhibitor, Kazal type 4 SEQ ID NOS: 12724-12725 SPINK5 Serine peptidase inhibitor, Kazal type 5 SEQ
ID NOS: 12726-12731 SPINK6 Serine peptidase inhibitor, Kazal type 6 SEQ ID NOS: 12732-12734
SPINK7 Serine peptidase inhibitor, Kazal type 7 (putative) SEQ ID NOS: 12735-12736 SPINK8 Serine
peptidase inhibitor, Kazal type 8 (putative) SEQ ID NO: 12737 SPINK9 Serine peptidase inhibitor, Kazal
type 9 SEQ ID NOS: 12738-12739 SPINT1 Serine peptidase inhibitor, Kunitz type 1 SEQ ID NOS:
12740-12747 SPINT2 Serine peptidase inhibitor, Kunitz type, 2 SEQ ID NOS: 12748-12755 SPINT3
Serine peptidase inhibitor, Kunitz type, 3 SEQ ID NO: 12756 SPINT4 Serine peptidase inhibitor, Kunitz
type 4 SEQ ID NO: 12757 SPOCK1 Sparc/osteonectin, cwcv and kazal-like domains SEQ ID NOS:
12758-12761 proteoglycan (testican) 1 SPOCK2 Sparc/osteonectin, cwcv and kazal-like domains SEQ ID
NOS: 12762-12765 proteoglycan (testican) 2 SPOCK3 Sparc/osteonectin, cwcv and kazal-like domains
SEQ ID NOS: 12766-12791 proteoglycan (testican) 3 SPON1 Spondin 1, extracellular matrix protein SEQ
ID NO: 12792 SPON2 Spondin 2, extracellular matrix protein SEQ ID NOS: 12793-12802 SPP1 Secreted
phosphoprotein 1 SEQ ID NOS: 12803-12807 SPP2 Secreted phosphoprotein 2, 24 kDa SEQ ID NOS:
12808-12810 SPRN Shadow of prion protein homolog (zebrafish) SEQ ID NO: 12811 SPRYD3 SPRY
domain containing 3 SEQ ID NOS: 12812-12815 SPRYD4 SPRY domain containing 4 SEQ ID NO:
12816 SPTY2D1- SPTY2D1 antisense RNA 1 SEQ ID NOS: 12817-12822 AS1 SPX Spexin hormone
SEQ ID NOS: 12823-12824 SRGN Serglycin SEQ ID NO: 12825 SRL Sarcalumenin SEQ ID NOS:
12826-12828 SRP14 Signal recognition particle 14 kDa (homologous SEQ ID NOS: 12829-12832 Alu
RNA binding protein) SRPX Sushi-repeat containing protein, X-linked SEQ ID NOS: 12833-12836
SRPX2 Sushi-repeat containing protein, X-linked 2 SEQ ID NOS: 12837-12840 SSC4D Scavenger
receptor cysteine rich family, 4 domains SEQ ID NO: 12841 SSC5D Scavenger receptor cysteine rich
family, 5 domains SEQ ID NOS: 12842-12845 SSPO SCO-spondin SEQ ID NO: 12846 SSR2 Signal
sequence receptor, beta (translocon- SEQ ID NOS: 12847-12856 associated protein beta) SST
Somatostatin SEQ ID NO: 12857 ST3GAL1 ST3 beta-galactoside alpha-2,3-sialyltransferase 1 SEQ ID
NOS: 12858-12865 ST3GAL4 ST3 beta-galactoside alpha-2,3-sialyltransferase 4 SEQ ID NOS: 12866-
12881 ST6GAL1 ST6 beta-galactosamide alpha-2,6-sialyltranferase 1 SEQ ID NOS: 12882-12897
ST6GALNAC2 ST6 (alpha-N-acetyl-neuraminyl-2,3-beta- SEQ ID NOS: 12898-12902 galactosvl-1,3)-N-
acetylgalactosaminide alpha-2,6- sialyltransferase 2 ST6GALNAC5 ST6 (alpha-N-acetyl-neuraminyl-2,3-
beta- SEQ ID NOS: 12903-12904 galactosyl-1,3)-N-acetylgalactosaminide alpha-2,6- sialyltransferase 5
ST6GALNAC6 ST6 (alpha-N-acetyl-neuraminyl-2,3-beta- SEQ ID NOS: 12905-12912 galactosyl-1,3)-N-
acelylgalaclosaminide alpha-2,6- sialyltransferase 6 ST8SIA2 ST8 alpha-N-acetyl-neuraminide alpha-2,8-
SEQ ID NOS: 12913-12915 sialyltransferase 2 ST8SIA4 ST8 alpha-N-acetyl-neuraminide alpha-2,8- SEQ
ID NOS: 12916-12918 sialyltransferase 4 ST8SIA6 ST8 alpha-N-acetyl-neuraminide alpha-2,8- SEQ ID
NOS: 12919-12920 sialyltransferase 6 STARD7 StAR-related lipid transfer (START) domain SEQ ID
NOS: 12921-12922 containing 7 STATH Statherin SEQ ID NOS: 12923-12925 STC1 Stanniocalcin 1
SEQ ID NOS: 12926-12927 STC2 Stanniocalcin 2 SEQ ID NOS: 12928-12930 STMND1 Stathmin
domain containing 1 SEQ ID NOS: 12931-12932 C7orf73 Chromosome 7 open reading frame 73 SEQ ID
NOS: 12933-12934 STOML2 Stomatin (EPB72)-like 2 SEQ ID NOS: 12935-12938 STOX1 Storkhead
box 1 SEQ ID NOS: 12939-12943 STRC Stereocilin SEQ ID NOS: 12944-12949 SUCLG1 Succinate-
CoA ligase, alpha subunit SEQ ID NOS: 12950-12951 SUDS3 SDS3 homolog, SIN3A corepressor
complex SEQ ID NO: 12952 component SULF1 Sulfatase 1 SEQ ID NOS: 12953-12963 SULF2
Sulfatase 2 SEQ ID NOS: 12964-12968 SUMF1 Sulfatase modifying factor 1 SEQ ID NOS: 12969-12973
SUMF2 Sulfatase modifying factor 2 SEQ ID NOS: 12974-12987 SUSD1 Sushi domain containing 1
SEQ ID NOS: 12988-12993 SUSD5 Sushi domain containing 5 SEQ ID NOS: 12994-12995 SVEP1
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Sushi, von Willebrand factor type A, EGF and SEQ ID NOS: 12996-12998 pentraxin domain containing 1
SWSAP1 SWIM-type zinc finger 7 associated protein 1 SEQ ID NO: 12999 SYAP1 Synapse associated
protein 1 SEQ ID NO: 13000 SYCN Syncoilin SEQ ID NO: 13001 TAC1 Tachykinin, precursor 1 SEQ ID
NOS: 13002-13004 TAC3 Tachykinin 3 SEQ ID NOS: 13005-13014 TAC4 Tachykinin 4 (hemokinin)
SEQ ID NOS: 13015-13020 TAGLN2 Transgelin 2 SEQ ID NOS: 13021-13024 TAPBP TAP binding
protein (tapasin) SEQ ID NOS: 13025-13030 TAPBPL TAP binding protein-like SEQ ID NOS: 13031-
13032 TBL2 Transducin (beta)-like 2 SEQ ID NOS: 13033-13045 TBX10 T-box 10 SEQ ID NO: 13046
TCF12 Transcription factor 12 SEQ ID NOS: 13047-13060 TCN1 Transcobalamin I (vitamin B12,
binding protein, R SEQ ID NO: 13061 binder family) TCN2 Transcobalamin II SEQ ID NOS: 13062-
13065 TCTN1 Tectonic family member 1 SEQ ID NOS: 13066-13084 TCTN3 Tectonic family member 3
SEQ ID NOS: 13085-13089 TDP2 Tyrosyl-DNA phosphodiesterase 2 SEQ ID NOS: 13090-13091
C14orf80 Chromosome 14 open reading frame 80 SEQ ID NOS: 13092-13105 TEK TEK tyrosine kinase,
endothelial SEQ ID NOS: 13106-13110 TEPP Testis, prostate and placenta expressed SEQ ID NOS:
13111-13112 TEX101 Testis expressed 101 SEQ ID NOS: 13113-13114 TEX264 Testis expressed 264
SEQ ID NOS: 13115-13126 C1orf234 Chromosome 1 open reading frame 234 SEQ ID NOS: 13127-
13129 TF Transferrin SEQ ID NOS: 13130-13136 TFAM Transcription factor A, mitochondrial SEQ ID
NOS: 13137-13139 TFF1 Trefoil factor 1 SEQ ID NO: 13140 TFF2 Trefoil factor 2 SEQ ID NO: 13141
TFF3 Trefoil factor 3 (intestinal) SEQ ID NOS: 13142-13144 TFPI Tissue factor pathway inhibitor
(lipoprotein- SEQ ID NOS: 13145-13154 associated coagulation inhibitor) TFPI2 Tissue factor pathway
inhibitor 2 SEQ ID NOS: 13155-13156 TG Thyroglobulin SEQ ID NOS: 13157-13166 TGFB1
Transforming growth factor, beta 1 SEQ ID NOS: 13167-13168 TGFB2 Transforming growth factor, beta
2 SEQ ID NOS: 13169-13170 TGFB3 Transforming growth factor, beta 3 SEQ ID NOS: 13171-13172
TGFBI Transforming growth factor, beta-induced, 68 kDa SEQ ID NOS: 13173-13180 TGFBR1
Transforming growth factor, beta receptor III SEQ ID NOS: 13181-13190 TGFBR3 Transforming growth
factor, beta receptor III SEQ ID NOS: 13191-13197 THBS1 Thrombospondin 1 SEQ ED NOS: 13198-
13199 THBS2 Thrombospondin 2 SEQ ID NOS: 13200-13202 THBS3 Thrombospondin 3 SEQ ID NOS:
13203-13207 THBS4 Thrombospondin 4 SEQ ID NOS: 13208-13209 THOC3 THO complex 3 SEQ ID
NOS: 13210-13219 THPO Thrombopoietin SEQ ID NOS: 13220-13225 THSD4 Thrombospondin, type I,
domain containing 4 SEQ ID NOS: 13226-13229 THY1 Thy-1 cell surface antigen SEQ ID NOS: 13230-
13235 TIE1 Tyrosine kinase with immunoglobulin-like and SEQ ID NOS: 13236-13237 EGF-like
domains 1 TIMMDC1 Translocase of inner mitochondrial membrane SEQ ID NOS: 13238-13245 domain
containing 1 TIMP1 TIMP metallopeptidase inhibitor 1 SEQ ID NOS: 13246-13250 TIMP2 TIMP
metallopeptidase inhibitor 2 SEQ ID NOS: 13251-13255 TIMP3 TIMP metallopeptidase inhibitor 3 SEQ
ID NO: 13256 TIMP4 TIMP metallopeptidase inhibitor 4 SEQ ID NO: 13257 TINAGL1 Tubulointerstitial
nephritis antigen-like 1 SEQ ID NOS: 13258-13260 TINF2 TERF1 (TRF1)-interacting nuclear factor 2
SEQ ID NOS: 13261-13270 TLL2 Tolloid-like 2 SEQ ID NO: 13271 TLR1 Toll-like receptor 1 SEQ ID
NOS: 13272-13277 TLR3 Toll-like receptor 3 SEQ ID NOS: 13278-13280 TM2D2 TM2 domain
containing 2 SEQ ID NOS: 13281-13286 TM2D3 TM2 domain containing 3 SEQ ID NOS: 13287-13294
TM7SF3 Transmembrane 7 superfamily member 3 SEQ ID NOS: 13295-13309 TM9SF1 Transmembrane
9 superfamily member 1 SEQ ID NOS: 13310-13320 TMCO6 Transmembrane and coiled-coil domains 6
SEQ ID NOS: 13321-13328 TMED1 Transmembrane p24 trafficking protein 1 SEQ ID NOS: 13329-
13335 TMED2 Transmembrane p24 trafficking protein 2 SEQ ID NOS: 13336-13338 TMED3
Transmembrane p24 trafficking protein 3 SEQ ID NOS: 13339-13342 TMED4 Transmembrane p24
trafficking protein 4 SEQ ID NOS: 13343-13345 TMED5 Transmembrane p24 trafficking protein 5 SEQ
ID NOS: 13346-13349 TMED7 Transmembrane p24 trafficking protein 7 SEQ ID NOS: 13350-13351
TMED7-TMED7-TICAM2 readthrough SEQ ID NOS: 13352-13353 TICAM2 TMEM108
Transmembrane protein 108 SEQ ID NOS: 13354-13362 TMEM116 Transmembrane protein 116 SEQ ID
NOS: 13363-13374 TMEM119 Transmembrane protein 119 SEQ ID NOS: 13375-13378 TMEM155
Transmembrane protein 155 SEQ ID NOS: 13379-13382 TMEM168 Transmembrane protein 168 SEQ ID
NOS: 13383-13388 TMEM178A Transmembrane protein 178A SEQ ID NOS: 13389-13390 TMEM179
Transmembrane protein 179 SEQ ID NOS: 13391-13396 TMEM196 Transmembrane protein 196 SEQ ID
NOS: 13397-13401 TMEM199 Transmembrane protein 199 SEQ ID NOS: 13402-13405 TMEM205
Transmembrane protein 205 SEQ ID NOS: 13406-13419 TMEM213 Transmembrane protein 213 SEQ ID
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NOS: 13420-13423 TMEM25 Transmembrane protein 25 SEQ ID NOS: 13424-13440 TMEM30C
Transmembrane protein 30C SEQ ID NO: 13441 TMEM38B Transmembrane protein 38B SEQ ID NOS:
13442-13446 TMEM44 Transmembrane protein 44 SEQ ID NOS: 13447-13456 TMEM52
Transmembrane protein 52 SEQ ID NOS: 13457-13461 TMEM52B Transmembrane protein 52B SEQ ID
NOS: 13462-13464 TMEM59 Transmembrane protein 59 SEQ ID NOS: 13465-13472 TMEM67
Transmembrane protein 67 SEQ ID NOS: 13473-13484 TMEM70 Transmembrane protein 70 SEQ ID
NOS: 13485-13487 TMEM87A Transmembrane protein 87A SEQ ID NOS: 13488-13497 TMEM94
Transmembrane protein 94 SEQ ID NOS: 13498-13513 TMEM95 Transmembrane protein 95 SEQ ID
NOS: 13514-13516 TMIGD1 Transmembrane and immunoglobulin domain SEQ ID NOS: 13517-13518
containing 1 TMPRSS12 Transmembrane (C-terminal) protease, serine 12 SEQ ID NOS: 13519-13520
TMPRSS5 Transmembrane protease, serine 5 SEQ ID NOS: 13521-13532 TMUB1 Transmembrane and
ubiquitin-like domain SEQ ID NOS: 13533-13539 containing 1 TMX2 Thioredoxin-related
transmembrane protein 2 SEQ ID NOS: 13540-13547 TMX3 Thioredoxin-related transmembrane protein
3 SEQ ID NOS: 13548-13555 TNC Tenascin C SEQ ID NOS: 13556-13564 TNFAIP6 Tumor necrosis
factor, alpha-induced protein 6 SEQ ID NO: 13565 TNFRSF11A Tumor necrosis factor receptor
superfamily, SEQ ID NOS: 13566-13570 member 11a, NFKB activator TNFRSF11B Tumor necrosis
factor receptor superfamily, SEQ ID NOS: 13571-13572 member 11b TNFRSF12A Tumor necrosis factor
receptor superfamily, SEQ ID NOS: 13573-13578 member 12A TNFRSF14 Tumor necrosis factor
receptor superfamily, SEQ ID NOS: 13579-13585 member 14 TNFRSF18 Tumor necrosis factor receptor
superfamily, SEQ ID NOS: 13586-13589 member 18 TNFRSF1A Tumor necrosis factor receptor
superfamily, SEQ ID NOS: 13590-13598 member 1A TNFRSF1B Tumor necrosis factor receptor
superfamily, SEQ ID NOS: 13599-13600 member 1B TNFRSF25 Tumor necrosis factor receptor
superfamily, SEQ ID NOS: 13601-13612 member 25 TNFRSF6B Tumor necrosis factor receptor
superfamily, SEQ ID NO: 13613 member 6b, decoy TNFSF11 Tumor necrosis factor (ligand) superfamily,
SEQ ID NOS: 13614-13618 member 11 TNFSF12 Tumor necrosis factor (ligand) superfamily, SEQ ID
NOS: 13619-13620 member 12, TNFSF12-TNFSF13 readthrough SEQ ID NO: 13621
TNFSF13 TNFSF15 Tumor necrosis factor (ligand) superfamily, SEQ ID NOS: 13622-13623 member 15
TNN Tenascin N SEQ ID NOS: 13624-13626 TNR Tenascin R SEQ ID NOS: 13627-13629 TNXB
Tenascin XB SEQ ID NOS: 13630-13636 FAM179B Family with sequence similarity 179, member B
SEQ ID NOS: 13637-13642 TOMM7 Translocase of outer mitochondrial membrane 7 SEQ ID NOS:
13643-13646 homolog (yeast) TOP1MT Topoisomerase (DMA) I, mitochondrial SEQ ID NOS: 13647-
13661 TOR1A Torsin family 1, member A (torsin A) SEQ ID NO: 13662 TOR1B Torsin family 1, member
B (torsin B) SEQ ID NOS: 13663-13664 TOR2A Torsin family 2, member A SEQ ID NOS: 13665-13671
TOR3A Torsin family 3, member A SEQ ID NOS: 13672-13676 TPD52 Tumor protein D52 SEQ ID NOS:
13677-13689 TPO Thyroid peroxidase SEQ ID NOS: 13690-13700 TPP1 Tripeptidyl peptidase I SEQ ID
NOS: 13701-13718 TPSAB1 Tryptase alpha/beta 1 SEQ ID NOS: 13719-13721 TPSB2 Tryptase beta 2
(gene/pseudogene) SEQ ID NOS: 13722-13724 TPSD1 Tryptase delta 1 SEQ ID NOS: 13725-13726
TPST1 Tyrosylprotein sulfotransferase 1 SEQ ID NOS: 13727-13729 TPST2 Tyrosylprotein
sulfotransferase 2 SEQ ID NOS: 13730-13738 TRABD2A TraB domain containing 2A SEQ ID NOS:
13739-13741 TRABD2B TraB domain containing 2B SEQ ID NO: 13742 TREH Trehalase (brush-border
membrane glycoprotein) SEQ ID NOS: 13743-13745 TREM1 Triggering receptor expressed on myeloid
cells 1 SEQ ID NOS: 13746-13749 TREM2 Triggering receptor expressed on myeloid cells 2 SEQ ID
NOS: 13750-13752 TRH Thyrotropin-releasing hormone SEQ ID NOS: 13753-13754 TRIM24 Tripartite
motif containing 24 SEQ ID NOS: 13755-13756 TRIM28 Tripartite motif containing 28 SEQ ID NOS:
13757-13762 TRIO Trio Rho guanine nucleotide exchange factor SEQ ID NOS: 13763-13769 TRNP1
TMF1-regulated nuclear protein 1 SEQ ID NOS: 13770-13771 TSC22D4 TSC22 domain family, member
4 SEQ ID NOS: 13772-13775 TSHB Thyroid stimulating hormone, beta SEQ ID NOS: 13776-13777
TSHR Thyroid stimulating hormone receptor SEQ ID NOS: 13778-13785 TSKU Tsukushi, small leucine
rich proteoglycan SEQ ID NOS: 13786-13790 TSLP Thymic stromal lymphopoietin SEQ ID NOS:
13791-13793 TSPAN3 Tetraspanin 3 SEQ ID NOS: 13794-13799 TSPAN31 Tetraspanin 31 SEQ ID NOS:
13800-13806 TSPEAR Thrombospondin-type laminin G domain and EAR SEQ ID NOS: 13807-13810
repeats TTC13 Tetratricopeptide repeat domain 13 SEQ ID NOS: 13811-13817 TTC19 Tetratricopeptide
repeat domain 19 SEQ ID NOS: 13818-13823 TTC9B Tetratricopeptide repeat domain 9B SEQ ID NO:
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13824 TTLL11 Tubulin tyrosine ligase-like family member 11 SEQ ID NOS: 13825-13829 TTR
Transthyretin SEQ ID NOS: 13830-13832 TWSG1 Twisted gastrulation BMP signaling modulator 1 SEQ
ID NOS: 13833-13835 TXNDC12 Thioredoxin domain containing 12 (endoplasmic SEQ ID NOS: 13836-
13838 reticulum) TXNDC15 Thioredoxin domain containing 15 SEQ ID NOS: 13839-13845 TXNDC5
Thioredoxin domain containing 5 (endoplasmic SEQ ID NOS: 13846-13847 reticulum) TXNRD2
Thioredoxin reductase 2 SEQ ID NOS: 13848-13860 TYRP1 Tyrosinase-related protein 1 SEQ ID NOS:
13861-13863 UBAC2 UBA domain containing 2 SEQ ID NOS: 13864-13868 UBALD1 UBA-like
domain containing 1 SEQ ID NOS: 13869-13877 UBAP2 Ubiquitin associated protein 2 SEQ ID NOS:
13878-13884 UBXN8 UBX domain protein 8 SEQ ID NOS: 13885-13891 UCMA Upper zone of growth
plate and cartilage matrix SEO ID NOS: 13892-13893 associated UCN Urocortin SEO ID NO: 13894
UCN2 Urocortin 2 SEQ ID NO: 13895 UCN3 Urocortin 3 SEQ ID NO: 13896 UGGT2 UDP-glucose
glycoprotein glucosyltransferase 2 SEQ ID NOS: 13897-13902 UGT1A10 UDP glucuronosyltransferase 1
family, polypeptide SEQ ID NOS: 13903-13904 A10 UGT2A1 UDP glucuronosyltransferase 2 family,
polypeptide SEQ ID NOS: 13905-13909 A1, complex locus UGT2B11 UDP glucuronosyltransferase 2
family, polypeptide SEQ ID NO: 13910 B11 UGT2B28 UDP glucuronosyltransferase 2 family,
polypeptide SEQ ID NOS: 13911-13912 B28 UGT2B4 UDP glucuronosyltransferase 2 family,
polypeptide SEQ ID NOS: 13913-13916 B4 UGT2B7 UDP glucuronosyltransferase 2 family, polypeptide
SEQ ID NOS: 13917-13920 B7 UGT3A1 UDP glycosyltransferase 3 family, polypeptide A1 SEQ ID
NOS: 13921-13926 UGT3A2 UDP glycosyltransferase 3 family, polypeptide A2 SEQ ID NOS: 13927-
13930 UGT8 UDP glycosyltransfcrasc 8 SEO ID NOS: 13931-13933 ULBP3 UL16 binding protein 3
SEQ ID NOS: 13934-13935 UMOD Uromodulin SEQ ID NOS: 13936-13947 UNC5C Unc-5 netrin
receptor C SEQ ID NOS: 13948-13952 UPK3B Uroplakin 3B SEQ ID NOS: 13953-13955 USP11
Ubiquitin specific peptidase 11 SEQ ID NOS: 13956-13959 USP14 Ubiquitin specific peptidase 14
(tRNA-guanine SEQ ID NOS: 13960-13966 transglycosylase) USP3 Ubiquitin specific peptidase 3 SEQ
ID NOS: 13967-13982 CIRH1A Cirrhosis, autosomal recessive 1A (cirhin) SEQ ID NOS: 13983-13992
UTS2 Urotensin 2 SEQ ID NOS: 13993-13995 UTS2B Urotensin 2B SEQ ID NOS: 13996-14001 UTY
Ubiquitously transcribed tetratricopeptide repeat SEQ ID NOS: 14002-14014 containing. Y-linked UXS1
UDP-glucuronate decarboxylase 1 SEQ ID NOS: 14015-14022 VASH1 Vasohibin 1 SEQ ID NOS: 14023-
14025 VCAN Versican SEQ ID NOS: 14026-14032 VEGFA Vascular endothelial growth factor A SEQ ID
NOS: 14033-14058 VEGFB Vascular endothelial growth factor B SEQ ID NOS: 14059-14061 VEGFC
Vascular endothelial growth factor C SEQ ID NO: 14062 FIGF C-fos induced growth factor (vascular
endothelial SEQ ID NO: 14063 growth factor D) VGF VGF nerve growth factor inducible SEQ ID NOS:
14064-14066 VIP Vasoactive intestinal peptide SEQ ID NOS: 14067-14069 VIPR2 Vasoactive intestinal
peptide receptor 2 SEQ ID NOS: 14070-14073 VIT Vitrin SEQ ID NOS: 14074-14081 VKORC1 Vitamin
K epoxide reductase complex, subunit 1 SEQ ID NOS: 14082-14089 VLDLR Very low density lipoprotein
receptor SEQ ID NOS: 14090-14092 VMO1 Vitelline membrane outer layer 1 homolog SEQ ID NOS:
14093-14096 (chicken) VNN1 Vanin 1 SEQ ID NO: 14097 VNN2 Vanin 2 SEQ ID NOS: 14098-14111
VNN3 Vanin 3 SEQ ID NOS: 14112-14123 VOPP1 Vesicular, overexpressed in cancer, prosurvival SEQ
ID NOS: 14124-14136 protein 1 VPREB1 Pre-B lymphocyte 1 SEQ ID NOS: 14137-14138 VPREB3 Pre-
B lymphocyte 3 SEQ ID NOS: 14139-14140 VPS37B Vacuolar protein sorting 37 homolog B SEQ ID
NOS: 14141-14143 (S. cerevisiae) VPS51 Vacuolar protein sorting 51 homolog SEQ ID NOS: 14144-
14155 (S. cerevisiae) VSIG1 V-set and immunoglobulin domain containing 1 SEQ ID NOS: 14156-14158
VSIG10 V-set and immunoglobulin domain containing 10 SEQ ID NOS: 14159-14160 VSTM1 V-set and
transmembrane domain containing 1 SEQ ID NOS: 14161-14167 VSTM2A V-set and transmembrane
domain containing 2A SEQ ID NOS: 14168-14171 VSTM2B V-set and transmembrane domain containing
2B SEQ ID NO: 14172 VSTM2L V-set and transmembrane domain containing 2 like SEQ ID NOS:
14173-14175 VSTM4 V-set and transmembrane domain containing 4 SEQ ID NOS: 14176-14177 VTN
Vitronectin SEQ ID NOS: 14178-14179 VWA1 Von Willebrand factor A domain containing 1 SEQ ID
NOS: 14180-14183 VWA2 Von Willebrand factor A domain containing 2 SEQ ID NOS: 14184-14185
VWA5B2 Von Willebrand factor A domain containing 5B2 SEQ ID NOS: 14186-14187 VWA7 Von
Willebrand factor A domain containing 7 SEQ ID NO: 14188 VWC2 Von Willebrand factor C domain
containing 2 SEQ ED NO: 14189 VWC2L Von Willebrand factor C domain containing SEQ ID NOS:
14190-14191 protein 2-like VWCE Von Willebrand factor C and EGF domains SEQ ID NOS: 14192-
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14196 VWDE Von Willebrand factor D and EGF domains SEQ ID NOS: 14197-14202 VWF Von
Willebrand factor SEQ ID NOS: 14203-14205 WDR25 WD repeat domain 25 SEQ ID NOS: 14206-14212
WDR81 WD repeat domain 81 SEQ ID NOS: 14213-14222 WDR90 WD repeat domain 90 SEQ ID NOS:
14223-14230 WFDC1 WAP four-disulfide core domain 1 SEQ ID NOS: 14231-14233 WFDC10A WAP
four-disulfide core domain 10A SEQ ID NO: 14234 WFDC10B WAP four-disulfide core domain 10B
SEQ ID NOS: 14235-14236 WFDC11 WAP four-disulfide core domain 11 SEQ ID NOS: 14237-14239
WFDC12 WAP four-disulfide core domain 12 SEQ ID NO: 14240 WFDC13 WAP four-disulfide core
domain 13 SEQ ID NO: 14241 WFDC2 WAP four-disulfide core domain 2 SEQ ID NOS: 14242-14246
WFDC3 WAP four-disulfide core domain 3 SEQ ID NOS: 14247-14250 WFDC5 WAP four-disulfide core
domain 5 SEQ ID NOS: 14251-14252 WFDC6 WAP four-disulfide core domain 6 SEQ ID NOS: 14253-
14254 WFDC8 WAP four-disulfide core domain 8 SEQ ID NOS: 14255-14256 WFIKKN1 WAP,
follistatin/kazal, immunoglobulin, kunitz SEQ ID NO: 14257 and netrin domain containing 1 WFIKKN2
WAP, follistatin/kazal, immunoglobulin, kunitz SEQ ID NOS: 14258-14259 and netrin domain containing
2 DFNB31 Deafness, autosomal recessive 31 SEQ ID NOS: 14260-14263 WIF1 WNT inhibitory fac tor I
SEQ ID NOS: 14264-14266 WISP1 WNT1 inducible signaling pathway protein 1 SEQ ID NOS: 14267-
14271 WISP2 WNT1 inducible signaling pathway protein 2 SEQ ID NOS: 14272-14274 WISP3 WNT1
inducible signaling pathway protein 3 SEQ ID NOS: 14275-14282 WNK1 WNK lysine deficient protein
kinase 1 SEQ ID NOS: 14283-14296 WNT1 Wingless-type MMTV integration site family, SEQ ID NOS:
14297-14298 member 1 WNT10B Wingless-type MMTV integration site family, SEQ ID NOS: 14299-
14303 member 10B WNT11 Wingless-type MMTV integration site family, SEQ ID NOS: 14304-14306
member 11 WNT16 Wingless-type MMTV integration site family, SEQ ID NOS: 14307-14308 member
16 WNT2 Wingless-type MMTV integration site family SEQ ID NOS: 14309-14311 member 2 WNT3
Wingless-type MMTV integration site family, SEQ ID NO: 14312 member 3 WNT3A Wingless-type
MMTV integration site family, SEQ ID NO: 14313 member 3A WNT5A Wingless-type MMTV
integration site family, SEQ ID NOS: 14314-14317 member 5A WNT5B Wingless-type MMTV
integration site family, SEQ ID NOS: 14318-14324 member 5B WNT6 Wingless-type MMTV integration
site family, SEQ ID NO: 14325 member 6 WNT7A Wingless-type MMTV integration site family, SEQ ID
NO: 14326 member 7A WNT7B Wingless-type MMTV integration site family, SEQ ID NOS: 14327-
14331 member 7B WNT8A Wingless-type MMTV integration site family, SEQ ID NOS: 14332-14335
member 8A WNT8B Wingless-type MMTV integration site family, SEQ ID NO: 14336 member 8B
WNT9A Wingless-type MMTV integration site family, SEQ ID NO: 14337 member 9A WNT9B
Wingless-type MMTV integration site family, SEQ ID NOS: 14338-14340 member 9B WSB1 WD repeat
and SOCS box containing 1 SEQ ID NOS: 14341-14350 WSCD1 WSC domain containing 1 SEQ ID
NOS: 14351-14360 WSCD2 WSC domain containing 2 SEQ ID NOS: 14361-14364 XCL1 Chemokine (C
motif) ligand 1 SEQ ID NO: 14365 XCL2 Chemokine (C motif) ligand 2 SEQ ID NO: 14366 XPNPEP2
X-prolyl aminopeptidase (aminopeptidase P) 2, SEQ ID NOS: 14367-14368 membrane-bound XXYLT1
Xyloside xylosyltransferase I SEQ ID NOS: 14369-14374 XYLT1 Xylosyltransferase I SEQ ID NO:
14375 XYLT2 Xylosyltransferase II SEQ ID NOS: 14376-14381 ZFYVE21 Zinc finger, FYVE domain
containing 21 SEQ ID NOS: 14382-14386 ZG16 Zymogen granule protein 16 SEQ ID NO: 14387 ZG16B
Zymogen granule protein 16B SEQ ID NOS: 14388-14391 ZIC4 Zic family member 4 SEQ ID NOS:
14392-14400 ZNF207 Zinc finger protein 207 SEQ ID NOS: 14401-14411 ZNF26 Zinc finger protein 26
SEQ ID NOS: 14412-14415 ZNF34 Zinc finger protein 34 SEQ ID NOS: 14416-14419 ZNF419 Zinc
finger protein 419 SEQ ID NOS: 14420-14434 ZNF433 Zinc finger protein 433 SEQ ID NOS: 14435-
14444 ZNF449 Zinc finger protein 449 SEQ ID NOS: 14445-14446 ZNF488 Zinc finger protein 488 SEQ
ID NOS: 14447-14448 ZNF511 Zinc finger protein 511 SEQ ID NOS: 14449-14450 ZNF570 Zinc finger
protein 570 SEQ ID NOS: 14451-14456 ZNF691 Zinc finger protein 691 SEQ ID NOS: 14457-14464
ZNF98 Zinc finger protein 98 SEQ ID NOS: 14465-14468 ZPBP Zona pellucida binding protein SEQ ID
NOS: 14469-14472 ZPBP2 Zona pellucida binding protein 2 SEQ ID NOS: 14473-14476 ZSCAN29 Zinc
finger and SCAN domain containing 29 SEQ ID NOS: 14477-14483
Expression of Cell Markers
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(471) In certain embodiments of the disclosure, T cells are modified to express detectable markers or indicators. In some embodiments, these detectable markers include, but are not limited to, fluorescent proteins. Non-limiting examples of fluorescent proteins include TagBFP, mTagBFP2, Azurite, EBFP2,

mKalama1, Sirius, Sapphire, T-Sapphire, ECFP, Cerulean, SCFP3A, mTurquoise, mTurquoise2, monomeric Midorishi-Cyan, TagCFP, mTFP1, EGFP, Emerald, Superfolder GFP, monomeric Azami Green, mUKG, mWasabi, Clover, mNeonGreen, EYFP, Citrine, Venus, SYFP2, TagYFP, monomeric Kusabira Orange, mKok, mKO2, mOrange, mOrange2, mRaspberry, mCherr, mStrawberry, mTangerine, tdTomato, TagRFP, TagFRP-T, mApple, mRuby, mRuby2, mPlum, HcRed-Tandem, mKate2, mNeptune, NiRFP, TagRFP657, IFP1.4, mRFP, mKeima Red, LSS-mKate1, LSS-mKate2, mBeRFPand spectrally shifted variants thereof. In some embodiments of the disclosure, the detectable marker or indicator comprises luciferase. In some embodiments, the detectable marker or indicator is codon optimized for expression in humans. In some embodiments, the detectable marker or indicator is an intracellular marker or indicator. In some embodiments, the detectable marker or indicator is a cytoplasmic marker or indicator. In some embodiments, the detectable marker or indicator is a nuclear marker or indicator. In some embodiments, the detectable marker or indicator is a mitochondrial marker or indicator. In some embodiments, the detectable marker or indicator is a cell surface marker. In some embodiments, particularly those embodiments where the markers or indicators are cell surface markers, the marker or indicator may be tethered to the membrane of the cell. Cells modified to express markers with the compositions and methods of the disclosure can be used as indicator cells in vivo, ex vivo, in vitro and in situ. In certain embodiments of the disclosure, a marker or indicator is under the control of an inducible promoter of the disclosure such that when the inducible promoter is targeted, the promoter induces expression of the marker or indicator.

(472) Inducible Promoters

- (473) In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding an NFκB promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding an interferon (IFN) promoter or a sequence encoding an interleukin-2 promoter. In certain embodiments, the interferon (IFN) promoter is an IFNγ promoter. In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a cytokine or a chemokine. In certain embodiments, the cytokine or chemokine comprises IL2, IL3, IL4, IL5, IL6, IL10, IL12, IL13, IL17A/F, IL21, IL22, 1L23, transforming growth factor beta (TGFβ), colony stimulating factor 2 (GM-CSF), interferon gamma (IFNγ), Tumor necrosis factor (TNFα), LTα, perforin, Granzyme C (Gzmc), Granzyme B (Gzmb), C-C motif chemokine ligand 5 (CCL5), C-C motif chemokine ligand 4 (Ccl4), C-C motif chemokine ligand 3 (Ccl3), X-C motif chemokine ligand 1 (Xcl1) and LIF interleukin 6 family cytokine (Lif).
- (474) In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a gene comprising a surface protein involved in cell differentiation, activation, exhaustion and function. In certain embodiments, the gene comprises CD69, CD71, CTLA4. PD-1, TIGIT, LAG3, TIM-3, GITR, MHCII, COX-2, FASL and 4-1BB.
- (475) In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a gene involved in CD metabolism and differentiation. In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of Nr4a1, Nr4a3, Tnfrsf9 (4-1BB), Sema7a, Zfp3612, Gadd45b, Dusp5, Dusp6 and Neto2. (476) Nucleic Acid Molecules
- (477) Nucleic acid molecules of the disclosure encoding protein scaffolds can be in the form of RNA, such as mRNA, hnRNA, tRNA or any other form, or in the form of DNA, including, but not limited to, cDNA and genomic DNA obtained by cloning or produced synthetically, or any combinations thereof. The DNA can be triple-stranded, double-stranded or single-stranded, or any combination thereof. Any portion of at least one strand of the DNA or RNA can be the coding strand, also known as the sense strand, or it can be the non-coding strand, also referred to as the anti-sense strand.
- (478) Isolated nucleic acid molecules of the disclosure can include nucleic acid molecules comprising an open reading frame (ORF), optionally, with one or more introns, e.g., but not limited to, at least one specified portion of at least one protein scaffold; nucleic acid molecules comprising the coding sequence for a protein scaffold or loop region that binds to the target protein; and nucleic acid molecules which comprise a nucleotide sequence substantially different from those described above but which, due to the degeneracy of the genetic code, still encode the protein scaffold as described herein and/or as known in the

art. Of course, the genetic code is well known in the art. Thus, it would be routine for one skilled in the art to generate such degenerate nucleic acid variants that code for specific protein scaffolds of the present invention. See, e.g., Ausubel, et al., supra, and such nucleic acid variants are included in the present invention.

(479) As indicated herein, nucleic acid molecules of the disclosure which comprise a nucleic acid encoding a protein scaffold can include, but are not limited to, those encoding the amino acid sequence of a protein scaffold fragment, by itself; the coding sequence for the entire protein scaffold or a portion thereof; the coding sequence for a protein scaffold, fragment or portion, as well as additional sequences, such as the coding sequence of at least one signal leader or fusion peptide, with or without the aforementioned additional coding sequences, such as at least one intron, together with additional, non-coding sequences, including but not limited to, non-coding 5' and 3' sequences, such as the transcribed, non-translated sequences that play a role in transcription, mRNA processing, including splicing and polyadenylation signals (for example, ribosome binding and stability of mRNA); an additional coding sequence that codes for additional amino acids, such as those that provide additional functionalities. Thus, the sequence encoding a protein scaffold can be fused to a marker sequence, such as a sequence encoding a peptide that facilitates purification of the fused protein scaffold comprising a protein scaffold fragment or portion.

(480) Polynucleotides Selectively Hybridizing to a Polynucleotide as Described Herein

(481) The disclosure provides isolated nucleic acids that hybridize under selective hybridization conditions to a polynucleotide disclosed herein. Thus, the polynucleotides of this embodiment can be used for isolating, detecting, and/or quantifying nucleic acids comprising such polynucleotides. For example, polynucleotides of the present invention can be used to identify, isolate, or amplify partial or full-length clones in a deposited library. In some embodiments, the polynucleotides are genomic or cDNA sequences isolated, or otherwise complementary to, a cDNA from a human or mammalian nucleic acid library. (482) Preferably, the cDNA library comprises at least 80% full-length sequences, preferably, at least 85% or 90% full-length sequences, and, more preferably, at least 95% full-length sequences. The cDNA libraries can be normalized to increase the representation of rare sequences. Low or moderate stringency hybridization conditions are typically, but not exclusively, employed with sequences having a reduced sequence identity relative to complementary sequences. Moderate and high stringency conditions can optionally be employed for sequences of greater identity. Low stringency conditions allow selective hybridization of sequences having about 70% sequence identity and can be employed to identify orthologous or paralogous sequences.

(483) Optionally, polynucleotides of this invention will encode at least a portion of a protein scaffold encoded by the polynucleotides described herein. The polynucleotides of this invention embrace nucleic acid sequences that can be employed for selective hybridization to a polynucleotide encoding a protein scaffold of the present invention. See, e.g., Ausubel, supra; Colligan, supra, each entirely incorporated herein by reference.

(484) Construction of Nucleic Acids

(485) The isolated nucleic acids of the disclosure can be made using (a) recombinant methods, (b) synthetic techniques, (c) purification techniques, and/or (d) combinations thereof, as well-known in the art. (486) The nucleic acids can conveniently comprise sequences in addition to a polynucleotide of the present invention. For example, a multi-cloning site comprising one or more endonuclease restriction sites can be inserted into the nucleic acid to aid in isolation of the polynucleotide. Also, translatable sequences can be inserted to aid in the isolation of the translated polynucleotide of the disclosure. For example, a hexa-histidine marker sequence provides a convenient means to purify the proteins of the disclosure. The nucleic acid of the disclosure, excluding the coding sequence, is optionally a vector, adapter, or linker for cloning and/or expression of a polynucleotide of the disclosure.

(487) Additional sequences can be added to such cloning and/or expression sequences to optimize their function in cloning and/or expression, to aid in isolation of the polynucleotide, or to improve the introduction of the polynucleotide into a cell. Use of cloning vectors, expression vectors, adapters, and linkers is well known in the art. (See, e.g., Ausubel, supra; or Sambrook, supra).

(488) Recombinant Methods for Constructing Nucleic Acids

(489) The isolated nucleic acid compositions of this disclosure, such as RNA, cDNA, genomic DNA, or

any combination thereof, can be obtained from biological sources using any number of cloning methodologies known to those of skill in the art. In some embodiments, oligonucleotide probes that selectively hybridize, under stringent conditions, to the polynucleotides of the present invention are used to identify the desired sequence in a cDNA or genomic DNA library. The isolation of RNA, and construction of cDNA and genomic libraries are well known to those of ordinary skill in the art. (See, e.g., Ausubel, supra; or Sambrook, supra).

(490) Nucleic Acid Screening and Isolation Methods

(491) A cDNA or genomic library can be screened using a probe based upon the sequence of a polynucleotide of the disclosure. Probes can be used to hybridize with genomic DNA or cDNA sequences to isolate homologous genes in the same or different organisms. Those of skill in the art will appreciate that various degrees of stringency of hybridization can be employed in the assay; and either the hybridization or the wash medium can be stringent. As the conditions for hybridization become more stringent, there must be a greater degree of complementarity between the probe and the target for duplex formation to occur. The degree of stringency can be controlled by one or more of temperature, ionic strength, pH and the presence of a partially denaturing solvent, such as formamide. For example, the stringency of hybridization is conveniently varied by changing the polarity of the reactant solution through, for example, manipulation of the concentration of formamide within the range of 0% to 50%. The degree of complementarity (sequence identity) required for detectable binding will vary in accordance with the stringency of the hybridization medium and/or wash medium. The degree of complementarity will optimally be 100%, or 70-100%, or any range or value therein. However, it should be understood that minor sequence variations in the probes and primers can be compensated for by reducing the stringency of the hybridization and/or wash medium.

(492) Methods of amplification of RNA or DNA are well known in the art and can be used according to the disclosure without undue experimentation, based on the teaching and guidance presented herein. (493) Known methods of DNA or RNA amplification include, but are not limited to, polymerase chain reaction (PCR) and related amplification processes (see, e.g., U.S. Pat. Nos. 4,683,195, 4,683,202, 4,800,159, 4,965,188, to Mullis, et al.; U.S. Pat. Nos. 4,795,699 and 4,921,794 to Tabor, et al; U.S. Pat. No. 5,142,033 to Innis; U.S. Pat. No. 5,122,464 to Wilson, et al.; U.S. Pat. No. 5,091,310 to Innis; U.S. Pat. No. 5,066,584 to Gyllensten, et al; U.S. Pat. No. 4,889,818 to Gelfand, et al; U.S. Pat. No. 4,994,370 to Silver, et al; U.S. Pat. No. 4,766,067 to Biswas; U.S. Pat. No. 4,656,134 to Ringold) and RNA mediated amplification that uses anti-sense RNA to the target sequence as a template for double-stranded DNA synthesis (U.S. Pat. No. 5,130,238 to Malek, et al, with the tradename NASBA), the entire contents of which references are incorporated herein by reference. (See, e.g., Ausubel, supra; or Sambrook, supra.) (494) For instance, polymerase chain reaction (PCR) technology can be used to amplify the sequences of polynucleotides of the disclosure and related genes directly from genomic DNA or cDNA libraries. PCR and other in vitro amplification methods can also be useful, for example, to clone nucleic acid sequences that code for proteins to be expressed, to make nucleic acids to use as probes for detecting the presence of the desired mRNA in samples, for nucleic acid sequencing, or for other purposes. Examples of techniques sufficient to direct persons of skill through in vitro amplification methods are found in Berger, supra, Sambrook, supra, and Ausubel, supra, as well as Mullis, et al., U.S. Pat. No. 4,683,202 (1987); and Innis, et al., PCR Protocols A Guide to Methods and Applications, Eds., Academic Press Inc., San Diego, Calif. (1990). Commercially available kits for genomic PCR amplification are known in the art. See, e.g., Advantage-GC Genomic PCR Kit (Clontech). Additionally, e.g., the T4 gene 32 protein (Boehringer Mannheim) can be used to improve yield of long PCR products.

(495) Synthetic Methods for Constructing Nucleic Acids

(496) The isolated nucleic acids of the disclosure can also be prepared by direct chemical synthesis by known methods (see, e.g., Ausubel, et al., supra). Chemical synthesis generally produces a single-stranded oligonucleotide, which can be converted into double-stranded DNA by hybridization with a complementary sequence, or by polymerization with a DNA polymerase using the single strand as a template. One of skill in the art will recognize that while chemical synthesis of DNA can be limited to sequences of about 100 or more bases, longer sequences can be obtained by the ligation of shorter sequences.

(497) Recombinant Expression Cassettes

(498) The disclosure further provides recombinant expression cassettes comprising a nucleic acid of the disclosure. A nucleic acid sequence of the disclosure, for example, a cDNA or a genomic sequence encoding a protein scaffold of the disclosure, can be used to construct a recombinant expression cassette that can be introduced into at least one desired host cell. A recombinant expression cassette will typically comprise a polynucleotide of the disclosure operably linked to transcriptional initiation regulatory sequences that will direct the transcription of the polynucleotide in the intended host cell. Both heterologous and non-heterologous (i.e., endogenous) promoters can be employed to direct expression of the nucleic acids of the disclosure.

(499) In some embodiments, isolated nucleic acids that serve as promoter, enhancer, or other elements can be introduced in the appropriate position (upstream, downstream or in the intron) of a non-heterologous form of a polynucleotide of the disclosure so as to up or down regulate expression of a polynucleotide of the disclosure. For example, endogenous promoters can be altered in vivo or in vitro by mutation, deletion and/or substitution.

(500) Vectors and Host Cells

- (501) The disclosure also relates to vectors that include isolated nucleic acid molecules of the disclosure, host cells that are genetically engineered with the recombinant vectors, and the production of at least one protein scaffold by recombinant techniques, as is well known in the art. See, e.g., Sambrook, et al., supra; Ausubel, et al., supra, each entirely incorporated herein by reference.
- (502) For example, the PB-EF1a vector may be used. The vector comprises the following nucleotide sequence:

(503) TABLE-US-00133 (SEQ ID NO. 17073)

cagcagcgagccgcccggggctccgctccggtccggcgctccccccgcatccccgagccggcagcgtgcggggacagcccgggca cggggaaggtggcacgggatcgctttcctctgaacgcttctcgctgctctttgagcctgcagacacctggggggatacggggaaa agttgactgtgcctttcgatcgaaccatggacagttagctttgcaaagatggataaagttttaaacagagaggaatctttgcagc taatggaccttctaggtcttgaaaggagtgggaattggctccggtgcccgtcagtgggcagagcgcacatcgcccacagtccccg agaagttggggggggggggcgaattgaaccggtgcctagagaaggtggcgcggggtaaactgggaaagtgatgtcgtgtactg gctccgcctttttcccgagggtggggggagaaccgtatataagtgcagtagtcgccgtgaacgttctttttcgcaacgggtttgcc gccagaacacaggtaagtgccgtgtgtgttcccgcgggcctggcctctttacgggttatggcccttgcgtgccttgaattactt cccttcgcctcgtgcttgagttgaggcctggcctgggcgctggggccgccgcgtgcgaatctggtggcaccttcgcgcctgtctc gctgctttcgataagtctctagccatttaaaatttttgatgacctgctgcgacgctttttttctggcaagatagtcttgtaaatg ccgccgtgtatcgccccgccctgggcggcaaggctggcccggtcggcaccagttgcgtgagcggaaagatggccgcttcccggcc gtcctcagccgtcgcttcatgtgactccacggagtaccgggcgccgtccaggcacctcgattagttctcgagcttttggagtacg acttgatgtaattctccttggaatttgccctttttgagtttggatcttggttcattctcaagcctcagacagtggttcaaagttt ttttcttccatttcaggtgtcgtgagaattctaatacgactcactatagggtgtgctgtctcatcattttggcaaagattggcca ccaagcttgtcctgcaggagggtcgacgcctctagacgggcggccgctccggatccacgggtaccgatcacatatgcctttaatt aaacactagttctatagtgtcacctaaattccctttagtgagggttaatggccgtaggccgccagaattgggtccagacatgata agatacattgatgagtttggacaaaccacaactagaatgcagtgaaaaaaatgctttatttgtgaaatttgtgatgctattgctt ggaggttttttcggactctaggacctgcgcatgcgcttggcgtaatcatggtcatagctgtttcctgttttccccgtatcccccc aggtgtctgcaggctcaaagagcagcgagaagcgttcagaggaaagcgatcccgtgccaccttccccgtgcccgggctgtccccg gacgtaattacatccctggggggctttgggggggggggtgtccctctcaccgcggtggagctccagcttttgttcgaattggggccc aattatcgtatgagttaaatcttaaaagtcacgtaaaagataatcatgcgtcattttgactcacgcggtcgttatagttcaaaat

cagtgacacttaccgcattgacaagcacgcctcacgggagctccaagcggcgactgagatgtcctaaatgcacagcgacggattc cct taags gcgcct att gcgt tgcgct cact gcccgctt tccag tcgggaaacct gtcgt gccag ctgcat taat gaat cggccalland gaat cggccalland gcgccag ctgcat taat gcgcag ctgcag ctacgcgcggggagaggcggtttgcgtattgggcgctcttccgcttcctcgctcactgactcgctgcgctcggtcgttcggctgcgg cgagcggtatcagctcactcaaaggcggtaatacggttatccacagaatcaggggataacgcaggaaagaacatgaccaaaatcc cttaacgtgagttttcgttccactgagcgtcagaccccgtagaaaagatcaaaggatcttcttgagatcctttttttctgcgcgt gtaactggcttcagcagagcgcagataccaaatactgttcttctagtgtagccgtagttaggccaccacttcaagaactctgtag caccgcctacatacctcgctctgctaatcctgttaccagtggctgctgccagtggcgataagtcgtgtcttaccgggttggactc aagacgatagttaccggataaggcgcagcggtcgggctgaacggggggttcgtgcacacagcccagcttggagcgaacgacctac accgaactgagatacctacagcgtgagctatgagaaagcgccacgcttcccgaagggagaaaggcggacaggtatccggtaagcg gcagggtcggaacaggagagcgcacgagggagcttccagggggaaacgcctggtatctttatagtcctgtcgggtttcgccacct ctggccttttgctggccttttgctcacatgagattatcaaaaaggatcttcacctagatccttttaaattaaaaatgaagtttta aatcaatctaaagtatatatgagtaaacttggtctgacagtcagaagaactcgtcaagaaggcgatagaaggcgatgcgctgcga atcgggagcggcgataccgtaaagcacgaggaagcggtcagcccattcgccgccaagctcttcagcaatatcacgggtagccaac gctatgtcctgatagcggtccgccacacccagccggccacagtcgatgaatccagaaaagcggccattttccaccatgatattcg gcaagcaggcatcgccatgggtcacgacgagatcctcgccgtcgggcatgctcgccttgagcctggcgaacagttcggctggcgc gcttggtggtcgaatgggcaggtagccggatcaagcgtatgcagccgccgcattgcatcagccatgatggatactttctcggcag gagcaaggtgagatgacaggagatcctgccccggcacttcgcccaatagcagccagtcccttcccgcttcagtgacaacgtcgag tcggtcttgacaaaaagaaccgggcgcccctgcgctgacagccggaacacggcggcatcagagcagccgattgtctgttgtgccc agtcatagccgaatagcctctccacccaagcggccggagaacctgcgtgcaatccatcttgttcaatcataatattattgaagca tttatcagggttcgtctcgtcccggtctcctcccaatgcatgtcaatattggccattagccatattattcattggttatatagca taaatcaatattggctattggccattgcatacgttgtatctatatcataata.

(504) The polynucleotides can optionally be joined to a vector containing a selectable marker for propagation in a host. Generally, a plasmid or nanoplasmid vector is introduced in a precipitate, such as a calcium phosphate precipitate, or in a complex with a charged lipid. If the vector is a virus, it can be packaged in vitro using an appropriate packaging cell line and then transduced into host cells. (505) The DNA insert should be operatively linked to an appropriate promoter. The expression constructs will further contain sites for transcription initiation, termination and, in the transcribed region, a ribosome binding site for translation. The coding portion of the mature transcripts expressed by the constructs will preferably include a translation initiating at the beginning and a termination codon (e.g., UAA, UGA or UAG) appropriately positioned at the end of the mRNA to be translated, with UAA and UAG preferred for mammalian or eukaryotic cell expression.

(506) Expression vectors will preferably but optionally include at least one selectable marker. Such markers include, e.g., but are not limited to, ampicillin, zeocin (sh bla gene), puromycin (pac gene), hygromycin B (hygB gene), G418/Geneticin (neo gene), mycophenolic acid, or glutamine synthetase (GS, U.S. Pat. Nos. 5,122,464; 5,770,359; 5,827,739), blasticidin (bsd gene), resistance genes for eukaryotic cell culture as well as ampicillin, zeocin (Sh bla gene), puromycin (pac gene), hygromycin B (hvgB gene), G418/Geneticin (neo gene), kanamycin, spectinomycin, streptomycin, carbenicillin, bleomycin, erythromycin, polymyxin B, or tetracycline resistance genes for culturing in *E. coli* and other bacteria or prokaryotics (the above patents are entirely incorporated hereby by reference). Appropriate culture mediums and conditions for the above-described host cells are known in the art. Suitable vectors will be readily apparent to the skilled artisan. Introduction of a vector construct into a host cell can be effected by calcium phosphate transfection. DEAE-dextran mediated transfection, cationic lipid-mediated transfection, electroporation, transduction, infection or other known methods. Such methods are described in the art, such as Sambrook, supra, Chapters 1-4 and 16-18; Ausubel, supra, Chapters 1, 9, 13, 15, 16. (507) Expression vectors will preferably but optionally include at least one selectable cell surface marker for isolation of cells modified by the compositions and methods of the disclosure. Selectable cell surface markers of the disclosure comprise surface proteins, glycoproteins, or group of proteins that distinguish a cell or subset of cells from another defined subset of cells. Preferably the selectable cell surface marker

distinguishes those cells modified by a composition or method of the disclosure from those cells that are not modified by a composition or method of the disclosure. Such cell surface markers include, e.g., but are not limited to, "cluster of designation" or "classification determinant" proteins (often abbreviated as "CD") such as a truncated or full length form of CD19, CD271, CD34, CD22, CD20, CD33, CD52, or any combination thereof. Cell surface markers further include the suicide gene marker RQR8 (Philip B et al. Blood. 2014 Aug. 21; 124(8):1277-87).

- (508) Expression vectors will preferably but optionally include at least one selectable drug resistance marker for isolation of cells modified by the compositions and methods of the disclosure. Selectable drug resistance markers of the disclosure may comprise wild-type or mutant Neo, DHFR, TYMS, FRANCF, RAD51C, GCS, MDR1, ALDH1, NKX2.2, or any combination thereof.
- (509) At least one protein scaffold of the disclosure can be expressed in a modified form, such as a fusion protein, and can include not only secretion signals, but also additional heterologous functional regions. For instance, a region of additional amino acids, particularly charged amino acids, can be added to the N-terminus of a protein scaffold to improve stability and persistence in the host cell, during purification, or during subsequent handling and storage. Also, peptide moieties can be added to a protein scaffold of the disclosure to facilitate purification. Such regions can be removed prior to final preparation of a protein scaffold or at least one fragment thereof. Such methods are described in many standard laboratory manuals, such as Sambrook, supra, Chapters 17.29-17.42 and 18.1-18.74; Ausubel, supra. Chapters 16, 17 and 18.
- (510) Those of ordinary skill in the art are knowledgeable in the numerous expression systems available for expression of a nucleic acid encoding a protein of the disclosure. Alternatively, nucleic acids of the disclosure can be expressed in a host cell by turning on (by manipulation) in a host cell that contains endogenous DNA encoding a protein scaffold of the disclosure. Such methods are well known in the art, e.g., as described in U.S. Pat. Nos. 5,580,734, 5,641,670, 5,733,746, and 5,733,761, entirely incorporated herein by reference.
- (511) Illustrative of cell cultures useful for the production of the protein scaffolds, specified portions or variants thereof, are bacterial, yeast, and mammalian cells as known in the art. Mammalian cell systems often will be in the form of monolayers of cells although mammalian cell suspensions or bioreactors can also be used. A number of suitable host cell lines capable of expressing intact glycosylated proteins have been developed in the art, and include the COS-1 (e.g., ATCC CRL 1650), COS-7 (e.g., ATCC CRL-1651), HEK293, BHK21 (e.g., ATCC CRL-10), CHO (e.g., ATCC CRL 1610) and BSC-1 (e.g., ATCC CRL-26) cell lines, Cos-7 cells, CHO cells, hep G2 cells, P3X63Ag8.653, SP2/0-Ag14, 293 cells, HeLa cells and the like, which are readily available from, for example, American Type Culture Collection, Manassas, Va. (www.atcc.org). Preferred host cells include cells of lymphoid origin, such as myeloma and lymphoma cells. Particularly preferred host cells are P3X63Ag8.653 cells (ATCC Accession Number CRL-1580) and SP2/0-Ag14 cells (ATCC Accession Number CRL-1851). In a particularly preferred embodiment, the recombinant cell is a P3X63Ab8.653 or an SP2/0-Ag14 cell.
- (512) Expression vectors for these cells can include one or more of the following expression control sequences, such as, but not limited to, an origin of replication; a promoter (e.g., late or early SV40 promoters, the CMV promoter (U.S. Pat. Nos. 5,168,062; 5,385,839), an HSV tk promoter, a pgk (phosphoglycerate kinase) promoter, an EF-1 alpha promoter (U.S. Pat. No. 5,266,491), at least one human promoter; an enhancer, and/or processing information sites, such as ribosome binding sites, RNA splice sites, polyadenylation sites (e.g., an SV40 large T Ag poly A addition site), and transcriptional terminator sequences. See, e.g., Ausubel et al., supra; Sambrook, et al., supra. Other cells useful for production of nucleic acids or proteins of the present invention are known and/or available, for instance, from the American Type Culture Collection Catalogue of Cell Lines and Hybridomas (www.atcc.org) or other known or commercial sources.
- (513) When eukaryotic host cells are employed, polyadenylation or transcription terminator sequences are typically incorporated into the vector. An example of a terminator sequence is the polyadenlyation sequence from the bovine growth hormone gene. Sequences for accurate splicing of the transcript can also be included. An example of a splicing sequence is the VP1 intron from SV40 (Sprague, et al., J. Virol. 45:773-781 (1983)). Additionally, gene sequences to control replication in the host cell can be incorporated into the vector, as known in the art.

(514) Amino Acid Codes

- (515) The amino acids that make up protein scaffolds of the disclosure are often abbreviated. The amino acid designations can be indicated by designating the amino acid by its single letter code, its three letter code, name, or three nucleotide codon(s) as is well understood in the art (see Alberts, B., et al., Molecular Biology of The Cell, Third Ed., Garland Publishing, Inc., New York, 1994). A protein scaffold of the disclosure can include one or more amino acid substitutions, deletions or additions, either from natural mutations or human manipulation, as specified herein. Amino acids in a protein scaffold of the disclosure that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis (e.g., Ausubel, supra, Chapters 8, 15; Cunningham and Wells, Science 244:1081-1085 (1989)). The latter procedure introduces single alanine mutations at every residue in the molecule. The resulting mutant molecules are then tested for biological activity, such as, but not limited to, at least one neutralizing activity. Sites that are critical for protein scaffold binding can also be identified by structural analysis, such as crystallization, nuclear magnetic resonance or photoaffinity labeling (Smith, et al., J. Mol. Biol. 224:899-904 (1992) and de Vos, et al., Science 255:306-312 (1992)). (516) As those of skill will appreciate, the invention includes at least one biologically active protein scaffold of the disclosure. Biologically active protein scaffolds have a specific activity at least 20%, 30%, or 40%, and, preferably, at least 50%, 60%, or 70%, and, most preferably, at least 80%, 90%, or 95%-99% or more of the specific activity of the native (non-synthetic), endogenous or related and known protein scaffold. Methods of assaying and quantifying measures of enzymatic activity and substrate specificity are well known to those of skill in the art.
- (517) In another aspect, the disclosure relates to protein scaffolds and fragments, as described herein, which are modified by the covalent attachment of an organic moiety. Such modification can produce a protein scaffold fragment with improved pharmacokinetic properties (e.g., increased in vivo serum half-life). The organic moiety can be a linear or branched hydrophilic polymeric group, fatty acid group, or fatty acid ester group. In particular embodiments, the hydrophilic polymeric group can have a molecular weight of about 800 to about 120,000 Daltons and can be a polyalkane glycol (e.g., polyethylene glycol (PEG), polypropylene glycol (PPG)), carbohydrate polymer, amino acid polymer or polyvinyl pyrolidone, and the fatty acid or fatty acid ester group can comprise from about eight to about forty carbon atoms. (518) T Cell Isolation From a Leukopheresis Product
- (519) A leukapheresis product or blood may be collected from a subject at clinical site using a closed system and standard methods (e.g., a COBE Spectra Apheresis System) Preferably, the product is collected according to standard hospital or institutional Leukapheresis procedures in standard Leukapheresis collection bags. For example, in preferred embodiments of the methods of the disclosure, no additional anticoagulants or blood additives (heparin, etc.) are included beyond those normally used during leukapheresis.
- (520) Alternatively, white blood cells (WBC)/Peripheral Blood Mononuclear Cells (PBMC) (using Biosafe Sepax 2 (Closed/Automated)) or T cells (using CliniMACS® Prodigy (Closed/Automated)) may be isolated directly from whole blood. However, in certain subjects (e.g. those diagnosed and/or treated for cancer), the WBC/PBMC yield may be significantly lower when isolated from whole blood than when isolated by leukapheresis.
- (521) Either the leukapheresis procedure and/or the direct cell isolation procedure may be used for any subject of the disclosure.
- (522) The leukapheresis product, blood. WBC/PBMC composition and/or T-cell composition should be packed in insulated containers and should be kept at controlled room temperature (+19° C. to +25° C.) according to standard hospital of institutional blood collection procedures approved for use with the clinical protocol. The leukapheresis product, blood, WBC/PBMC composition and/or T-cell composition should not be refrigerated.
- (523) The cell concentration leukapheresis product, blood. WBC/PBMC composition and/or T-cell composition should not exceed 0.2×10.sup.9 cells per mL during transportation. Intense mixing of the leukapheresis product, blood. WBC/PBMC composition and/or T-cell composition should be avoided. (524) If the leukapheresis product, blood, WBC/PBMC composition and/or T-cell composition has to be stored, e.g. overnight, it should be kept at controlled room temperature (same as above). During storage, the concentration of the leukapheresis product, blood, WBC/PBMC composition and/or T-cell composition

should never exceed 0.2×10.sup.9 cell per mL.

(525) Preferably, cells of the leukapheresis product, blood, WBC/PBMC composition and/or T-cell composition should be stored in autologous plasma. In certain embodiments, if the cell concentration of the leukapheresis product, blood. WBC/PBMC composition and/or T-cell composition is higher than 0.2×10.sup.9 cell per mL, the product should be diluted with autologous plasma.

- (526) Preferably, the leukapheresis product, blood, WBC/PBMC composition and/or T-cell composition should not be older than 24 hours when starting the labeling and separation procedure. The leukapheresis product, blood, WBC/PBMC composition and/or T-cell composition may be processed and/or prepared for cell labeling using a closed and/or automated system (e.g., CliniMACS Prodigy).
- (527) An automated system may perform additional buffy coat isolation, possibly by ficolation, and/or washing of the cellular product (e.g., the leukapheresis product, blood. WBC/PBMC composition and/or T cell composition).
- (528) A closed and/or automated system may be used to prepare and label cells for T-Cell isolation (from, for example, the leukapheresis product, blood, WBC/PBMC composition and/or T cell composition). (529) Although WBC/PBMCs may be nucleofected directly (which is easier and saves additional steps), the methods of the disclosure may include first isolating T cells prior to nucleofection. The easier strategy of directly nucleofecting PBMC requires selective expansion of CAR+ cells that is mediated via CAR signaling, which by itself is proving to be an inferior expansion method that directly reduces the in vivo efficiency of the product by rendering T cells functionally exhausted. The product may be a heterogeneous composition of CAR+ cells including T cells, NK cells, NKT cells, monocytes, or any combination thereof, which increases the variability in product from patient to patient and makes dosing and CRS management more difficult. Since T cells are thought to be the primary effectors in tumor suppression and killing, T cell isolation for the manufacture of an autologous product may result in significant benefits over the other more heterogeneous composition.
- (530) T cells may be isolated directly, by enrichment of labeled cells or depletion of labeled cells in a oneway labeling procedure or, indirectly, in a two-step labeling procedure. According to certain enrichment strategies of the disclosure, T cells may be collected in a Cell Collection Bag and the non-labeled cells (non-target cells) in a Negative Fraction Bag. In contrast to an enrichment strategy of the disclosure, the non-labeled cells (target cells) are collected in a Cell Collection Bag and the labeled cells (non-target cells) are collected in a Negative Fraction Bag or in the Non-Target Cell Bag, respectively. Selection reagents may include, but are not limited to, antibody-coated beads. Antibody-coated beads may either be removed prior to a modification and/or an expansion step, or, retained on the cells prior to a modification and/or an expansion step. One or more of the following non-limiting examples of cellular markers may be used to isolate T-cells: CD3, CD4, CD8, CD25, anti-biotin, CD1c, CD3/CD19, CD3/CD56, CD14, CD19, CD34, CD45RA, CD56, CD62L, CD133, CD137, CD271, CD304, IFN-gamma, TCR alpha/beta, and/or any combination thereof. Methods for the isolation of T-cells may include one or more reagents that specifically bind and/or detectably-label one or more of the following non-limiting examples of cellular markers may be used to isolate T-cells. CD3, CD4, CD8, CD25, anti-biotin, CD1c, CD3/CD19, CD3/CD56, CD14, CD19, CD34, CD45RA, CD56, CD62L, CD133, CD137. CD271, CD304, IFNgamma, TCR alpha/beta, and/or any combination thereof. These reagents may or may not be "Good Manufacturing Practices" ("GMP") grade. Reagents may include, but are not limited to, Thermo DynaBeads and Miltenyi CliniMACS products. Methods of isolating T-cells of the disclosure may include multiple iterations of labeling and/or isolation steps. At any point in the methods of isolating T-cells of the disclosure, unwanted cells and/or unwanted cell types may be depleted from a T cell product composition of the disclosure by positively or negatively selecting for the unwanted cells and/or unwanted cell types. A T cell product composition of the disclosure may contain additional cell types that may express CD4, CD8, and/or another T cell marker(s).
- (531) Methods of the disclosure for nucleofection of T cells may eliminate the step of T cell isolation by, for example, a process for nucleofection of T cells in a population or composition of WBC/PBMCs that, following nucleofection, includes an isolation step or a selective expansion step via TCR signaling. (532) Certain cell populations may be depleted by positive or negative selection before or after T cell enrichment and/or sorting. Examples of cell compositions that may be depleted from a cell product composition may include myeloid cells, CD25+ regulatory T cells (T Regs), dendritic cells, macrophages,

- red blood cells, mast cells, gamma-delta T cells, natural killer (NK) cells, a Natural Killer (NK)-like cell (e.g. a Cytokine Induced Killer (CIK) cell), induced natural killer (iNK) T cells, NK T cells, B cells, or any combination thereof.
- (533) T cell product compositions of the disclosure may include CD4+ and CD8+ T-Cells. CD4+ and CD8+ T-Cells may be isolated into separate collection bags during an isolation or selection procedure. CD4+ T cells and CD8+ T cells may be further treated separately, or treated after reconstitution (combination into the same composition) at a particular ratio.
- (534) The particular ratio at which CD4+ T cells and CD8+ T cells may be reconstituted may depend upon the type and efficacy of expansion technology used, cell medium, and/or growth conditions utilized for expansion of T-cell product compositions. Examples of possible CD4+: CD8+ ratios include, but are not limited to, 50%:50%, 60%:40%, 40%:60% 75%:25% and 25%:75%.
- (535) CD8+ T cells exhibit a potent capacity for tumor cell killing, while CD4+ T cells provide many of the cytokines required to support CD8+ T cell proliferative capacity and function. Because T cells isolated from normal donors are predominantly CD4+, the T-cell product compositions are artificially adjusted in vitro with respect to the CD4+:CD8+ ratio to improve upon the ratio of CD4+ T cells to CD8+ T cells that would otherwise be present in vivo. An optimized ratio may also be used for the ex vivo expansion of the autologous T-cell product composition. In view of the artificially adjusted CD4+:CD8+ ratio of the T-cell product composition, it is important to note that the product compositions of the disclosure may be significantly different and provide significantly greater advantage than any endogenously-occurring population of T-cells.
- (536) Preferred methods for T cell isolation may include a negative selection strategy for yielding untouched pan T cell, meaning that the resultant T-cell composition includes T-cells that have not been manipulated and that contain an endogenously-occurring variety/ratio of T-cells.
- (537) Reagents that may be used for positive or negative selection include, but are not limited to, magnetic cell separation beads. Magnetic cell separation beads may or may not be removed or depleted from selected populations of CD4+ T cells, CD8+ T cells, or a mixed population of both CD4+ and CD8+ T cells before performing the next step in a T-cell isolation method of the disclosure.
- (538) T cell compositions and T cell product compositions may be prepared for cryopreservation, storage in standard T Cell Culture Medium. and/or genetic modification.
- (539) T cell compositions, T cell product compositions, unstimulated T cell compositions, resting T cell compositions or any portion thereof may be cryopreserved using a standard cryopreservation method optimized for storing and recovering human cells with high recovery, viability, phenotype, and/or functional capacity. Commercially-available cryopreservation media and/or protocols may be used. Cryopreservation methods of the disclosure may include a DMSO free cryopreservant (e.g. CryoSOfree<sup>TM</sup> DMSO-free Cryopreservation Medium) reduce freezing-related toxicity.
- (540) T cell compositions, T cell product compositions, unstimulated T cell compositions, resting T cell compositions or any portion thereof may be stored in a culture medium. T cell culture media of the disclosure may be optimized for cell storage, cell genetic modification, cell phenotype and/or cell expansion. T cell culture media of the disclosure may include one or more antibiotics. Because the inclusion of an antibiotic within a cell culture media may decrease transfection efficiency and/or cell yield following genetic modification via nucleofection, the specific antibiotics (or combinations thereof) and their respective concentration(s) may be altered for optimal transfection efficiency and/or cell yield following genetic modification via nucleofection.
- (541) T cell culture media of the disclosure may include serum, and, moreover, the serum composition and concentration may be altered for optimal cell outcomes. Human AB serum is preferred over FBS/FCS for culture of T cells because, although contemplated for use in T cell culture media of the disclosure, FBS/FCS may introduce xeno-proteins. Serum may be isolated form the blood of the subject for whom the T-cell composition in culture is intended for administration, thus, a T cell culture medium of the disclosure may comprise autologous serum. Serum-free media or serum-substitute may also be used in T-cell culture media of the disclosure. In certain embodiments of the T-cell culture media and methods of the disclosure, serum-free media or serum-substitute may provide advantages over supplementing the medium with xeno-serum, including, but not limited to, healthier cells that have greater viability, nucleofect with higher efficiency, exhibit greater viability post-nucleofection, display a more desirable cell phenotype, and/or

greater/faster expansion upon addition of expansion technologies.

(542) T cell culture media may include a commercially-available cell growth media. Exemplary commercially-available cell growth media include, but are not limited to, PBS, HBSS, OptiMEM, DMEM, RPMI 1640, AIM-V, X-VIVO 15, CellGro DC Medium, CTS OpTimizer T Cell Expansion SFM, TexMACS Medium, PRIME-XV T Cell Expansion Medium, ImmunoCult-XF T Cell Expansion Medium, or any combination thereof.

(543) T cell compositions, T cell product compositions, unstimulated T cell compositions, resting T cell compositions or any portion thereof may be prepared for genetic modification. Preparation of T cell compositions, T cell product compositions, unstimulated T cell compositions, resting T cell compositions or any portion thereof for genetic modification may include cell washing and/or resuspension in a desired nucleofection buffer. Cryopreserved T-cell compositions may be thawed and prepared for genetic modification by nucleofection. Cryopreserved cells may be thawed according to standard or known protocols. Thawing and preparation of cryopreserved cells may be optimized to yield cells that have greater viability, nucleofect with higher efficiency, exhibit greater viability post-nucleofection, display a more desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. For example, Grifols Albutein (25% human albumin) may be used in the thawing and/or preparation process.

(544) Genetic Modification of an Autologous T Cell Product Composition

(545) T cell compositions, T cell product compositions, unstimulated T cell compositions, resting T cell compositions or any portion thereof may be genetically modified using, for example, a nucleofection strategy such as electroporation. The total number of cells to be nucleofected, the total volume of the nucleofection reaction, and the precise timing of the preparation of the sample may be optimized to yield cells that have greater viability, nucleofect with higher efficiency, exhibit greater viability post-nucleofection, display a more desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies.

(546) Nucleofection and/or electroporation may be accomplished using, for example. Lonza Amaxa, MaxCyte PulseAgile, Harvard Apparatus BTX, and/or Invitrogen Neon. Non-metal electrode systems, including, but not limited to, plastic polymer electrodes, may be preferred for nucleofection. (547) Prior to genetic modification by nucleofection. T cell compositions, T cell product compositions, unstimulated T cell compositions, resting T cell compositions or any portion thereof may be resuspended in a nucleofection buffer. Nucleofection buffers of the disclosure include commercially-available nucleofection buffers. Nucleofection buffers of the disclosure may be optimized to yield cells that have greater viability, nucleofect with higher efficiency, exhibit greater viability post-nucleofection, display a more desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. Nucleofection buffers of the disclosure may include, but are not limited to, PBS, HBSS, OptiMEM, BTXpress. Amaxa Nucleofector. Human T cell nucleofection buffer and any combination thereof. Nucleofection buffers of the disclosure may comprise one or more supplemental factors to yield cells that have greater viability, nucleofect with higher efficiency, exhibit greater viability post-nucleofection, display a more desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. Exemplary supplemental factors include, but are not limited to, recombinant human cytokines, chemokines, interleukins and any combination thereof. Exemplary cytokines, chemokines, and interleukins include, but are not limited to, IL2, IL7, IL12, IL15, IL21, IL1, IL3, IL4, IL5, IL6, IL8, CXCL8, IL9, IL10, IL11, IL13, IL14, IL16, IL17, IL18, IL19, IL20, IL22, IL23, IL25, IL26, IL27, IL28, IL29, IL30, IL31, IL32, IL33, IL35, IL36, GM-CSF, IFN-gamma, IL-1 alpha/IL-1F1, IL-1 beta/IL-1F2, IL-12 p70, IL-12/IL-35 p35, IL-13, IL-17/IL-17A, IL-17A/F Heterodimer, IL-17F, IL-18/IL-1F4, IL-23, IL-24, IL-32, IL-32 beta, IL-32 gamma, IL-33, LAP (TGF-beta 1), Lymphotoxin-alpha/TNF-beta, TGFbeta, TNF-alpha, TRANCE/TNFSF11/RANK L and any combination thereof. Exemplary supplemental factors include, but are not limited to, salts, minerals, metabolites or any combination thereof. Exemplary salts, minerals, and metabolites include, but are not limited to, HEPES, Nicotinamide, Heparin, Sodium Pyruvate, L-Glutamine, MEM Non-Essential Amino Acid Solution, Ascorbic Acid, Nucleosides, FBS/FCS, Human serum, serum-substitute, anti-biotics, pH adjusters, Earle's Salts, 2-Mercaptoethanol, Human transferrin, Recombinant human insulin, Human serum albumin, Nucleofector PLUS Supplement, KCL, MgCl2, Na2HPO4, NAH2PO4, Sodium lactobionate, Manitol, Sodium succinate, Sodium Chloride, ClNa, Glucose, Ca(NO3)2, Tris/HC, K2HPO4, KH2PO4, Polyethylenimine, Poly-ethylene-glycol, Poloxamer 188, Poloxamer 181, Poloxamer 407, Poly-vinylpyrrolidone, Pop313, Crown-5, and any combination thereof. Exemplary supplemental factors include, but are not limited to, media such as PBS, HBSS, OptiMEM, DMEM, RPMI 1640, AIM-V, X-VIVO 15, CellGro DC Medium, CTS OpTimizer T Cell Expansion SFM, TexMACS Medium, PRIME-XV T Cell Expansion Medium ImmunoCult-XF T Cell Expansion Medium and any combination thereof. Exemplary supplemental factors include, but are not limited to, inhibitors of cellular DNA sensing, metabolism, differentiation, signal transduction, the apoptotic pathway and combinations thereof. Exemplary inhibitors include, but are not limited to, inhibitors of TLR9, MyD88, IRAK, TRAF6, TRAF3, IRF-7, NF-KB, Type 1 Interferons, proinflammatory cytokines, cGAS, STING, Sec5, TBK1, IRF-3, RNA pol III, RIG-1, IPS-1, FADD, RIP1, TRAF3, AIM2, ASC, Caspasel, Pro-L1B, PI3K, Akt, Wnt3A, inhibitors of glycogen synthase kinase-30 (GSK-3 β) (e.g. TWS119), Bafilomycin, Chloroquine. Quinacrine, AC-YVAD-CMK, Z-VAD-FMK, Z-ETD-FMK and any combination thereof. Exemplary supplemental factors include, but are not limited to, reagents that modify or stabilize one or more nucleic acids in a way to enhance cellular delivery, enhance nuclear delivery or transport, enhance the facilitated transport of nucleic acid into the nucleus, enhance degradation of epi-chromosomal nucleic acid, and/or decrease DNA-mediated toxicity. Exemplary reagents that modify or stabilize one or more nucleic acids include, but are not limited to, pH modifiers, DNA-binding proteins, lipids, phospholipids, CaPO4, net neutral charge DNA binding peptides with or without NLS sequences. TREX1 enzyme, and any combination thereof.

- (548) Transposition reagents, including a transposon and a transposase, may be added to a nucleofection reaction of the disclosure prior to, simultaneously with, or after an addition of cells to a nucleofection buffer (optionally, contained within a nucleofection reaction vial or cuvette). Transposons of the disclosure may comprise plasmid DNA, nanoplasmid, linearized plasmid DNA, a PCR product, DOGGYBONE<sup>TM</sup> DNA, an mRNA template, a single or double-stranded DNA, a protein-nucleic acid combination or any combination thereof. Transposons of the disclosure may comprised one or more sequences that encode one or more TTAA site(s), one or more inverted terminal repeat(s)(ITRs), one or more long terminal repeat(s) (LTRs), one or more insulator(s), one or more promotor(s), one or more full-length or truncated gene(s), one or more polyA signal(s), one or more self-cleaving 2A peptide cleavage site(s), one or more internal ribosome entry site(s) (IRES), one or more enhancer(s), one or more regulator(s), one or more replication origin(s), and any combination thereof.
- (549) Transposons of the disclosure may comprise one or more sequences that encode one or more full-length or truncated gene(s). Full-length and/or truncated gene(s) introduced by transposons of the disclosure may encode one or more of a signal peptide, a Centyrin, a single chain variable fragment (scFv), a hinge, a transmembrane domain, a costimulatory domain, a chimeric antigen receptor (CAR), a chimeric T-cell receptor (CAR-T), a CARTyrin (a CAR-T comprising a Centyrin), a receptor, a ligand, a cytokine, a drug resistance gene, a tumor antigen, an allo or auto antigen, an enzyme, a protein, a peptide, a poly-peptide, a fluorescent protein, a mutein or any combination thereof.
- (550) Transposons of the disclosure may be prepared in water, TAE, TBE, PBS, HBSS, media, a supplemental factor of the disclosure or any combination thereof.
- (551) Transposons of the disclosure may be designed to optimize clinical safety and/or improve manufacturability. As a non-limiting example, transposons of the disclosure may be designed to optimize clinical safety and/or improve manufacturability by eliminating unnecessary sequences or regions and/or including a non-antibiotic selection marker. Transposons of the disclosure may or may not be GMP grade. (552) Transposase enzymes of the disclosure may be encoded by one or more sequences of plasmid DNA, nanoplasmid DNA, mRNA, protein, protein-nucleic acid combination or any combination thereof. (553) Transposase enzymes of the disclosure may be prepared in water. TAE, TBE, PBS, HBSS, media, a supplemental factor of the disclosure or any combination thereof. Transposase enzymes of the disclosure or the sequences/constructs encoding or delivering them may or may not be GMP grade.
- (554) Transposons and transposase enzymes of the disclosure may be delivered to a cell by am means. (555) Although compositions and methods of the disclosure include delivery of a transposon and/or transposase of the disclosure to a cell by plasmid DNA (pDNA) or nanoplasmid DNA, the use of a plasmid or a nanoplasmid for delivery may allow the transposon and/or transposase to be integrated into the chromosomal DNA of the cell, which may lead to continued transposase expression. Accordingly,

transposon and/or transposase enzymes of the disclosure may be delivered to a cell as either mRNA or protein to remove any possibility for chromosomal integration.

(556) Transposons and transposases of the disclosure may be pre-incubated alone or in combination with one another prior to the introduction of the transposon and/or transposase into a nucleofection reaction. The absolute amounts of each of the transposon and the transposase, as well as the relative amounts, e.g., a ratio of transposon to transposase may be optimized.

(557) Following preparation of nucleofection reaction, optionally, in a vial or cuvette, the reaction may be loaded into a nucleofector apparatus and activated for delivery of an electric pulse according to the manufacturer's protocol. Electric pulse conditions used for delivery of a transposon and/or a transposase of the disclosure (or a sequence encoding a transposon and/or a transposase of the disclosure) to a cell may be optimized for yielding cells with enhanced viability, higher nucleofection efficiency, greater viability post-nucleofection, desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. When using Amaxa nucleofector technology, each of the various nucleofection programs for the Amaxa 2B or 4D nucleofector are contemplated.

(558) Following a nucleofection reaction of the disclosure, cells may be gently added to a cell medium. For example, when T cells undergo the nucleofection reaction, the T cells may be added to a T cell medium. Post-nucleofection cell media of the disclosure may comprise any one or more commerciallyavailable media. Post-nucleofection cell media of the disclosure (including post-nucleofection T cell media of the disclosure) may be optimized to yield cells with greater viability, higher nucleofection efficiency, exhibit greater viability post-nucleofection, display a more desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. Post-nucleofection cell media of the disclosure (including post-nucleofection T cell media of the disclosure) may comprise PBS, HBSS, OptiMEM, DMEM, RPMI 1640, AIM-V, X-VIVO 15, CellGro DC Medium, CTS OpTimizer T Cell Expansion SFM, TexMACS Medium. PRIME-XV T Cell Expansion Medium, ImmunoCult-XF T Cell Expansion Medium and any combination thereof. Post-nucleofection cell media of the disclosure (including post-nucleofection T cell media of the disclosure) may comprise one or more supplemental factors of the disclosure to enhance viability, nucleofection efficiency, viability post-nucleofection, cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. Exemplary supplemental factors include, but are not limited to, recombinant human cytokines, chemokines, interleukins and any combination thereof. Exemplary cytokines, chemokines, and interleukins include, but are not limited to, IL2, IL7, IL12, IL15, IL21, IL1, IL3, IL4, IL5, IL6, IL8, CXCL8, IL9, IL10, IL11, IL13, IL14, IL16, IL17, IL18, IL19, IL20, IL22, IL23, IL25, IL26, IL27, IL28, 1L29, IL30, IL31, IL32, IL33, IL35, IL36, GM-CSF, IFN-gamma, IL-1 alpha/IL-1F1, IL-1 beta IL-1F2, IL-12 p70, IL-12/IL-35 p35, IL-13, IL-17, IL-17A, IL-17A/F Heterodimer, IL-17F, IL-18/IL-1F4, IL-23, IL-24, IL-32, IL-32 beta, IL-32 gamma, IL-33, LAP (TGF-beta 1), Lymphotoxin-alpha-TNF-beta, TGF-beta, TNF-alpha. TRANCE/TNFSF11/RANK L and any combination thereof. Exemplary supplemental factors include, but are not limited to, salts, minerals, metabolites or any combination thereof. Exemplary salts, minerals, and metabolites include, but are not limited to, HEPES, Nicotinamide, Heparin, Sodium Pyruvate, L-Glutamine, MEM Non-Essential Amino Acid Solution, Ascorbic Acid, Nucleosides, FBS/FCS, Human serum, serum-substitute, antibiotics, pH adjusters, Earle's Salts, 2-Mercaptoethanol, Human transferrin, Recombinant human insulin, Human serum albumin, Nucleofector PLUS Supplement, KCL, MgCl2, Na2HPO4, NAH2PO4, Sodium lactobionate, Manitol, Sodium succinate, Sodium Chloride. ClNa, Glucose, Ca(NO3)2, Tris/HCl, K2HPO4, KH2PO4, Polyethylenimine, Poly-ethylene-glycol, Poloxamer 188, Poloxamer 181, Poloxamer 407, Poly-vinylpyrrolidone, Pop313, Crown-5, and any combination thereof. Exemplary supplemental factors include, but are not limited to, media such as PBS, HBSS, OptiMEM, DMEM, RPMI 1640, AIM-V, X-VIVO 15, CellGro DC Medium, CTS OpTimizer T Cell Expansion SFM. TexMACS Medium, PRIME-XV T Cell Expansion Medium, ImmunoCult-XF T Cell Expansion Medium and any combination thereof. Exemplary supplemental factors include, but are not limited to, inhibitors of cellular DNA sensing, metabolism, differentiation, signal transduction, the apoptotic pathway and combinations thereof. Exemplary inhibitors include, but are not limited to, inhibitors of TLR9, MyD88, IRAK, TRAF6, TRAF3, IRF-7, NF-KB, Type 1 Interferons, pro-inflammatory cytokines, cGAS, STING, Sec5, TBK1, IRF-3, RNA pol III, RIG-1, IPS-1, FADD, RIP1, TRAF3, AIM2, ASC, Caspasel, Pro-IL1B, PI3K, Akt, Wnt3A, inhibitors of glycogen synthase kinase-3β(GSK-3 β) (e.g. TWS119), Bafilomycin, Chloroquine,

Quinacrine, AC-YVAD-CMK, Z-VAD-FMK, Z-IETD-FMK and any combination thereof. Exemplary supplemental factors include, but are not limited to, reagents that modify or stabilize one or more nucleic acids in a way to enhance cellular delivery, enhance nuclear delivery or transport, enhance the facilitated transport of nucleic acid into the nucleus, enhance degradation of epi-chromosomal nucleic acid, and/or decrease DNA-mediated toxicity. Exemplary reagents that modify or stabilize one or more nucleic acids include, but are not limited to, pH modifiers, DNA-binding proteins, lipids, phospholipids, CaPO4, net neutral charge DNA binding peptides with or without NLS sequences, TREX1 enzyme, and any combination thereof.

- (559) Post-nucleofection cell media of the disclosure (including post-nucleofection T cell media of the disclosure) may be used at room temperature or pre-warmed to, for example to between 32° C. to 37° C., inclusive of the endpoints. Post-nucleofection cell media of the disclosure (including post-nucleofection T cell media of the disclosure) may be pre-warmed to any temperature that maintains or enhances cell viability and/or expression of a transposon or portion thereof of the disclosure.
- (560) Post-nucleofection cell media of the disclosure (including post-nucleofection T cell media of the disclosure) may be contained in tissue culture flasks or dishes, G-Rex flasks. Bioreactor or cell culture bags, or any other standard receptacle. Post-nucleofection cell cultures of the disclosure (including post-nucleofection T cell cultures of the disclosure) may be may be kept still, or, alternatively, they may be perturbed (e.g. rocked, swirled, or shaken).
- (561) Post-nucleofection cell cultures may comprise genetically-modified cells Post-nucleofection T cell cultures may comprise genetically-modified T cells. Genetically modified cells of the disclosure may be either rested for a defined period of time or stimulated for expansion by, for example, the addition of a T Cell Expander technology. In certain embodiments, genetically modified cells of the disclosure may be either rested for a defined period of time or immediately stimulated for expansion by, for example, the addition of a T Cell Expander technology. Genetically modified cells of the disclosure may be rested to allow them sufficient time to acclimate, time for transposition to occur, and/or time for positive or negative selection, resulting in cells with enhanced viability, higher nucleofection efficiency, greater viability post-nucleofection, desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. Genetically modified cells of the disclosure may be rested, for example, for 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or more hours. In certain embodiments, genetically modified cells of the disclosure may be rested, for example, for an overnight. In certain aspects, an overnight is about 12 hours. Genetically modified cells of the disclosure may be rested, for example, for 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or more days.
- (562) Genetically modified cells of the disclosure may be selected following a nucleofection reaction and prior to addition of an expander technology. For optimal selection of genetically-modified cells, the cells may be allowed to rest in a post-nucleofection cell medium for at least 2-14 days to facilitate identification of modified cells (e.g., differentiation of modified from non-modified cells).
- (563) As early as 24-hours post-nucleofection, expression of a CAR/CARTyrin and selection marker of the disclosure may be detectable in modified T cells upon successful nucleofection of a transposon of the disclosure. Due to epi-chromosomal expression of the transposon, expression of a selection marker alone may not differentiate modified T cells (those cells in which the transposon has been successfully integrated) from unmodified T cells (those cells in which the transposon was not successfully integrated). When epi-chromosomal expression of the transposon obscures the detection of modified cells by the selection marker, the nucleofected cells (both modified and unmodified cells) may be rested for a period of time (e.g. 2-14 days) to allow the cells to cease expression or lose all epi-chromosomal transposon expression. Following this extended resting period, only modified T cells should remain positive for expression of selection marker. The length of this extended resting period may be optimized for each nucleofection reaction and selection process. When epi-chromosomal expression of the transposon obscures the detection of modified cells by the selection marker, selection may be performed without this extended resting period, however, an additional selection step may be included at a later time point (e.g. either during or after the expansion stage).
- (564) Selection of genetically modified cells of the disclosure may be performed by any means. In certain embodiments of the methods of the disclosure, selection of genetically modified cells of the disclosure may be performed by isolating cells expressing a specific selection marker. Selection markers of the

disclosure may be encoded by one or more sequences in the transposon. Selection markers of the disclosure may be expressed by the modified cell as a result of successful transposition (i.e., not encoded by one or more sequences in the transposon). In certain embodiments, genetically modified cells of the disclosure contain a selection marker that confers resistance to a deleterious compound of the postnucleofection cell medium. The deleterious compound may comprise, for example, an antibiotic or a drug that, absent the resistance conferred by the selection marker to the modified cells, would result in cell death. Exemplary selection markers include, but are not limited to, wild type (WT) or mutant forms of one or more of the following genes: neo, DHFR, TYMS, ALDH, MDR1, MGMT, FANCF, RAD51C, GCS, and NKX2.2. Exemplary selection markers include, but are not limited to, a surface-expressed selection marker or surface-expressed tag may be targeted by Ab-coated magnetic bead technology or column selection, respectively. A cleavable tag such as those used in protein purification may be added to a selection marker of the disclosure for efficient column selection, washing, and elution. In certain embodiments, selection markers of the disclosure are not expressed by the modified cells (including modified T cells) endogenously and, therefore, may be useful in the physical isolation of modified cells (by, for example, cell sorting techniques). Exemplary selection markers of the disclosure are not expressed by the modified cells (including modified T cells) endogenously include, but are not limited to, full-length, mutated, or truncated forms of CD271, CD19 CD52, CD34, RQR8, CD22, CD20, CD33 and any combination thereof.

(565) Genetically modified cells of the disclosure may be selective expanded following a nucleofection reaction. In certain embodiments, modified T cells comprising a CAR/CARTyrin may be selectively expanded by CAR/CARTyrin stimulation. Modified T cells comprising a CAR/CARTyrin may be stimulated by contact with a target-covered reagent (e.g. a tumor line or a normal cell line expressing a target or expander beads covered in a target). Alternatively, modified T cells comprising a CAR/CARTyrin may be stimulated by contact with an irradiated tumor cell, an irradiated allogeneic normal cell, an irradiated autologous PBMC. To minimize contamination of cell product compositions of the disclosure with a target-expressing cell used for stimulation, for example, when the cell product composition may be administered directly to a subject, the stimulation may be performed using expander beads coated with CAR/CARTyrin target protein. Selective expansion of modified T cells comprising a CAR/CARTyrin by CAR/CARTyrin stimulation may be optimized to avoid functionally-exhausting the modified T-cells. (566) Selected genetically-modified cells of the disclosure may be cryopreserved, rested for a defined period of time, or stimulated for expansion by the addition of a Cell Expander technology. Selected genetically-modified cells of the disclosure may be cryopreserved, rested for a defined period of time, or immediately stimulated for expansion by the addition of a Cell Expander technology. When the selected genetically-modified cells are T cells, the T cells may be stimulated for expansion by the addition of a T-Cell Expander technology. Selected genetically modified cells of the disclosure may be rested, for example, for 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or more hours. In certain embodiments, selected genetically modified cells of the disclosure may be rested, for example, for an overnight. In certain aspects, an overnight is about 12 hours. Selected genetically modified cells of the disclosure may be rested, for example, for 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or more days. Selected genetically modified cells of the disclosure may be rested for any period of time resulting in cells with enhanced viability, higher nucleofection efficiency, greater viability post-nucleofection, desirable cell phenotype, and/or greater/faster expansion upon addition of expansion technologies. (567) Selected genetically-modified cells (including selected genetically-modified T cells of the disclosure) may be cryopreserved using any standard cryopreservation method, which may be optimized for storing and/or recovering human cells with high recovery, viability, phenotype, and/or functional capacity. Cryopreservation methods of the disclosure may include commercially-available cryopreservation media and/or protocols.

(568) A transposition efficiency of selected genetically-modified cells (including selected genetically-modified T cells of the disclosure) may be assessed by any means. For example, prior to the application of an expander technology, expression of the transposon by selected genetically-modified cells (including selected genetically-modified T cells of the disclosure) may be measured by fluorescence-activated cell sorting (FACS). Determination of a transposition efficiency of selected genetically-modified cells (including selected genetically-modified T cells of the disclosure) may include determining a percentage

- of selected cells expressing the transposon (e.g. a CAR). Alternatively, or in addition, a purity of T cells, a Mean Fluorescence Intensity (MFI) of the transposon expression (e.g. CAR expression), an ability of a CAR (delivered in the transposon) to mediate degranulation and/or killing of a target cell expressing the CAR ligand, and/or a phenotype of selected genetical-modified cells (including selected genetically-modified T cells of the disclosure) may be assessed by any means.
- (569) Cell product compositions of the disclosure may be released for administration to a subject upon meeting certain release criteria Exemplary release criteria may include, but are not limited to, a particular percentage of modified, selected and/or expanded T cells expressing detectable levels of a CAR on the cell surface.
- (570) Genetic Modification of an Autologous T Cell Product Composition
- (571) Genetically-modified cells (including genetically-modified T cells) of the disclosure may be expanded using an expander technology. Expander technologies of the disclosure may comprise a commercially-available expander technology. Exemplary expander technologies of the disclosure include stimulation a genetically-modified T cell of the disclosure via the TCR While all means for stimulation of a genetically-modified T cell of the disclosure are contemplated, stimulation a genetically-modified T cell of the disclosure via the TCR is a preferred method, yielding a product with a superior level of killing capacity.
- (572) To stimulate a genetically-modified T cell of the disclosure via the TCR, Thermo Expander DynaBeads may be used at a 3:1 bead to T cell ratio. If the expander beads are not biodegradable, the beads may be removed from the expander composition. For example, the beads may be removed from the expander composition after about 5 days. To stimulate a genetically-modified T cell of the disclosure via the TCR, a Miltenyi T Cell Activation/Expansion Reagent may be used. To stimulate a genetically-modified T cell of the disclosure via the TCR, StemCell Technologies' ImmunoCult Human CD3/CD28 or CD3/CD28/CD2 T Cell Activator Reagent may be used. This technology may be preferred since the soluble tetrameric antibody complexes would degrade after a period and would not require removal from the process.
- (573) Artificial antigen presenting cells (APCs) may be engineered to co-express the target antigen and may be used to stimulate a cell or T-cell of the disclosure through a TCR and/or CAR of the disclosure. Artificial APCs may comprise or may be derived from a tumor cell line (including, for example, the immortalized myelogenous leukemia line K562) and may be engineered to co-express multiple costimulatory molecules or technologies (such as CD28, 4-1BBL, CD64, mbIL-21, mbIL-15, CAR target molecule, etc.). When artificial APCs of the disclosure are combined with costimulatory molecules, conditions may be optimized to prevent the development or emergence of an undesirable phenotype and functional capacity, namely terminally-differentiated effector T cells.
- (574) Irradiated PBMCs (auto or allo) may express some target antigens, such as CD19, and may be used to stimulate a cell or T-cell of the disclosure through a TCR and/or CAR of the disclosure. Alternatively, or in addition, irradiated tumor cells may express some target antigens and may be used to stimulate a cell or T-cell of the disclosure through a TCR and/or CAR of the disclosure.
- (575) Plate-bound and/or soluble anti-CD3, anti-CD2 and/or anti-CD28 stimulate may be used to stimulate a cell or T-cell of the disclosure through a TCR and/or CAR of the disclosure.
- (576) Antigen-coated beads may display target protein and may be used to stimulate a cell or T-cell of the disclosure through a TCR and/or CAR of the disclosure. Alternatively, or in addition, expander beads coated with a CAR/CARTyrin target protein may be used to stimulate a cell or T-cell of the disclosure through a TCR and/or CAR of the disclosure.
- (577) Expansion methods drawn to stimulation of a cell or T-cell of the disclosure through the TCR or CAR/CARTyrin and via surface-expressed CD2, CD3, CD28, 4-1BB, and/or other markers on genetically-modified T cells.
- (578) An expansion technology may be applied to a cell of the disclosure immediately post-nucleofection until approximately 24 hours post-nucleofection. While various cell media may be used during an expansion procedure, a desirable T Cell Expansion Media of the disclosure may yield cells with, for example, greater viability, cell phenotype, total expansion, or greater capacity for in vivo persistence, engraftment, and/or CAR-mediated killing. Cell media of the disclosure may be optimized to improve/enhance expansion, phenotype, and function of genetically-modified cells of the disclosure. A

preferred phenotype of expanded T cells may include a mixture of T stem cell memory, T central, and T effector memory cells. Expander Dynabeads may yield mainly central memory T cells which may lead to superior performance in the clinic.

(579) Exemplary T cell expansion media of the disclosure may include, in part or in total, PBS, HBSS, OptiMEM, DMEM, RPMI 1640, AIM-V, X-VIVO 15, CellGro DC Medium. CTS OpTimizer T Cell Expansion SFM. TexMACS Medium, PRIME-XV T Cell Expansion Medium, ImmunoCult-XF T Cell Expansion Medium, or any combination thereof. T cell expansion media of the disclosure may further include one or more supplemental factors. Supplemental factors that may be included in a T cell expansion media of the disclosure enhance viability, cell phenotype, total expansion, or increase capacity for in vivo persistence, engraftment, and/or CAR-mediated killing. Supplemental factors that may be included in a T cell expansion media of the disclosure include, but are not limited to, recombinant human cytokines, chemokines, and/or interleukins such as IL2, IL7, IL12, IL15, IL21, IL1, IL3, IL4, IL5, IL6, IL8, CXCL8, IL9, IL10, IL11, IL13, IL14, IL16, IL17, IL18, IL19, IL20, IL22, IL23, IL25, IL26, IL27, IL28, IL29, IL30, IL31, IL32, IL33, IL35, IL36, GM-CSF, IFN-gamma, IL-1 alpha/IL-1F1, IL-1 beta/IL-1F2, IL-12 p70, IL-12/IL-35 p35, IL-13, IL-17/IL-17A, IL-17A/F Heterodimer, IL-17F, IL-18/IL-1F4, IL-23, IL-24, IL-32, IL-32 beta, IL-32 gamma. IL-33, LAP (TGF-beta 1), Lymphotoxin-alpha/TNF-beta, TGF-beta, TNF-alpha, TRANCE/TNFSF11/RANK L, or any combination thereof. Supplemental factors that may be included in a T cell expansion media of the disclosure include, but are not limited to, salts, minerals, and/or metabolites such as HEPES, Nicotinamide, Heparin, Sodium Pyruvate, L-Glutamine, MEM Non-Essential Amino Acid Solution, Ascorbic Acid, Nucleosides, FBS/FCS, Human serum, serum-substitute, anti-biotics, pH adjusters, Earle's Salts, 2-Mercaptoethanol, Human transferrin, Recombinant human insulin, Human serum albumin, Nucleofector PLUS Supplement, KCL, MgCl2, Na2HPO4, NAH2PO4, Sodium lactobionate, Manitol, Sodium succinate, Sodium Chloride, ClNa, Glucose, Ca(NO3)2, Tris/HC, K2HPO4, KH2PO4, Polyethylenimine, Poly-ethylene-glycol, Poloxamer 188, Poloxamer 181, Poloxamer 407, Poly-vinylpyrrolidone, Pop313, Crown-5 or any combination thereof. Supplemental factors that may be included in a T cell expansion media of the disclosure include, but are not limited to, inhibitors of cellular DNA sensing, metabolism, differentiation, signal transduction, and/or the apoptotic pathway such as inhibitors of TLR9, MyD88, IRAK, TRAF6, TRAF3, IRF-7, NF-KB, Type 1 Interferons, proinflammatory cytokines, cGAS, STING, Sec5, TBK1, IRF-3, RNA pol III, RIG-1, IPS-1, FADD, RIP1, TRAF3, AIM2, ASC, Caspasel, Pro-IL1B, PI3K Akt, Wnt3A, inhibitors of glycogen synthase kinase-3β (GSK-3 β) (e.g. TWS119), Bafilomycin, Chloroquine, Quinacrine, AC-YVAD-CMK, Z-VAD-FMK, Z-IETD-FMK, or any combination thereof. (580) Supplemental factors that may be included in a T cell expansion media of the disclosure include, but

are not limited to, reagents that modify or stabilize nucleic acids in a way to enhance cellular delivery, enhance nuclear delivery or transport, enhance the facilitated transport of nucleic acid into the nucleus, enhance degradation of epi-chromosomal nucleic acid, and/or decrease DNA-mediated toxicity, such as pH modifiers, DNA-binding proteins, lipids, phospholipids, CaPO4, net neutral charge DNA binding peptides with or without NLS sequences. TREX1 enzyme, or any combination thereof. (581) Genetically-modified cells of the disclosure may be selected during the expansion process by the use of selectable drugs or compounds. For example, in certain embodiments, when a transposon of the disclosure may encode a selection marker that confers to genetically-modified cells resistance to a drug added to the culture medium, selection may occur during the expansion process and may require approximately 1-14 days of culture for selection to occur. Examples of drug resistance genes that may be used as selection markers encoded by a transposon of the disclosure, include, but are not limited to, wild type (WT) or mutant forms of the genes neo, DHFR, TYMS, ALDH, MDR1, MGMT, FANCF, RAD51C, GCS, NKX2.2, or any combination thereof. Examples of corresponding drugs or compounds that may be added to the culture medium to which a selection marker may confer resistance include, but are not limited to, G418, Puromycin, Ampicillin, Kanamycin, Methotrexate, Mephalan, Temozolomide, Vincristine, Etoposide, Doxorubicin, Bendamustine, Fludarabine, Aredia (Pamidronate Disodium), Becenum (Carmustine), BiCNU (Carmustine), Bortezomib, Carfilzomib, Carmubris (Carmustine), Carmustine, Clafen (Cyclophosphamide), Cyclophosphamide, Cytoxan (Cyclophosphamide), Daratumumab, Darzalex (Daratumumab), Doxil (Doxorubicin Hydrochloride Liposome), Doxorubicin Hydrochloride Liposome,

Dox-SL (Doxorubicin Hydrochloride Liposome), Elotuzumab, Empliciti (Elotuzumab), Evacet

(Doxorubicin Hydrochloride Liposome), Farydak (Panobinostat), ixazomib Citrate, Kyprolis (Carfilzomib), Lenalidomide, LipoDox (Doxorubicin Hydrochloride Liposome), Mozobil (Plerixafor), Neosar (Cyclophosphamide), Ninlaro (Ixazomib Citrate), Pamidronate Disodium, Panobinostat, Plerixafor, Pomalidomide, Pomalyst (Pomalidomide), Revlimid (Lenalidomide), Synovir (Thalidomide), Thalidomide, Thalomid (Thalidomide), Velcade (Bortezomib), Zoledronic Acid, Zometa (Zoledronic Acid), or any combination thereof.

- (582) A T-Cell Expansion process of the disclosure may occur in a cell culture bag in a WAVE Bioreactor, a G-Rex flask, or in any other suitable container and/or reactor.
- (583) A cell or T-cell culture of the disclosure may be kept steady, rocked, swirled, or shaken.
- (584) A cell or T-cell expansion process of the disclosure may optimize certain conditions, including, but not limited to culture duration, cell concentration, schedule for T cell medium addition/removal, cell size, total cell number, cell phenotype, purity of cell population, percentage of genetically-modified cells in growing cell population, use and composition of supplements, the addition/removal of expander technologies, or any combination thereof.
- (585) A cell or T-cell expansion process of the disclosure may continue until a predefined endpoint prior to formulation of the resultant expanded cell population. For example, a cell or T-cell expansion process of the disclosure may continue for a predetermined amount of time: at least, 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24 hours; at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 days; at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12 weeks; at least 1, 2, 3, 4, 5, 6, months, or at least 1 year. A cell or T-cell expansion process of the disclosure may continue until the resultant culture reaches a predetermined overall cell density: 1, 10, 100, 1000, 104, 105, 106, 107, 108, 109, 1010 cells per volume (µl, ml, L) or any density in between. A cell or T-cell expansion process of the disclosure may continue until the genetically-modified cells of a resultant culture demonstrate a predetermined level of expression of a transposon of the disclosure: 1%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, or 100% or any percentage in between of a threshold level of expression (a minimum, maximum or mean level of expression indicating the resultant genetically-modified cells are clinically-efficacious). A cell or T-cell expansion process of the disclosure may continue until the proportion of genetically-modified cells of a resultant culture to the proportion of unmodified cells reaches a predetermined threshold: at least 1:10, 1:9, 1:8, 1:7, 1:6, 1:5, 1:4, 1:3, 1:2, 1:1, 2:1, 2:1, 4:1, 5:1, 6:1, 7:1, 8:1, 9:1 10:1 or any ratio in between. (586) Analysis of Genetically-Modified Autologous T Cells for Release
- (587) A percentage of genetically-modified cells may be assessed during or after an expansion process of the disclosure. Cellular expression of a transposon by a genetically-modified cell of the disclosure may be measured by fluorescence-activated cell sorting (FACS) For example, FACS may be used to determine a percentage of cells or T cells expressing a CAR of the disclosure. Alternatively, or in addition, a purity of genetically-modified cells or T cells, the Mean Fluorescence Intensity (MFI) of a CAR expressed by a genetically-modified cell or T cell of the disclosure, an ability of the CAR to mediate degranulation and/or killing of a target cell expressing the CAR ligand, and/or a phenotype of CAR+ T cells may be assessed. (588) Compositions of the disclosure intended for administration to a subject may be required to meet one or more "release criteria" that indicate that the composition is safe and efficacious for formulation as a pharmaceutical product and/or administration to a subject. Release criteria may include a requirement that a composition of the disclosure (e.g. a T-cell product of the disclosure) comprises a particular percentage of T cells expressing detectable levels of a CAR of the disclosure on their cell surface.
- (589) The expansion process should be continued until a specific criterion has been met (e.g. achieving a certain total number of cells, achieving a particular population of memory cells, achieving a population of a specific size).
- (590) Certain criterion signal a point at which the expansion process should end. For example, cells should be formulated, reactivated, or cryopreserved once they reach a cell size of 300 fL (otherwise, cells reaching a size above this threshold may start to die). Cryopreservation immediately once a population of cells reaches an average cell size of less than 300 fL may yield better cell recovery upon thawing and culture because the cells haven't yet reached a fully quiescent state prior to cryopreservation (a fully quiescent size is approximately 180 fL). Prior to expansion, T cells of the disclosure may have a cell size of about 180 f, but may more than quadruple their cell size to approximately 900 fL at 3 days post-expansion. Over the next 6-12 days, the population of T-cells will slowly decrease cell size to full

quiescence at 180 f.

(591) A process for preparing a cell population for formulation may include, but is not limited to the steps of, concentrating the cells of the cell population, washing the cells, and/or further selection of the cells via drug resistance or magnetic bead sorting against a particular surface-expressed marker. A process for preparing a cell population for formulation may further include a sorting step to ensure the safety and purity of the final product. For example, if a tumor cell from a patient has been used to stimulate a genetically-modified T-cell of the disclosure or that have been genetically-modified in order to stimulate a genetically-modified T-cell of the disclosure that is being prepared for formulation, it is critical that no tumor cells from the patient are included in the final product.

- (592) Cell Product Infusion and/or Cryopreservation for Infusion
- (593) A pharmaceutical formulation of the disclosure may be distributed into bags for infusion, cryopreservation, and/or storage.
- (594) A pharmaceutical formulation of the disclosure may be cryopreserved using a standard protocol and, optionally, an infusible cryopreservation medium. For example, a DMSO free cryopreservant (e.g. CryoSOfree™, DMSO-free Cryopreservation Medium) may be used to reduce freezing-related toxicity. A cryopreserved pharmaceutical formulation of the disclosure may be stored for infusion to a patient at a later date. An effective treatment may require multiple administrations of a pharmaceutical formulation of the disclosure and, therefore, pharmaceutical formulations may be packaged in pre-aliquoted "doses" that may be stored frozen but separated for thawing of individual doses.
- (595) A pharmaceutical formulation of the disclosure may be stored at room temperature. An effective treatment may require multiple administrations of a pharmaceutical formulation of the disclosure and, therefore, pharmaceutical formulations may be packaged in pre-aliquoted "doses" that may be stored together but separated for administration of individual doses.
- (596) A pharmaceutical formulation of the disclosure may be archived for subsequent re-expansion and/or selection for generation of additional doses to the same patient in the case of an allogenic therapy who may need an administration at a future date following, for example, a remission and relapse of a condition. (597) Formulations
- (598) As noted above, the disclosure provides for stable formulations, which preferably comprise a phosphate buffer with saline or a chosen salt, as well as preserved solutions and formulations containing a preservative as well as multi-use preserved formulations suitable for pharmaceutical or veterinary use, comprising at least one protein scaffold in a pharmaceutically acceptable formulation. Preserved formulations contain at least one known preservative or optionally selected from the group consisting of at least one phenol, m-cresol, p-cresol, o-cresol, chlorocresol, benzyl alcohol, phenylmercuric nitrite, phenoxyethanol, formaldehyde, chlorobutanol, magnesium chloride (e.g., hexahydrate), alkylparaben (methyl, ethyl, propyl, butyl and the like), benzalkonium chloride, benzethonium chloride, sodium dehydroacetate and thimerosal, polymers, or mixtures thereof in an aqueous diluent. Any suitable concentration or mixture can be used as known in the art, such as about 0.0015%, or any range, value, or fraction therein. Non-limiting examples include, no preservative, about 0.1-2% m-cresol (e.g., 0.2, 0.3, 0.4, 0.5, 0.9, 1.0%), about 0.1-3% benzyl alcohol (e.g., 0.5, 0.9, 1.1, 1.5, 1.9, 2.0, 2.5%), about 0.001-0.5% thimerosal (e.g., 0.005, 0.01), about 0.001-2.0% phenol (e.g., 0.05, 0.25, 0.28, 0.5, 0.9, 1.0%), 0.0005-1.0% alkylparaben(s) (e.g., 0.00075, 0.0009, 0.001, 0.002, 0.005, 0.0075, 0.009, 0.01, 0.2.0.05, 0.075, 0.09, 0.1, 0.2, 0.3, 0.5, 0.75, 0.9, 1.0%), and the like.
- (599) As noted above, the invention provides an article of manufacture, comprising packaging material and at least one vial comprising a solution of at least one protein scaffold with the prescribed buffers and/or preservatives, optionally in an aqueous diluent, wherein said packaging material comprises a label that indicates that such solution can be held over a period of 1, 2, 3, 4, 5, 6, 9, 12, 18, 20, 24, 30, 36, 40, 48, 54, 60, 66, 72 hours or greater. The invention further comprises an article of manufacture, comprising packaging material, a first vial comprising lyophilized at least one protein scaffold, and a second vial comprises a label that instructs a patient to reconstitute the at least one protein scaffold in the aqueous diluent to form a solution that can be held over a period of twenty-four hours or greater.
- (600) The at least one protein scaffold used in accordance with the present invention can be produced by recombinant means, including from mammalian cell or transgenic preparations, or can be purified from

other biological sources, as described herein or as known in the art.

(601) The range of at least one protein scaffold in the product of the present invention includes amounts yielding upon reconstitution, if in a wet/dry system, concentrations from about 1.0  $\mu$ g/ml to about 1000 mg/ml, although lower and higher concentrations are operable and are dependent on the intended delivery vehicle, e.g., solution formulations will differ from transdermal patch, pulmonary, transmucosal, or osmotic or micro pump methods.

(602) Preferably, the aqueous diluent optionally further comprises a pharmaceutically acceptable preservative. Preferred preservatives include those selected from the group consisting of phenol, m-cresol, p-cresol, o-cresol, chlorocresol, benzyl alcohol, alkylparaben (methyl, ethyl, propyl, butyl and the like), benzalkonium chloride, benzethonium chloride, sodium dehydroacetate and thimerosal, or mixtures thereof. The concentration of preservative used in the formulation is a concentration sufficient to yield an anti-microbial effect. Such concentrations are dependent on the preservative selected and are readily determined by the skilled artisan.

(603) Other excipients, e.g., isotonicity agents, buffers, antioxidants, and preservative enhancers, can be optionally and preferably added to the diluent. An isotonicity agent, such as glycerin, is commonly used at known concentrations. A physiologically tolerated buffer is preferably added to provide improved pH control. The formulations can cover a wide range of pHs, such as from about pH 4 to about pH 10, and preferred ranges from about pH 5 to about pH 9, and a most preferred range of about 6.0 to about 8.0. Preferably, the formulations of the present invention have a pH between about 6.8 and about 7.8. Preferred buffers include phosphate buffers, most preferably, sodium phosphate, particularly, phosphate buffered saline (PBS).

(604) Other additives, such as a pharmaceutically acceptable solubilizers like Tween 20 (polyoxyethylene (20) sorbitan monolaurate), Tween 40 (polyoxyethylene (20) sorbitan monopalmitate), Tween 80 (polyoxyethylene (20) sorbitan monooleate), Pluronic F68 (polyoxyethylene polyoxypropylene block copolymers), and PEG (polyethylene glycol) or non-ionic surfactants, such as polysorbate 20 or 80 or poloxamer 184 or 188, Pluronic® polyls, other block co-polymers, and chelators, such as EDTA and EGTA, can optionally be added to the formulations or compositions to reduce aggregation. These additives are particularly useful if a pump or plastic container is used to administer the formulation. The presence of pharmaceutically acceptable surfactant mitigates the propensity for the protein to aggregate. (605) The formulations of the present invention can be prepared by a process which comprises mixing at least one protein scaffold and a preservative selected from the group consisting of phenol, m-cresol, pcresol, o-cresol, chlorocresol, benzyl alcohol, alkylparaben, (methyl, ethyl, propyl, butyl and the like), benzalkonium chloride, benzethonium chloride, sodium dehydroacetate and thimerosal or mixtures thereof in an aqueous diluent. Mixing the at least one protein scaffold and preservative in an aqueous diluent is carried out using conventional dissolution and mixing procedures. To prepare a suitable formulation, for example, a measured amount of at least one protein scaffold in buffered solution is combined with the desired preservative in a buffered solution in quantities sufficient to provide the protein and preservative at the desired concentrations. Variations of this process would be recognized by one of ordinary skill in the art. For example, the order the components are added, whether additional additives are used, the temperature and pH at which the formulation is prepared, are all factors that can be optimized for the concentration and means of administration used.

(606) The claimed formulations can be provided to patients as clear solutions or as dual vials comprising a vial of lyophilized at least one protein scaffold that is reconstituted with a second vial containing water, a preservative and/or excipients, preferably, a phosphate buffer and/or saline and a chosen salt, in an aqueous diluent. Either a single solution vial or dual vial requiring reconstitution can be reused multiple times and can suffice for a single or multiple cycles of patient treatment and thus can provide a more convenient treatment regimen than currently available.

(607) The present claimed articles of manufacture are useful for administration over a period ranging from immediate to twenty-four hours or greater. Accordingly, the presently claimed articles of manufacture offer significant advantages to the patient. Formulations of the invention can optionally be safely stored at temperatures of from about 2° C. to about 40° C. and retain the biological activity of the protein for extended periods of time, thus allowing a package label indicating that the solution can be held and/or used over a period of 6, 12, 18, 24, 36, 48, 72, or 96 hours or greater. If preserved diluent is used, such

label can include use up to 1-12 months, one-half, one and a half, and/or two years.

(608) The solutions of at least one protein scaffold of the invention can be prepared by a process that comprises mixing at least one protein scaffold in an aqueous diluent. Mixing is carried out using conventional dissolution and mixing procedures. To prepare a suitable diluent, for example, a measured amount of at least one protein scaffold in water or buffer is combined in quantities sufficient to provide the protein and, optionally, a preservative or buffer at the desired concentrations. Variations of this process would be recognized by one of ordinary skill in the art. For example, the order the components are added, whether additional additives are used, the temperature and pH at which the formulation is prepared, are all factors that can be optimized for the concentration and means of administration used.

- (609) The claimed products can be provided to patients as clear solutions or as dual vials comprising a vial of lyophilized at least one protein scaffold that is reconstituted with a second vial containing the aqueous diluent. Either a single solution vial or dual vial requiring reconstitution can be reused multiple times and can suffice for a single or multiple cycles of patient treatment and thus provides a more convenient treatment regimen than currently available.
- (610) The claimed products can be provided indirectly to patients by providing to pharmacies, clinics, or other such institutions and facilities, clear solutions or dual vials comprising a vial of lyophilized at least one protein scaffold that is reconstituted with a second vial containing the aqueous diluent. The clear solution in this case can be up to one liter or even larger in size, providing a large reservoir from which smaller portions of the at least one protein scaffold solution can be retrieved one or multiple times for transfer into smaller vials and provided by the pharmacy or clinic to their customers and/or patients. (611) Recognized devices comprising single vial systems include pen-injector devices for delivery of a solution, such as BD Pens, BD Autojector®, Humaject®, NovoPen®, B-D® Pen, AutoPen®, and OptiPen®, GenotropinPen®, Genotronorm Pen®, Humatro Pen®, Reco-Pen®, Roferon Pen®, Biojector®, Iject®, J-tip Needle-Free Injector®, Intraject®, Medi-Ject®, e.g., as made or developed by Becton Dickinson (Franklin Lakes. N.J., www.bectondickenson.com), Disetronic (Burgdorf, Switzerland, www.disetronic.com, Bioject, Portland, Oreg. (www.bioject.com); National Medical Products, Weston Medical (Peterborough, UK, www.weston-medical.com), Medi-Ject Corp (Minneapolis, Minn., www.mediject.com), and similarly suitable devices. Recognized devices comprising a dual vial system include those pen-injector systems for reconstituting a lyophilized drug in a cartridge for delivery of the reconstituted solution, such as the HumatroPen®. Examples of other devices suitable include pre-filled syringes, auto-injectors, needle free injectors and needle free IV infusion sets.
- (612) The products presently claimed include packaging material. The packaging material provides, in addition to the information required by the regulatory agencies, the conditions under which the product can be used. The packaging material of the present invention provides instructions to the patient to reconstitute at least one protein scaffold in the aqueous diluent to form a solution and to use the solution over a period of 2-24 hours or greater for the two vial, wet/dry, product. For the single vial, solution product, the label indicates that such solution can be used over a period of 2-24 hours or greater. The presently claimed products are useful for human pharmaceutical product use.
- (613) The formulations of the present invention can be prepared by a process that comprises mixing at least one protein scaffold and a selected buffer, preferably, a phosphate buffer containing saline or a chosen salt. Mixing at least one protein scaffold and buffer in an aqueous diluent is carried out using conventional dissolution and mixing procedures. To prepare a suitable formulation, for example, a measured amount of at least one protein scaffold in water or buffer is combined with the desired buffering agent in water in quantities sufficient to provide the protein and buffer at the desired concentrations. Variations of this process would be recognized by one of ordinary skill in the art. For example, the order the components are added, whether additional additives are used, the temperature and pH at which the formulation is prepared, are all factors that can be optimized for the concentration and means of administration used.
- (614) The claimed stable or preserved formulations can be provided to patients as clear solutions or as dual vials comprising a vial of lyophilized protein scaffold that is reconstituted with a second vial containing a preservative or buffer and excipients in an aqueous diluent. Either a single solution vial or dual vial requiring reconstitution can be reused multiple times and can suffice for a single or multiple cycles of patient treatment and thus provides a more convenient treatment regimen than currently

available.

(615) Other formulations or methods of stabilizing the protein scaffold may result in other than a clear solution of lyophilized powder comprising the protein scaffold. Among non-clear solutions are formulations comprising particulate suspensions, said particulates being a composition containing the protein scaffold in a structure of variable dimension and known variously as a microsphere, microparticle, nanoparticle, nanosphere, or liposome. Such relatively homogenous, essentially spherical, particulate formulations containing an active agent can be formed by contacting an aqueous phase containing the active agent and a polymer and a nonaqueous phase followed by evaporation of the nonaqueous phase to cause the coalescence of particles from the aqueous phase as taught in U.S. Pat. No. 4,589,330. Porous microparticles can be prepared using a first phase containing active agent and a polymer dispersed in a continuous solvent and removing said solvent from the suspension by freeze-drying or dilution-extractionprecipitation as taught in U.S. Pat. No. 4,818,542. Preferred polymers for such preparations are natural or synthetic copolymers or polymers selected from the group consisting of gelatin agar, starch, arabinogalactan, albumin, collagen, polyglycolic acid, polylactic aced, glycolide-L(-) lactide poly(epsilon-caprolactone, poly(epsilon-caprolactone-CO-lactic acid), poly(epsilon-caprolactone-CO-gly colic acid), poly(β-hydroxy butyric acid), polyethylene oxide, polyethylene, poly(alkyl-2-cyanoacrylate), poly(hydroxyethyl methacrylate), polyamides, poly(amino acids), poly(2-hydroxyethyl DL-aspartamide), poly(ester urea), poly(L-phenylalanine/ethylene glycol/1,6-diisocyanatohexane) and poly(methyl methacrylate). Particularly preferred polymers are polyesters, such as polyglycolic acid, polylactic aced, glycolide-L(-) lactide poly(epsilon-caprolactone, poly(epsilon-caprolactone-CO-lactic acid), and poly(epsilon-caprolactone-CO-glycolic acid. Solvents useful for dissolving the polymer and/or the active include: water, hexafluoroisopropanol, methylenechloride, tetrahydrofuran, hexane, benzene, or hexafluoroacetone sesquihydrate. The process of dispersing the active containing phase with a second phase may include pressure forcing said first phase through an orifice in a nozzle to affect droplet formation.

- (616) Dry powder formulations may result from processes other than lyophilization, such as by spray drying or solvent extraction by evaporation or by precipitation of a crystalline composition followed by one or more steps to remove aqueous or nonaqueous solvent. Preparation of a spray-dried protein scaffold preparation is taught in U.S. Pat. No. 6,019,968. The protein scaffold-based dry powder compositions may be produced by spray drying solutions or slurries of the protein scaffold and, optionally, excipients, in a solvent wider conditions to provide a respirable dry powder. Solvents may include polar compounds, such as water and ethanol, which may be readily dried. Protein scaffold stability may be enhanced by performing the spray drying procedures in the absence of oxygen, such as under a nitrogen blanket or by using nitrogen as the drying gas. Another relatively dry formulation is a dispersion of a plurality of perforated microstructures dispersed in a suspension medium that typically comprises a hydrofluoroalkane propellant as taught in WO 9916419. The stabilized dispersions may be administered to the lung of a patient using a metered dose inhaler. Equipment useful in the commercial manufacture of spray dried medicaments are manufactured by Buchi Ltd. or Niro Corp.
- (617) At least one protein scaffold in either the stable or preserved formulations or solutions described herein, can be administered to a patient in accordance with the present invention via a variety of delivery methods including SC or IM injection; transdermal, pulmonary, transmucosal, implant, osmotic pump, cartridge, micro pump, or other means appreciated by the skilled artisan, as well-known in the art. (618) Therapeutic Applications
- (619) The present invention also provides a method for modulating or treating a disease, in a cell, tissue, organ, animal, or patient, as known in the art or as described herein, using at least one protein scaffold of the present invention. e.g., administering or contacting the cell, tissue, organ, animal, or patient with a therapeutic effective amount of protein scaffold. The present invention also provides a method for modulating or treating a disease, in a cell, tissue, organ, animal, or patient.
- (620) Any method of the present invention can comprise administering an effective amount of a composition or pharmaceutical composition comprising at least one protein scaffold to a cell, tissue, organ, animal or patient in need of such modulation, treatment or therapy. Such a method can optionally further comprise co-administration or combination therapy for treating such diseases or disorders, wherein the administering of said at least one protein scaffold, specified portion or variant thereof, further comprises

administering, before concurrently, and/or after, at least one selected from at least one of an alkylating agent, an a mitotic inhibitor, and a radiopharmaceutical. Suitable dosages are well known in the art. See, e.g., Wells et al., eds., Pharmacotherapy Handbook, 2nd Edition, Appleton and Lange, Stamford, Conn. (2000)); PDR Pharmacopoeia, Tarascon Pocket Pharmacopoeia 2000, Deluxe Edition. Tarascon Publishing, Loma Linda, Calif. (2000); Nursing 2001 Handbook of Drugs, 21st edition, Springhouse Corp., Springhouse, Pa., 2001; Health Professional's Drug Guide 2001, ed., Shannon, Wilson, Stang, Prentice-Hall, Inc, Upper Saddle River, N.J., each of which references are entirely incorporated herein by reference.

- (621) Preferred doses can optionally include about 0.1-99 and/or 100-500 mg/kg/administration, or any range, value or fraction thereof, or to achieve a serum concentration of about 0.1-5000  $\mu$ g/ml serum concentration per single or multiple administration, or any range, value or fraction thereof. A preferred dosage range for the protein scaffold of the present invention is from about 1 mg/kg, up to about 3, about 6 or about 12 mg/kg of body weight of the patient.
- (622) Alternatively, the dosage administered can vary depending upon known factors, such as the pharmacodynamic characteristics of the particular agent, and its mode and route of administration, age, health, and weight of the recipient; nature and extent of symptoms, kind of concurrent treatment, frequency of treatment, and the effect desired. Usually a dosage of active ingredient can be about 0.1 to 100 milligrams per kilogram of body weight. Ordinarily 0.1 to 50, and preferably. 0.1 to 10 milligrams per kilogram per administration or in sustained release form is effective to obtain desired results.
- (623) As a non-limiting example, treatment of humans or animals can be provided as a one-time or periodic dosage of at least one protein scaffold of the present invention about 0.1 to 100 mg/kg or any range, value or fraction thereof per day, on at least one of day 1-40, or, alternatively or additionally, at least one of week 1-52, or, alternatively or additionally, at least one of 1-20 years, or any combination thereof, using single, infusion or repeated doses.
- (624) Dosage forms (composition) suitable for internal administration generally contain from about 0.001 milligram to about 500 milligrams of active ingredient per unit or container. In these pharmaceutical compositions the active ingredient will ordinarily be present in an amount of about 0.5-99.999% by weight based on the total weight of the composition.
- (625) For parenteral administration, the protein scaffold can be formulated as a solution, suspension, emulsion, particle, powder, or lyophilized powder in association, or separately provided, with a pharmaceutically acceptable parenteral vehicle. Examples of such vehicles are water, saline, Ringer's solution, dextrose solution, and about 1-10% human serum albumin. Liposomes and nonaqueous vehicles, such as fixed oils, can also be used. The vehicle or lyophilized powder can contain additives that maintain isotonicity (e.g., sodium chloride, mannitol) and chemical stability (e.g., buffers and preservatives). The formulation is sterilized by known or suitable techniques.
- (626) Suitable pharmaceutical carriers are described in the most recent edition of Remington's Pharmaceutical Sciences, A. Osol, a standard reference text in this field.
- (627) Alternative Administration
- (628) Many known and developed modes can be used according to the present invention for administering pharmaceutically effective amounts of at least one protein scaffold according to the present invention. While pulmonary administration is used in the following description, other modes of administration can be used according to the present invention with suitable results. Protein scaffolds of the present invention can be delivered in a carrier, as a solution, emulsion, colloid, or suspension, or as a dry powder, using any of a variety of devices and methods suitable for administration by inhalation or other modes described here within or known in the art.
- (629) Parenteral Formulations and Administration
- (630) Formulations for parenteral administration can contain as common excipients sterile water or saline, polyalkylene glycols, such as polyethylene glycol, oils of vegetable origin, hydrogenated naphthalenes and the like. Aqueous or oily suspensions for injection can be prepared by using an appropriate emulsifier or humidifier and a suspending agent, according to known methods. Agents for injection can be a non-toxic, non-orally administrable diluting agent, such as aqueous solution, a sterile injectable solution or suspension in a solvent. As the usable vehicle or solvent, water, Ringer's solution, isotonic saline, etc. are allowed; as an ordinary solvent or suspending solvent, sterile involatile oil can be used. For these

purposes, any kind of involatile oil and fatty acid can be used, including natural or synthetic or semisynthetic fatty oils or fatty acids; natural or synthetic or semisynthetic mono- or di- or tri-glycerides. Parental administration is known in the art and includes, but is not limited to, conventional means of injections, a gas pressured needle-less injection device as described in U.S. Pat. No. 5,851,198, and a laser perforator device as described in U.S. Pat. No. 5,839,446 entirely incorporated herein by reference. (631) Alternative Delivery

(632) The invention further relates to the administration of at least one protein scaffold by parenteral, subcutaneous, intramuscular, intravenous, intrarticular, intrabronchial, intraabdominal, intracapsular, intracartilaginous, intracavitary, intracelial, intracerebellar, intracerebroventricular, intracolic, intracervical, intragastric, intrahepatic, intramvocardial, intraosteal, intrapelvic, intrapericardiac, intraperitoneal, intrapleural, intraprostatic, intrapulmonary, intrarectal, intrarenal, intraretinal, intraspinal, intrasynovial, intrathoracic, intrauterine, intravesical, intralesional, bolus, vaginal, rectal, buccal, sublingual, intranasal, or transdermal means. At least one protein scaffold composition can be prepared for use for parenteral (subcutaneous, intramuscular or intravenous) or any other administration particularly in the form of liquid solutions or suspensions; for use in vaginal or rectal administration particularly in semisolid forms, such as, but not limited to, creams and suppositories; for buccal, or sublingual administration, such as, but not limited to, in the form of tablets or capsules; or intranasally, such as, but not limited to, the form of powders, nasal drops or aerosols or certain agents; or transdermally, such as not limited to a gel, ointment, lotion, suspension or patch delivery system with chemical enhancers such as dimethyl sulfoxide to either modify the skin structure or to increase the drug concentration in the transdermal patch (Junginger, et al. In "Drug Permeation Enhancement;" Hsieh, D. S., Eds., pp. 59-90 (Marcel Dekker, Inc. New York 1994, entirely incorporated herein by reference), or with oxidizing agents that enable the application of formulations containing proteins and peptides onto the skin (WO 98/53847), or applications of electric fields to create transient transport pathways, such as electroporation, or to increase the mobility of charged drugs through the skin, such as iontophoresis, or application of ultrasound, such as sonophoresis (U.S. Pat. Nos. 4,309,989 and 4,767,402) (the above publications and patents being entirely incorporated herein by reference).

(633) Infusion of Modified Cells as Adoptive Cell Therapy

(634) The disclosure provides modified cells that express one or more CARs and/or CARTyrins of the disclosure that have been selected and/or expanded for administration to a subject in need thereof. Modified cells of the disclosure may be formulated for storage at any temperature including room temperature and body temperature. Modified cells of the disclosure may be formulated for cryopreservation and subsequent thawing. Modified cells of the disclosure may be formulated in a pharmaceutically acceptable carrier for direct administration to a subject from sterile packaging. Modified cells of the disclosure may be formulated in a pharmaceutically acceptable carrier with an indicator of cell viability and/or CAR/CARTyrin expression level to ensure a minimal level of cell function and CAR/CARTyrin expression. Modified cells of the disclosure may be formulated in a pharmaceutically acceptable carrier at a prescribed density with one or more reagents to inhibit further expansion and/or prevent cell death.

(635) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises between 2×10.sup.5 and 5×10.sup.8 cells per kg of body weight of the patient per administration, or any range, value or fraction thereof.

(636) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises between 0.2×10.sup.6 to 20×10.sup.6 cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 0.2×10.sup.6 cells per kg of body weight of the patient per administration, 2×10.sup.6 cells per kg of body weight of the patient per administration, or any cells per kg of body weight of the patient per administration in between.

(637) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 1×10.sup.6 cells or about 1×10.sup.6 cells per kg of body weight of the patient per administration. (638) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 3×10.sup.6 cells or about 3×10.sup.6 cells per kg of body weight of the patient per administration. (639) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises between 0.7×10.sup.6 to 6.7×10.sup.6 cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 0.7×10.sup.6 cells per kg of body weight of the patient per administration, 6.7×10.sup.6 cells per kg of body weight of the patient per administration or any cells per kg of body weight of the patient per administration in between. (640) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises between 0.7×10.sup.6 to 16×10.sup.6 cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 0.7×10.sup.6 cells per kg of body weight of the patient per administration, 2×10.sup.6 cells per kg of body weight of the patient per admiration, 6×10.sup.6 cells per kg of body weight of the patient per administration, 10.7×10.sup.6 cells per kg of body weight of the patient per administration, 16×10.sup.6 cells per kg of body weight of the patient per administration or any cells per kg of body weight of the patient per administration in between. (641) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 1.2×10.sup.6 to 7.1×10.sup.6 cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 1.2×10.sup.6 cells per kg of body weight of the patient per administration, 7.1×10.sup.6 cells per kg of body weight of the patient per administration or any number of cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises between 2×10.sup.6 to 3×10.sup.6 cells per kg of body weight of the patient per administration.

(642) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 1106×10.sup.6 to 2106×10.sup.6 cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 1106×10.sup.6 cells per kg of body weight of the patient per administration, 2106×10.sup.6 cells per kg of body weight of the patient per administration or any number of cells per kg of body weight of the patient per administration in between. In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 0.7×10.sup.6 to 1.3×10.sup.6 cells per kg of body weight of the patient per administration. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises 0.7×10.sup.6 cells per kg of body weight of the patient per administration, 1.3×10.sup.6 cells per kg of body weight of the patient per administration or any number of cells per kg of body weight of the patient per administration in between.

(643) In certain embodiments of the disclosure, modified cells of the disclosure are delivered to a patient via injection or intravenous infusion. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises a single or multiple doses. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises a split dose. In certain embodiments, a therapeutically effective dose of a composition of the disclosure or of compositions comprising modified cells of the disclosure comprises an initial dose and a maintenance dose.

(644) In certain embodiments of the disclosure, the modified cells are T cells and the T cells may be sorted according to T cell markers prior to either in vitro expansion or formulation with a pharmaceutically acceptable carrier. In some embodiments, modified T cells may be sorted on using CD8+ and/or CD4+ markers.

- (645) Inducible Proapoptotic Polypeptides
- (646) Inducible proapoptotic polypeptides of the disclosure are superior to existing inducible polypeptides because the inducible proapoptotic polypeptides of the disclosure are far less immunogenic. While inducible proapoptotic polypeptides of the disclosure are recombinant polypeptides, and, therefore, non-naturally occurring, the sequences that are recombined to produce the inducible proapoptotic polypeptides of the disclosure do not comprise non-human sequences that the host human immune system could recognize as "non-self" and, consequently, induce an immune response in the subject receiving an inducible proapoptotic polypeptide of the disclosure, a cell comprising the inducible proapoptotic polypeptide or a composition comprising the inducible proapoptotic polypeptide.
- (647) The disclosure provides inducible proapoptotic polypeptides comprising a ligand binding region, a linker, and a proapoptotic peptide, wherein the inducible proapoptotic polypeptide does not comprise a non-human sequence. In certain embodiments, the non-human sequence comprises a restriction site. In certain embodiments, the proapoptotic peptide is a caspase polypeptide. In certain embodiments, the caspase 9 polypeptide is a truncated caspase 9 polypeptide. Inducible proapoptotic polypeptides of the disclosure may be non-naturally occurring.
- (648) Caspase polypeptides of the disclosure include, but are not limited to, caspase 1, caspase 2, caspase 3, caspase 4, caspase 5, caspase 6, caspase 7, caspase 8, caspase 9, caspase 10, caspase 11, caspase 12, and caspase 14. Caspase polypeptides of the disclosure include, but are not limited to, those caspase polypeptides associated with apoptosis including caspase 2, caspase 3, caspase 6, caspase 7, caspase 8, caspase 9, and caspase 10. Caspase polypeptides of the disclosure include, but are not limited to, those caspase polypeptides of the disclosure include, but are not limited to, those caspase polypeptides of the disclosure include, but are not limited to, those caspase polypeptides of the disclosure include, but are not limited to, those caspase polypeptides that execute apoptosis, including caspase 3, caspase 6, and caspase 7.
- (649) Caspase polypeptides of the disclosure may be encoded by an amino acid or a nucleic acid sequence having one or more modifications compared to a wild type amino acid or a nucleic acid sequence. The nucleic acid sequence encoding a caspase polypeptide of the disclosure may be codon optimized. The one or more modifications to an amino acid and/or nucleic acid sequence of a caspase polypeptide of the disclosure may increase an interaction, a cross-linking, a cross-activation, or an activation of the caspase polypeptide of the disclosure compared to a wild type amino acid or a nucleic acid sequence. Alternatively. or in addition, the one or more modifications to an amino acid and/or nucleic acid sequence of a caspase polypeptide of the disclosure may decrease the immunogenicity of the caspase poly peptide of the disclosure compared to a wild type amino acid or a nucleic acid sequence.
- (650) Caspase polypeptides of the disclosure may be truncated compared to a wild type caspase polypeptide. For example, a caspase polypeptide may be truncated to eliminate a sequence encoding a Caspase Activation and Recruitment Domain (CARD) to eliminate or minimize the possibility of activating a local inflammatory response in addition to initiating apoptosis in the cell comprising an inducible caspase polypeptide of the disclosure. The nucleic acid sequence encoding a caspase polypeptide of the disclosure may be spliced to form a variant amino acid sequence of the caspase polypeptide of the disclosure compared to a wild type caspase polypeptide. Caspase polypeptides of the disclosure may be encoded by recombinant and/or chimeric sequences. Recombinant and/or chimeric caspase polypeptides of

the disclosure may include sequences from one or more different caspase polypeptides. Alternatively, or in addition, recombinant and/or chimeric caspase poly peptides of the disclosure may include sequences from one or more species (e.g. a human sequence and a non-human sequence). Caspase polypeptides of the disclosure may be non-naturally occurring.

(651) The ligand binding region of an inducible proapoptotic polypeptide of the disclosure may include any polypeptide sequence that facilitates or promotes the dimerization of a first inducible proapoptotic polypeptide of the disclosure with a second inducible proapoptotic polypeptide of the disclosure, the dimerization of which activates or induces cross-linking of the proapoptotic polypeptides and initiation of apoptosis in the cell.

(652) The ligand-binding ("dimerization") region may comprise any polypeptide or functional domain thereof that will allow for induction using an endogenous or non-naturally occurring ligand (i.e., and induction agent), for example, a non-naturally occurring synthetic ligand. The ligand-binding region may be internal or external to the cellular membrane, depending upon the nature of the inducible proapoptotic polypeptide and the choice of ligand (i.e. induction agent). A wide variety of ligand-binding polypeptides and functional domains thereof, including receptors, are known. Ligand-binding regions of the disclosure may include one or more sequences from a receptor. Of particular interest are ligand-binding regions for which ligands (for example, small organic ligands) are known or may be readily produced. These ligandbinding regions or receptors may include, but are not limited to, the FKBPs and cyclophilin receptors, the steroid receptors, the tetracycline receptor, and the like, as well as "non-naturally occurring" receptors, which can be obtained from antibodies, particularly the heavy or light chain subunit, mutated sequences thereof, random amino acid sequences obtained by stochastic procedures, combinatorial syntheses, and the like. In certain embodiments, the ligand-binding region is selected from the group consisting of a FKBP ligand-binding region, a cyclophilin receptor ligand-binding region, a steroid receptor ligand-binding region, a cyclophilin receptors ligand-binding region, and a tetracycline receptor ligand-binding region. (653) The ligand-binding regions comprising one or more receptor domain(s) may be at least about 50 amino acids, and fewer than about 350 amino acids, usually fewer than 200 amino acids, either as the endogenous domain or truncated active portion thereof. The binding region may, for example, be small (<25 kDa, to allow efficient transfection in viral vectors), monomeric, nonimmunogenic, have synthetically accessible, cell permeable, nontoxic ligands that can be configured for dimerization. (654) The ligand-binding regions comprising one or more receptor domain(s) may be intracellular or extracellular depending upon the design of the inducible proapoptotic polypeptide and the availability of an appropriate ligand (i.e. induction agent). For hydrophobic ligands, the binding region can be on either side of the membrane, but for hydrophilic ligands, particularly protein ligands, the binding region will usually be external to the cell membrane, unless there is a transport system for internalizing the ligand in a form in which it is available for binding. For an intracellular receptor, the inducible proapoptotic polypeptide or a transposon or vector comprising the inducible proapoptotic polypeptide may encode a signal peptide and transmembrane domain 5' or 3' of the receptor domain sequence or may have a lipid attachment signal sequence 5' of the receptor domain sequence. Where the receptor domain is between the signal peptide and the transmembrane domain, the receptor domain will be extracellular. (655) Antibodies and antibody subunits, e.g., heavy or light chain, particularly fragments, more particularly all or part of the variable region, or fusions of heavy and light chain to create high-affinity binding, can be used as a ligand binding region of the disclosure. Antibodies that are contemplated include ones that are an ectopically expressed human product, such as an extracellular domain that would not trigger an immune response and generally not expressed in the periphery (i.e., outside the CNS/brain area). Such examples, include, but are not limited to low affinity nerve growth factor receptor (LNGFR), and embryonic surface proteins (i.e., carcinoembryonic antigen). Yet further, antibodies can be prepared against haptenic molecules, which are physiologically acceptable, and the individual antibody subunits screened for binding affinity. The cDNA encoding the subunits can be isolated and modified by deletion of the constant region, portions of the variable region, mutagenesis of the variable region, or the like, to obtain a binding protein domain that has the appropriate affinity for the ligand. In this way, almost any physiologically acceptable haptenic compound can be employed as the ligand or to provide an epitope for the ligand. Instead of antibody units, endogenous receptors can be employed, where the binding region or domain is known and there is a useful or known ligand for binding.

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(656) For multimerizing the receptor, the ligand for the ligand-binding region/receptor domains of the
inducible proapoptotic polypeptides may be multimeric in the sense that the ligand can have at least two
binding sites, with each of the binding sites capable of binding to a ligand receptor region (i.e. a ligand
having a first binding site capable of binding the ligand-binding region of a first inducible proapoptotic
polypeptide and a second binding site capable of binding the ligand-binding region of a second inducible
proapoptotic polypeptide, wherein the ligand-binding regions of the first and the second inducible
proapoptotic polypeptides are either identical or distinct). Thus, as used herein, the term "multimeric
ligand binding region" refers to a ligand-binding region of an inducible proapoptotic polypeptide of the
disclosure that binds to a multimeric ligand. Multimeric ligands of the disclosure include dimeric ligands.
A dimeric ligand of the disclosure may have two binding sites capable of binding to the ligand receptor
domain. In certain embodiments, multimeric ligands of the disclosure are a dimer or higher order
oligomer, usually not greater than about tetrameric, of small synthetic organic molecules, the individual
molecules typically being at least about 150 Da and less than about 5 kDa, usually less than about 3 kDa.
A variety of pairs of synthetic ligands and receptors can be employed. For example, in embodiments
involving endogenous receptors, dimeric FK506 can be used with an FKBP12 receptor, dimerized
cyclosporin A can be used with the cyclophilin receptor, dimerized estrogen with an estrogen receptor,
dimerized glucocorticoids with a glucocorticoid receptor, dimerized tetracycline with the tetracycline
receptor, dimerized vitamin D with the vitamin D receptor, and the like. Alternatively higher orders of the
ligands, e.g., trimeric can be used. For embodiments involving non-naturally occurring receptors, e.g.,
antibody subunits, modified antibody subunits, single chain antibodies comprised of heavy and light chain
variable regions in tandem, separated by a flexible linker, or modified receptors, and mutated sequences
thereof, and the like, any of a large variety of compounds can be used. A significant characteristic of the
units comprising a multimeric ligand of the disclosure is that each binding site is able to bind the receptor
with high affinity, and preferably, that they are able to be dimerized chemically. Also, methods are
available to balance the hydrophobicity/hydrophilicity of the ligands so that they are able to dissolve in
serum at functional levels, yet diffuse across plasma membranes for most applications.
(657) Activation of inducible proapoptotic polypeptides of the disclosure may be accomplished through,
for example, chemically induced dimerization (CID) mediated by an induction agent to produce a
conditionally controlled protein or polypeptide. Proapoptotic polypeptides of the disclosure not only
inducible, but the induction of these polypeptides is also reversible, due to the degradation of the labile
dimerizing agent or administration of a monomeric competitive inhibitor.
(658) In certain embodiments, the ligand binding region comprises a FK506 binding protein 12 (FKBP12)
polypeptide. In certain embodiments, the ligand binding region comprises a FKBP12 poly peptide having
a substitution of valine (V) for phenylalanine (F) at position 36 (F36V) In certain embodiments, in which
the ligand binding region comprises a FKBP12 polypeptide having a substitution of valine (V) for
phenylalanine (F) at position 36 (F36V), the induction agent may comprise AP1903, a synthetic drug
(CAS Index Name: 2-Piperidinecarboxylic acid, 1-[(2S)-1-oxo-2-(3,4,5-trimethoxyphenyl)butyl]-, 1,2-
ethanediylbis[imino(2-oxo-2,1-ethanediyl)oxy-3,1-phenylene[(1 R)-3-(3,4-
dimethoxyphenyl)propylidene]]ester, [2S-[1(R*),2R*[S*[3*[1(R*),2R*]]]]]-(9Cl) CAS Registry Number
195514-63-7; Molecular Formula: C78H98N4O20; Molecular Weight: 1411.65)). In certain embodiments,
in which the ligand binding region comprises a FKBP12 polypeptide having a substitution of valine (V)
for phenylalanine (F) at position 36 (F36V), the induction agent may comprise AP20187 (CAS Registry
Number: 195514-80-8 and Molecular Formula: C82H107N5O20). In certain embodiments, the induction
agent is an AP20187 analog, such as, for example, AP1510. As used herein, the induction agents AP20187,
AP1903 and AP1510 may be used interchangeably.
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(659) AP1903 API is manufactured by Alphora Research Inc, and AP1903 Drug Product for Injection is made by Formatech Inc. It is formulated as a 5 mg/mL solution of AP1903 in a 25% solution of the nonionic solubilizer Solutol HS 15 (250 mg/mL, BASF). At room temperature, this formulation is a clear, slightly yellow solution. Upon refrigeration, this formulation undergoes a reversible phase transition, resulting in a milky solution. This phase transition is reversed upon re-warming to room temperature. The fill is 2.33 mL in a 3 mL glass vial (approximately 10 mg AP1903 for Injection total per vial). Upon determining a need to administer AP1903, patients may be, for example, administered a single fixed dose of AP1903 for Injection (0.4 mg/kg) via IV infusion over 2 hours, using a non-DEHP, non-ethylene oxide

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sterilized infusion set. The dose of AP1903 is calculated individually for all patients, and is not be
recalculated unless body weight fluctuates by ≥10%. The calculated dose is diluted in 100 mL in 0.9%
normal saline before infusion. In a previous Phase I study of AP1903, 24 healthy volunteers were treated
with single doses of AP1903 for Injection at dose levels of 0.01, 0.05, 0.1, 0.5 and 1.0 mg/kg infused IV
over 2 hours. AP1903 plasma levels were directly proportional to dose, with mean Cmax values ranging
from approximately 10-1275 ng/mL over the 0.01-1.0 mg/kg dose range. Following the initial infusion
period, blood concentrations demonstrated a rapid distribution phase, with plasma levels reduced to
approximately 18, 7, and 1% of maximal concentration at 0.5, 2 and 10 hours post-dose, respectively.
AP1903 for Injection was shown to be safe and well tolerated at all dose levels and demonstrated a
favorable pharmacokinetic profile. Iuliucci J D, et al., J Clin Pharmacol. 41: 870-9, 2001.
(660) The fixed dose of AP1903 for injection used, for example, may be 0.4 mg/kg intravenously infused
over 2 hours. The amount of AP1903 needed in vitro for effective signaling of cells is 10-100 nM (1600
Da MW). This equates to 16-160 \mug/L or {tilde over ( )}0.016-1.6 \mug/kg (1.6-160 \mug/kg). Doses up to 1
mg/kg were well-tolerated in the Phase I study of AP1903 described above. Therefore, 0.4 mg/kg may be a
safe and effective dose of AP1903 for this Phase I study in combination with the therapeutic cells.
(661) The amino acid and/or nucleic acid sequence encoding ligand binding of the disclosure may contain
sequence one or more modifications compared to a wild type amino acid or nucleic acid sequence. For
example, the amino acid and/or nucleic acid sequence encoding ligand binding region of the disclosure
may be a codon-optimized sequence. The one or more modifications may increase the binding affinity of a
ligand (e.g. an induction agent) for the ligand binding region of the disclosure compared to a wild type
polypeptide. Alternatively, or in addition, the one or more modifications may decrease the immunogenicity
of the ligand binding region of the disclosure compared to a wild type polypeptide. Ligand binding regions
of the disclosure and/or induction agents of the disclosure may be non-naturally occurring.
(662) Inducible proapoptotic polypeptides of the disclosure comprise a ligand binding region, a linker and
a proapoptotic peptide, wherein the inducible proapoptotic polypeptide does not comprise a non-human
sequence. In certain embodiments, the non-human sequence comprises a restriction site. The linker may
comprise any organic or inorganic material that permits, upon dimerization of the ligand binding region,
interaction, cross-linking, cross-activation, or activation of the proapoptotic polypeptides such that the
interaction or activation of the proapoptotic polypeptides initiates apoptosis in the cell. In certain
embodiments, the linker is a polypeptide. In certain embodiments, the linker is a polypeptide comprising a
G/S rich amino acid sequence (a "GS" linker). In certain embodiments, the linker is a polypeptide
comprising the amino acid sequence GGGGS (SEQ ID NO: 17014). In preferred embodiments, the linker
is a polypeptide and the nucleic acid encoding the poly peptide does not contain a restriction site for a
restriction endonuclease. Linkers of the disclosure may be non-naturally occurring.
(663) Inducible proapoptotic polypeptides of the disclosure may be expressed in a cell under the
transcriptional regulation of any promoter capable of initiating and/or regulating the expression of an
inducible proapoptotic polypeptide of the disclosure in that cell. The term "promoter" as used herein refers
to a promoter that acts as the initial binding site for RNA polymerase to transcribe a gene. For example,
inducible proapoptotic polypeptides of the disclosure may be expressed in a mammalian cell under the
transcriptional regulation of any promoter capable of initiating and/or regulating the expression of an
inducible proapoptotic polypeptide of the disclosure in a mammalian cell, including, but not limited to
native, endogenous, exogenous, and heterologous promoters. Preferred mammalian cells include human
cells. Thus, inducible proapoptotic polypeptides of the disclosure may be expressed in a human cell under
the transcriptional regulation of any promoter capable of initiating and/or regulating the expression of an
inducible proapoptotic polypeptide of the disclosure in a human cell, including, but not limited to, a
human promoter or a viral promoter. Exemplary promoters for expression in human cells include, but are
not limited to, a human cytomegalovirus (CMV) immediate early gene promoter, a SV40 early promoter, a
Rous sarcoma virus long terminal repeat, β-actin promoter, a rat insulin promoter and a glyceraldehyde-3-
phosphate dehydrogenase promoter, each of which may be used to obtain high-level expression of an
inducible proapoptotic polypeptide of the disclosure. The use of other viral or mammalian cellular or
bacterial phage promoters which are well known in the art to achieve expression of an inducible
proapoptotic polypeptide of the disclosure is contemplated as well, provided that the levels of expression
are sufficient for initiating apoptosis in a cell. By employing a promoter with well-known properties, the
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level and pattern of expression of the protein of interest following transfection or transformation can be optimized.

(664) Selection of a promoter that is regulated in response to specific physiologic or synthetic signals can permit inducible expression of the inducible proapoptotic polypeptide of the disclosure. The ecdysone system (Invitrogen, Carlsbad, Calif.) is one such system. This system is designed to allow regulated expression of a gene of interest in mammalian cells. It consists of a tightly regulated expression mechanism that allows virtually no basal level expression of a transgene, but over 200-fold inducibility. The system is based on the heterodimeric ecdysone receptor of *Drosophila*, and when ecdysone or an analog such as muristerone A binds to the receptor, the receptor activates a promoter to turn on expression of the downstream transgene high levels of mRNA transcripts are attained. In this system, both monomers of the heterodimeric receptor are constitutively expressed from one vector, whereas the ecdysoneresponsive promoter, which drives expression of the gene of interest, is on another plasmid. Engineering of this type of system into a vector of interest may therefore be useful. Another inducible system that may be useful is the Tet-Off<sup>TM</sup> or Tet-On<sup>TM</sup> system (Clontech. Palo Alto, Calif.) originally developed by Gossen and Bujard (Gossen and Bujard, Proc. Natl. Acad. Sci. USA, 89:5547-5551, 1992; Gossen et al., Science, 268:1766-1769, 1995). This system also allows high levels of gene expression to be regulated in response to tetracycline or tetracycline derivatives such as doxycycline. In the Tet-On™ system, gene expression is turned on in the presence of doxycycline, whereas in the Tet-Off<sup>TM</sup> system, gene expression is turned on in the absence of doxycycline. These systems are based on two regulatory elements derived from the tetracycline resistance operon of E. coli: the tetracycline operator sequence (to which the tetracycline repressor binds) and the tetracycline repressor protein. The gene of interest is cloned into a plasmid behind a promoter that has tetracycline-responsive elements present in it. A second plasmid contains a regulatory element called the tetracycline-controlled transactivator, which is composed, in the Tet-Off<sup>TM</sup> system, of the VP16 domain from the herpes simplex virus and the wild-type tetracycline repressor. Thus in the absence of doxycycline, transcription is constitutively on. In the Tet-On™ system, the tetracycline repressor is not wild type and in the presence of doxycycline activates transcription. For gene therapy vector production, the Tet-Off<sup>TM</sup> system may be used so that the producer cells could be grown in the presence of tetracycline or doxycycline and prevent expression of a potentially toxic transgene, but when the vector is introduced to the patient, the gene expression would be constitutively on. (665) In some circumstances, it is desirable to regulate expression of a transgene in a gene therapy vector. For example, different viral promoters with varying strengths of activity are utilized depending on the level of expression desired. In mammalian cells, the CMV immediate early promoter is often used to provide strong transcriptional activation. The CMV promoter is reviewed in Donnelly, J. J., et al., 1997. Annu. Rev. Immunol. 15:617-48. Modified versions of the CMV promoter that are less potent have also been used when reduced levels of expression of the transgene are desired. When expression of a transgene in hematopoietic cells is desired, retroviral promoters such as the LTRs from MLV or MMTV are often used. Other viral promoters that are used depending on the desired effect include SV40, RSV LTR, HIV-1 and HIV-2 LTR, adenovirus promoters such as from the E1A, E2A, or MLP region, AAV LTR, HSV-TK,

(666) In other examples, promoters may be selected that are developmentally regulated and are active in particular differentiated cells. Thus, for example, a promoter may not be active in a pluripotent stem cell, but, for example, where the pluripotent stem cell differentiates into a more mature cell, the promoter may then be activated.

and avian sarcoma virus.

(667) Similarly tissue specific promoters are used to effect transcription in specific tissues or cells so as to reduce potential toxicity or undesirable effects to non-targeted tissues. These promoters may result in reduced expression compared to a stronger promoter such as the CMV promoter, but may also result in more limited expression, and immunogenicity (Bojak, A., et al., 2002. Vaccine. 20:1975-79; Cazeaux, N., et al., 2002. Vaccine 20:3322-31). For example, tissue specific promoters such as the PSA associated promoter or prostate-specific glandular kallikrein, or the muscle creatine kinase gene may be used where appropriate.

(668) Examples of tissue specific or differentiation specific promoters include, but are not limited to, the following: B29 (B cells); CD14 (monocytic cells); CD43 (leukocytes and platelets); CD45 (hematopoietic cells); CD68 (macrophages); desmin (muscle); elastase-1 (pancreatic acinar cells); endoglin (endothelial

cells); fibronectin (differentiating cells, healing tissues); and Flt-1 (endothelial cells); GFAP (astrocytes). (669) In certain indications, it is desirable to activate transcription at specific times after administration of the gene therapy vector. This is done with such promoters as those that are hormone or cytokine regulatable Cytokine and inflammatory protein responsive promoters that can be used include K and T kininogen (Kageyama et al., (1987) J. Biol. Chem., 262, 2345-2351), c-fos, TNF-alpha, C-reactive protein (Arcone, et al., (1988) Nucl. Acids Res., 16(8), 3195-3207), haptoglobin (Oliviero et al., (1987) EMBO J., 6, 1905-1912), serum amyloid A2, C/EBP alpha, IL-1, IL-6 (Poli and Cortese. (1989) Proc. Nat'l Acad. Sci. USA, 86, 8202-8206), Complement C3 (Wilson et al., (1990) Mol. Cell. Biol., 6181-6191), IL-8, alpha-1 acid glycoprotein (Prowse and Baumann. (1988) Mol Cell Biol, 8, 42-51), alpha-1 antitrypsin, lipoprotein lipase (Zechner et al., Mol. Cell. Biol., 2394-2401, 1988), angiotensinogen (Ron, et al., (1991) Mol. Cell. Biol., 2887-2895), fibrinogen, c-jun (inducible by phorbol esters, TNF-alpha, UV radiation, retinoic acid, and hydrogen peroxide), collagenase (induced by phorbol esters and retinoic acid), metallothionein (heavy metal and glucocorticoid inducible), Stromelysin (inducible by phorbol ester, interleukin-1 and EGF), alpha-2 macroglobulin and alpha-1 anti-chymotrypsin. Other promoters include, for example, SV40, MMTV, Human Immunodeficiency Virus (MV), Moloney virus, ALV, Epstein Barr virus. Rous Sarcoma virus, human actin, myosin, hemoglobin, and creatine.

(670) It is envisioned that any of the above promoters alone or in combination with another can be useful depending on the action desired. Promoters, and other regulatory elements, are selected such that they are functional in the desired cells or tissue. In addition, this list of promoters should not be construed to be exhaustive or limiting; other promoters that are used in conjunction with the promoters and methods disclosed herein.

(671) Armored T-Cells "Knock Down" Strategy

(672) T-cells of the disclosure may be genetically modified to enhance their therapeutic potential. Alternatively, or in addition, T-cells of the disclosure may be modified to render them less sensitive to immunologic and/or metabolic checkpoints. Modifications of this type "armor" the T cells of the disclosure, which, following the modification, may be referred to here as "armored" T cells. Armored T cells of the disclosure may be produced by, for example, blocking and/or diluting specific endogenous checkpoint signals delivered to the T-cells (i.e. checkpoint inhibition) within the tumor immunosuppressive microenvironment, for example.

(673) In some embodiments, an armored T-cell of the disclosure is derived from a T cell, a NK cell, a hematopoietic progenitor cell, a peripheral blood (PB) derived T cell (including a T cell isolated or derived from G-CSF-mobilized peripheral blood), or an umbilical cord blood (UCB) derived T cell. In some embodiments, an armored T-cell of the disclosure comprises one or more of a chimeric ligand receptor (CLR comprising a protein scaffold, an antibody, an ScFv, or an antibody mimetic)/chimeric antigen receptor (CAR comprising a protein scaffold, an antibody, an ScFv. or an antibody mimetic), a CARTyrin (a CAR comprising a Centyrin), and/or a VCAR (a CAR comprising a camelid VHH or a single domain VH) of the disclosure. In some embodiments, an armored T-cell of the disclosure comprises an inducible proapoptotic polypeptide comprising (a) a ligand binding region, (b) a linker, and (c) a truncated caspase 9 poly peptide, wherein the inducible proapoptotic polypeptide does not comprise a non-human sequence. In some embodiments, the non-human sequence is a restriction site. In some embodiments, the ligand binding region inducible caspase polypeptide comprises a FK506 binding protein 12 (FKBP12) polypeptide. In some embodiments, the amino acid sequence of the FK506 binding protein 12 (FKBP12) polypeptide comprises a modification at position 36 of the sequence. In some embodiments, the modification is a substitution of valine (V) for phenylalanine (F) at position 36 (F36V). In some embodiments, an armored T-cell of the disclosure comprises an exogenous sequence. In some embodiments, the exogenous sequence comprises a sequence encoding a therapeutic protein. Exemplary therapeutic proteins may be nuclear, cytoplasmic, intracellular, transmembrane, cell-surface bound, or secreted proteins. Exemplary therapeutic proteins expressed by the armored T cell may modify an activity of the armored T cell or may modify an activity of a second cell. In some embodiments, an armored T-cell of the disclosure comprises a selection gene or a selection marker. In some embodiments, an armored Tcell of the disclosure comprises a synthetic gene expression cassette (also referred to herein as an inducible transgene construct).

(674) In some embodiments, a T-cell of the disclosure is modified to silence or reduce expression one or

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more gene(s) encoding receptor(s) of inhibitory checkpoint signals to produce an armored T-cell of the
disclosure. Examples of inhibitory checkpoint signals include, but are not limited to, a PD-L1 ligand
binding to a PD-1 receptor on a CAR-T cell of the disclosure or a TGFβ cytokine binding to a TGFβRII
receptor on a CAR-T cell. Receptors of inhibitory checkpoint signals are expressed on the cell surface or
within the cytoplasm of a T-cell. Silencing or reducing expressing of the gene encoding the receptor of the
inhibitory checkpoint signal results a loss of protein expression of the inhibitory checkpoint receptors on
the surface or within the cytoplasm of an armored T-cell of the disclosure. Thus, armored T cells of the
disclosure having silenced or reduced expression of one or more genes encoding an inhibitory checkpoint
receptor is resistant, non-receptive or insensitive to checkpoint signals. The armored T cell's resistance or
decreased sensitivity to inhibitory checkpoint signals enhances the armored T cell's therapeutic potential in
the presence of these inhibitory checkpoint signals. Inhibitory checkpoint signals include but are not
limited to the examples listed in Table 2. Exemplary inhibitory checkpoint signals that may be silenced in
an armored T cell of the disclosure include, but are not limited to, PD-1 and TGFβRII.
(675) TABLE-US-00134 TABLE 2 Exemplary Inhibitory Checkpoint Signals (and proteins that induce
immunosuppression). Full Name Abbreviation SEQ ID NO: Programmed cell death protein 1 PD1 14643-
14644 transforming growth factor β Receptor 1 TGFβR1 14645 transforming growth factor β Receptor 2
TGFβR2 14646 T-cell immunoglobulin and mucin-domain TIM3 14647 containing-3 Lymphocyte-
activation gene 3 LAG3 14648 Cytotoxic T-lymphocyte protein 4 CTLA4 14649 B- and T-lymphocyte
attenuator BTLA 14650 Killer cell immunoglobulin-like receptor KIR 14651 Alpha-2A adrenergic
receptor A2aR 14652 V-type immunoglobulin domain-containing VISTA 14653 suppressor of T-cell
activation T-cell immunoreceptor with Ig and ITIM TIGIT 14654 domains Programmed cell death 1 ligand
1 B7H1 or PD-L1 14655 Programmed cell death 1 ligand 2 B7DC or PD-L2 14656 T-lymphocyte
activation antigen CD80 B7-1 or CD80 14657 T-lymphocyte activation antigen CD86 B7-2 or CD86
14658 CD160 antigen CD160 14659 Leukocyte-associated immunoglobulin-like LAIR1 14660 receptor 1
T-cell immunoglobulin and mucin domain- TIM4 or TIMD4 14661 containing protein 4 Natural killer cell
receptor 2B4 2B4 or CD244 14662 Major Histocompatibility Complex type I MHC I 14663 Major
Histocompatibility Complex type II MHC II Putative 2-methylcitrate dehydratase receptor PDH1R T-cell
immunoglobulin and mucin domain 1 TIM1R receptor T-cell immunoglobulin and mucin domain 4
TIM4R receptor B7-H3 receptor B7H3R or CD176 Receptor B7-H4 receptor B7H4R Immunoglobulin-
like transcript (ILT) 3 receptor ILT3R phosphoinositide 3-kinase, subunit alpha PI3K alpha 14664
phosphoinositide 3-kinase, subunit gamma PI3K gamma 14665 Tyrosine-protein phosphatase non-receptor
type SHP2 or PTPN11 14666 11 Protein phosphatase 2, subunit gamma PP2A gamma 14667 Protein
phosphatase 2, subunit beta PP2A beta 14668 Protein phosphatase 2, subunit delta PP2A delta 14669
Protein phosphatase 2, subunit epsilon PP2A epsilon 14670 Protein phosphatase 2, subunit alpha PP2A
alpha 14671 T-cell Receptor, subunit alpha TCR alpha 14672 T-cell Receptor, subunit beta TCR beta
14673 T-cell Receptor, subunit zeta TCR zeta 14674 T-cell Receptor, subunit CD3 epsilon TCR CD3
epsilon 14675 T-cell Receptor, subunit CD3 gamma TCR CD3 gamma 14676 T-cell Receptor, subunit
CD3 delta TCR CD3 delta 14677 Cluster of Differentiation 28 CD28 14678 Galectins Galectins Galectin 9
Galectin 9 14679 High Mobility Group Box 1 HMGB1 14680 Arginase 1 ARG1 14681 Prostaglandin-
Endoperoxide Synthase 1 PTGS1 14682 Prostaglandin-Endoperoxide Synthase 2 PTGS2 14683 Mucin 1,
Cell Surface Associated MUC1 14684 Mucin 2, Oligomeric Mucus/Gel-Forming MUC2 14685 Mucin
3A, Cell Surface Associated MUC3A 14686 Mucin 3B, Cell Surface Associated MUC3B 14687 Mucin 4,
Cell Surface Associated MUC4 14688 Mucin 5AC, Oligomeric Mucus/Gel-Forming MUC5AC 14689
Mucin 5B, Oligomeric Mucus/Gel-Forming MUC5B 14690 Mucin 6, Oligomeric Mucus/Gel-Forming
MUC6 14691 Mucin 7, Secreted MUC7 14692 Mucin 8 MUC8 Mucin 12, Cell Surface Associated
MUC12 14693 Mucin 13, Cell Surface Associated MUC13 14694 Mucin 15, Cell Surface Associated
MUC15 14695 Mucin 16, Cell Surface Associated MUC16 14696 Mucin 17, Cell Surface Associated
MUC17 14697 Mucin 19, Oligomeric MUC19 14698 Mucin 20, Cell Surface Associated MUC20 14699
Mucin 21, Cell Surface Associated MUC21 14700 Mucin 22 MUC22 14701 Indoleamine 2,3-
Dioxygenase 1 IDO1 14702 Indoleamine 2,3-Dioxygenase 2 IDO2 14703 Inducible T Cell Costimulator
Ligand ICOSLG 14704 ROS Proto-Oncogene 1, Receptor Tyrosine ROS1 14705 Kinase Tumor Necrosis
Factor Receptor Superfamily 4-1BB, CD137, ILA or 14706 Member 9 TNFRSF9 4-1BB Ligand 4-1BB-L
14707 Glucocorticoid-induced TNFR family related GITR 14708 gene Glucocorticoid-induced TNFR
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family related GITRL 14709 gene ligand

(676) In some embodiments, a T-cell of the disclosure is modified to silence or reduce expression of one or more gene(s) encoding intracellular proteins involved in checkpoint signaling to produce an armored T-cell of the disclosure. The activity of a T-cell of the disclosure may be enhanced by targeting any intracellular signaling protein involved in a checkpoint signaling pathway thereby achieving checkpoint inhibition or interference to one or more checkpoint pathways. Intracellular signaling proteins involved in checkpoint signaling include, but are not limited to, exemplary intracellular signaling proteins listed in Table 3.

(677) TABLE-US-00135 TABLE 3 Exemplary Intracellular Signaling Proteins. Full Name Abbreviation SEQ ID NO: phosphoinositide 3-kinase, subunit alpha PI3K alpha 14710 phosphoinositide 3-kinase, subunit gamma PI3K gamma 14711 Tyrosine-protein phosphatase non-receptor type SHP2 or PTPN11 14712 11 Protein phosphatase 2, subunit gamma PP2A gamma 14713 Protein phosphatase 2, subunit beta PP2A beta 14714 Protein phosphatase 2, subunit delta PP2A delta 14715 Protein phosphatase 2, subunit epsilon PP2A epsilon 14716 Protein phosphatase 2, subunit alpha PP2A alpha 14717 RAC-alpha serine/threonine-protein kinase AKT or PKB 14718 Tyrosine-protein kinase ZAP-70 ZAP70 14719 Amino acid sequence (KIEELE)-containing KIEELE-domain domain protein containing proteins BCL2 associated athanogene 6 Bat3, Bag6 or Scythe 14720 B-cell lymphoma-extra large Bcl-xL 14721 Bcl-2-related protein A1 Bfl-1 or BCL2A1 14722

(678) In some embodiments, a T-cell of the disclosure is modified to silence or reduce expression of one or more gene(s) encoding a transcription factor that hinders the efficacy of a therapy to produce an armored T-cell of the disclosure. The activity of armored T-cells may be enhanced or modulated by silencing or reducing expression (or repressing a function) of a transcription factor that hinders the efficacy of therapy. Exemplary transcription factors that may be modified to silence or reduce expression or to repress a function thereof include, but are not limited to the exemplary transcription factors listed in Table 4. For example expression of a FOXP3 gene may be silenced or reduced in an armored T cell of the disclosure to prevent or reduce the formation of T regulatory CAR-T-cells (CAR-Treg cells), the expression or activity of which may reduce efficacy of a therapy.

(679) TABLE-US-00136 TABLE 4 Exemplary Transcription Factors. Full Name Abbreviation SEQ ID NO: activity-dependent neuroprotector homeobox ADNP 14723 ADNP homeobox 2 ADNP2 14724 AE binding protein 1 AEBP1 14725 AE binding protein 2 AEBP2 14726 AF4/FMR2 family member 1 AFF1 14727 AF4/FMR2 family member 2 AFF2 14728 AF4/FMR2 family member 3 AFF3 14729 AF4/FMR2 family member 4 AFF4 14730 AT-hook containing transcription factor 1 AHCTF1 14731 aryl hydrocarbon receptor AHR 14732 aryl-hydrocarbon receptor repressor AHRR 14733 autoimmune regulator AIRE 14734 AT-hook transcription factor AKNA 14735 ALX homeobox 1 ALX1 14736 ALX homeobox 3 ALX3 14737 ALX homeobox 4 ALX4 14738 ankyrin repeat and zinc finger domain containing 1 ANKZF1 14739 adaptor related protein complex 5 zeta 1 subunit AP5Z1 14740 androgen receptor AR 14741 arginine-fifty homeobox ARGFX 14742 Rho GTPase activating protein 35 ARHGAP35 14743 ATrich interaction domain 1A ARID1A 14744 AT-rich interaction domain 1B ARID1B 14745 AT-rich interaction domain 2 ARID2 14746 AT-rich interaction domain 3A ARID3A 14747 AT-rich interaction domain 3B ARID3B 14748 AT-rich interaction domain 3C ARID3C 14749 AT-rich interaction domain 4A ARID4A 14750 AT-rich interaction domain 4B ARID4B 14751 AT-rich interaction domain 5A ARID5A 14752 AT-rich interaction domain 5B ARID5B 14753 aryl hydrocarbon receptor nuclear translocator ARNT 14754 aryl hydrocarbon receptor nuclear translocator 2 ARNT2 14755 aryl hydrocarbon receptor nuclear translocator like ARNTL 14756 aryl hydrocarbon receptor nuclear translocator like 2 ARNTL2 14757 aristaless related homeobox ARX 14758 achaete-scute family bHLH transcription factor 1 ASCL1 14759 achaete-scute family bHLH transcription factor 2 ASCL2 14760 achaete-scute family bHLH transcription factor 3 ASCL3 14761 achaete-scute family bHLH transcription factor 4 ASCL4 14762 achaete-scute family bHLH transcription factor 5 ASCL5 14763 ash1 (absent, small, or homeotic)-like (Drosophila) ASH1L 14764 ash2 (absent, small, or homeotic)-like (Drosophila) ASH2L 14765 activating transcription factor 1 ATF1 14766 activating transcription factor 2 ATF2 14767 activating transcription factor 3 ATF3 14768 activating transcription factor 4 ATF4 14769 activating transcription factor 5 ATF5 14770 activating transcription factor 6 ATF6 14771 activating transcription factor 6 beta ATF6B 14772 activating transcription factor 7 ATF7 14773 atonal bHLH transcription factor 1 ATOH1 14774 atonal

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bHLH transcription factor 7 ATOH7 14775 atonal bHLH transcription factor 8 ATOH8 14776 alpha
thalassemia/mental retardation syndrome X- ATRX 14777 linked ataxin 7 ATXN7 14778 BTB and CNC
homology 1, basic leucine zipper BACH1 14779-14780 transcription factor1 BTB domain and CNC
homolog 2 BACH2 14781 BarH like homeobox 1 BARHL1 14782 BarH like homeobox 2 BARHL2
14783 BARX homeobox 1 BARX1 14784 BARX homeobox 2 BARX2 14785 Basic Leucine Zipper ATF-
Like Transcription Factor, Batf 14786 basic leucine zipper transcription factor, ATF-like BATF 14786
basic leucine zipper transcription factor, ATF-like 2 BATF2 14787 basic leucine zipper transcription
factor, ATF-like 3 BATF3 14788 bobby sox homolog (Drosophila) BBX 14789 B-cell CLL/lymphoma
11A BCL11A 14790 B-cell CLL/lymphoma 11B BCL11B 14791 B-cell CLL/lymphoma 3 BCL3 14792
B-cell CLL/lymphoma 6 BCL6 14793 B-cell CLL/lymphoma 6, member B BCL6B 14794 BCL2
associated transcription factor 1 BCLAF1 14795 basic helix-loop-helix family member a15 BHLHA15
14796 basic helix-loop-helix family member a9 BHLHA9 14797 basic helix-loop-helix domain
containing, class B, 9 BHLHB9 14798 basic helix-loop-helix family member e22 BHLHE22 14799 basic
helix-loop-helix family member e23 BHLHE23 14800 basic helix-loop-helix family member e40
BHLHE40 14801 basic helix-loop-helix family member e41 BHLHE41 14802 Beta-Interferon Gene
Positive-Regulatory Domain I Blimp-1 14803 Binding Factor bone morphogenetic protein 2 BMP2 14804
basonuclin 1 BNC1 14805 basonuclin 2 BNC2 14806 bolA family member 1 BOLA1 14807 bolA family
member 2 BOLA2 14808 bolA family member 3 BOLA3 14809 bromodomain PHD finger transcription
factor BPTF 14810 breast cancer 1 BRCA1 14811 brain specific homeobox BSX 14812 chromosome 20
open reading frame 194 C20orf194 14813 calmodulin binding transcription activator 1 CAMTA1 14814
calmodulin binding transcription activator 2 CAMTA2 14815 calcium regulated heat stable protein 1
CARHSP1 14816 castor zinc finger 1 CASZ1 14817 core-binding factor, beta subunit CBFB 14818
coiled-coil domain containing 79 CCDC79 14819 cell division cycle 5 like CDC5L 14820 caudal type
homeobox 1 CDX1 14821 caudal type homeobox 2 CDX2 14822 caudal type homeobox 4 CDX4 14823
CCAAT/enhancer binding protein alpha CEBPA 14824 CCAAT/enhancer binding protein beta CEBPB
14825 CCAAT/enhancer binding protein delta CEBPD 14826 CCAAT/enhancer binding protein epsilon
CEBPE 14827 CCAAT/enhancer binding protein gamma CEBPG 14828 CCAAT/enhancer binding
protein zeta CEBPZ 14829 centromere protein T CENPT 14830 ceramide synthase 3 CERS3 14831
ceramide synthase 6 CERS6 14832 chromosome alignment maintaining phosphoprotein 1 CHAMP1
14833 capicua transcriptional repressor CIC 14834 CDKN1A interacting zinc finger protein 1 CIZ1 14835
clock circadian regulator CLOCK 14836 CCR4-NOT transcription complex subunit 4 CNOT4 14837 CPX
chromosome region, candidate 1 CPXCR1 14838 cramped chromatin regulator homolog 1 CRAMP1
14839 cAMP responsive element binding protein 1 CREB1 14840 cAMP responsive element binding
protein 3 CREB3 14841 cAMP responsive element binding protein 3-like 1 CREB3L1 14842 cAMP
responsive element binding protein 3-like 2 CREB3L2 14843 cAMP responsive element binding protein
3-like 3 CREB3L3 14844 cAMP responsive element binding protein 3-like 4 CREB3L4 14845 cAMP
responsive element binding protein 5 CREB5 14846 CREB binding protein CREBBP 14847 cAMP
responsive element binding protein-like 2 CREBL2 14848 CREB3 regulatory factor CREBRF 14849
CREB/ATF bZIP transcription factor CREBZF 14850 cAMP responsive element modulator CREM 14851
cone-rod homeobox CRX 14852 cysteine-serine-rich nuclear protein 1 CSRNP1 14853 cysteine-serine-
rich nuclear protein 2 CSRNP2 14854 cysteine-serine-rich nuclear protein 3 CSRNP3 14855 CCCTC-
binding factor (zinc finger protein) CTCF 14856 CCCTC-binding factor like CTCFL 14857 cut-like
homeobox 1 CUX1 14858-14859 cut-like homeobox 2 CUX2 14860 CXXC finger protein 1 CXXC1
14861 dachshund family transcription factor 1 DACH1 14862 dachshund family transcription factor 2
DACH2 14863 D site of albumin promoter (albumin D-box) binding DBP 14864 protein developing brain
homeobox 1 DBX1 14865 developing brain homeobox 2 DBX2 14866 damage specific DNA binding
protein 2 DDB2 14867 DNA damage inducible transcript 3 DDIT3 14868 DEAF1, transcription factor
DEAF1 14869 distal-less homeobox 1 DLX1 14870 distal-less homeobox 2 DLX2 14871 distal-less
homeobox 3 DLX3 14872 distal-less homeobox 4 DLX4 14873 distal-less homeobox 5 DLX5 14874
distal-less homeobox 6 DLX6 14875 DNA methyltransferase 1 associated protein 1 DMAP1 14876
diencephalon/mesencephalon homeobox 1 DMBX1 14877 doublesex and mab-3 related transcription
factor 1 DMRT1 14878 doublesex and mab-3 related transcription factor 2 DMRT2 14879 doublesex and
mab-3 related transcription factor 3 DMRT3 14880 DMRT like family A1 DMRTA1 14881 DMRT like
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family A2 DMRTA2 14882 DMRT like family B with proline rich C-terminal 1 DMRTB1 14883 DMRT
like family C1 DMRTC1 14884 DMRT like family C1B DMRTC1B 14884 DMRT like family C2
DMRTC2 14885 cyclin D binding myb like transcription factor 1 DMTF1 14886 DnaJ heat shock protein
family (Hsp40) member C1 DNAJC1 14887 DnaJ heat shock protein family (Hsp40) member C2
DNAJC2 14888 DnaJ heat shock protein family (Hsp40) member C21 DNAJC21 14889 DNA (cytosine-
5-)-methyltransferase 1 DNMT1 14890 DNA (cytosine-5-)-methyltransferase 3 alpha DNMT3A 14891
DNA (cytosine-5-)-methyltransferase 3 beta DNMT3B 14892 DNA (cytosine-5-)-methyltransferase 3-like
DNMT3L 14893 double PHD fingers 1 DPF1 14894 double PHD fingers 2 DPF2 14895 double PHD
fingers 3 DPF3 14896 divergent-paired related homeobox DPRX 14897 down-regulator of transcription 1
DR1 14898 DR1 associated protein 1 DRAP1 14899 dorsal root ganglia homeobox DRGX 14900 double
homeobox 4 DUX4 14901 double homeobox 4 like 9 DUX4L9 14902 double homeobox A DUXA 14903
E2F transcription factor 1 E2F1 14904 E2F transcription factor 2 E2F2 14905 E2F transcription factor 3
E2F3 14906 E2F transcription factor 4 E2F4 14907 E2F transcription factor 5 E2F5 14908 E2F
transcription factor 6 E2F6 14909 E2F transcription factor 7 E2F7 14910 E2F transcription factor 8 E2F8
14911 E4F transcription factor 1 E4F1 14912 early B-cell factor 1 EBF1 14913 early B-cell factor 2 EBF2
14914 early B-cell factor 3 EBF3 14915 early B-cell factor 4 EBF4 14916 early growth response 1 EGR1
14917 early growth response 2 EGR2 14918 early growth response 3 EGR3 14919 early growth response
4 EGR4 14920 ets homologous factor EHF 14921 E74-like factor 1 (ets domain transcription factor) ELF1
14922 E74-like factor 2 (ets domain transcription factor) ELF2 14923 E74-like factor 3 (ets domain
transcription factor, ELF3 14924 epithelial-specific) E74-like factor 4 (ets domain transcription factor)
ELF4 14925 E74-like factor 5 (ets domain transcription factor) ELF5 14926 ELK1, member of ETS
oncogene family ELK1 14927 ELK3, ETS-domain protein (SRF accessory protein 2) ELK3 14928 ELK4,
ETS-domain protein (SRF accessory protein 1) ELKA 14929 ELM2 and Myb/SANT-like domain
containing 1 ELMSAN1 14930 empty spiracles homeobox 1 EMX1 14931 empty spiracles homeobox 2
EMX2 14932 engrailed homeobox 1 EN1 14933 engrailed homeobox 2 EN2 14934 enolase 1, (alpha)
ENO1 14935 eomesodermin EOMES 14936 endothelial PAS domain protein 1 EPAS1 14937 Ets2
repressor factor ERF 14938 v-ets avian erythroblastosis virus E26 oncogene ERG 14939-14940 homolog
estrogen receptor 1 ESR1 14941 estrogen receptor 2 (ER beta) ESR2 14942 estrogen related receptor
alpha ESRRA 14943 estrogen related receptor beta ESRRB 14944 estrogen related receptor gamma
ESRRG 14945 ESX homeobox 1 ESX1 14946 v-ets avian erythroblastosis virus E26 oncogene ETS1
14947 homolog 1 v-ets avian erythroblastosis virus E26 oncogene ETS2 14948 homolog 2 ets variant 1
ETV1 14949 ets variant 2 ETV2 14950 ets variant 3 ETV3 14951 ets variant 3-like ETV3L 14952 ets
variant 4 ETV4 14953 ets variant 5 ETV5 14954 ets variant 6 ETV6 14955 ets variant 7 ETV7 14956
even-skipped homeobox 1 EVX1 14957 even-skipped homeobox 2 EVX2 14958 enhancer of zeste 1
polycomb repressive complex 2 EZH1 14959 subunit enhancer of zeste 2 polycomb repressive complex 2
EZH2 14960 subunit family with sequence similarity 170 member A FAM170A 14961 Fer3-like bHLH
transcription factor FERD3L 14962 FEV (ETS oncogene family) FEV 14963 FEZ family zinc finger 1
FEZF1 14964 FEZ family zinc finger 2 FEZF2 14965 folliculogenesis specific bHLH transcription factor
FIGLA 14966 FLT3-interacting zinc finger 1 FIZ1 14967 Fli-1 proto-oncogene, ETS transcription factor
FLI1 14968 FBJ murine osteosarcoma viral oncogene homolog FOS 14969 FBJ murine osteosarcoma
viral oncogene homolog B FOSB 14970 FOS like antigen 1 FOSL1 14971 FOS like antigen 2 FOSL2
14972 forkhead box A1 FOXA1 14973 forkhead box A2 FOXA2 14974 forkhead box A3 FOXA3 14975
forkhead box B1 FOXB1 14976 forkhead box B2 FOXB2 14977 forkhead box C1 FOXC1 14978
forkhead box C2 FOXC2 14979 forkhead box D1 FOXD1 14980 forkhead box D2 FOXD2 14981
forkhead box D3 FOXD3 14982 forkhead box D4 FOXD4 14983 forkhead box D4-like 1 FOXD4L1
14984 forkhead box D4-like 3 FOXD4L3 14985 forkhead box D4-like 4 FOXD4L4 14986 forkhead box
D4-like 5 FOXD4L5 14987 forkhead box D4-like 6 FOXD4L6 14988 forkhead box E1 FOXE1 14989
forkhead box E3 FOXE3 14990 forkhead box F1 FOXF1 14991 forkhead box F2 FOXF2 14992 forkhead
box G1 FOXG1 14993 forkhead box H1 FOXH1 14994 forkhead box I1 FOXI1 14995 forkhead box I2
FOXI2 14996 forkhead box I3 FOXI3 14997 forkhead box J1 FOXJ1 14998 forkhead box J2 FOXJ2
14999 forkhead box J3 FOXJ3 15000 forkhead box K1 FOXK1 15001 forkhead box K2 FOXK2 15002
forkhead box L1 FOXL1 15003 forkhead box L2 FOXL2 15004 forkhead box M1 FOXM1 15005
forkhead box N1 FOXN1 15006 forkhead box N2 FOXN2 15007 forkhead box N3 FOXN3 15008
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forkhead box N4 FOXN4 15009 forkhead box O1 FOXO1 15010 forkhead box O3 FOXO3 15011
forkhead box O4 FOXO4 15012 forkhead box O6 FOXO6 15013 forkhead box P1 FOXP1 15014
forkhead box P2 FOXP3 15015 forkhead box P3 FOXP4 15016 forkhead box P4 FOXQ1 15017 forkhead
box Q1 FOXR1 15018 forkhead box R1 FOXR2 15019 forkhead box R2 FOXS1 15020 forkhead box S1
FOXP3 15021 far upstream element binding protein 1 FUBP1 15022 far upstream element (FUSE)
binding protein 3 FUBP3 15023 GA binding protein transcription factor alpha subunit GABPA 15024 GA
binding protein transcription factor, beta subunit 1 GABPB1 15025 GA binding protein transcription
factor, beta subunit 2 GABPB2 15026 GATA binding protein 1 (globin transcription factor 1) GATA1
15027 GATA binding protein 2 GATA2 15028 GATA binding protein 3 GATA3 15029 GATA binding
protein 4 GATA4 15030 GATA binding protein 5 GATA5 15031 GATA binding protein 6 GATA6 15032
GATA zinc finger domain containing 1 GATAD1 15033 GATA zinc finger domain containing 2A
GATAD2A 15034 GATA zinc finger domain containing 2B GATAD2B 15035 gastrulation brain
homeobox 1 GBX1 15036 gastrulation brain homeobox 2 GBX2 15037 GC-rich sequence DNA-binding
factor 2 GCFC2 15038 glial cells missing homolog 1 GCM1 15039 glial cells missing homolog 2 GCM2
15040 growth factor independent 1 transcription repressor GFI1 15041 growth factor independent 1B
transcription repressor GFI1B 15042 GLI family zinc finger 1 GLI1 15043 GLI family zinc finger 2 GLI2
15044 GLI family zinc finger 3 GLI3 15045 GLI family zinc finger 4 GLI4 15046 GLIS family zinc
finger 1 GLIS1 15047 GLIS family zinc finger 2 GLIS2 15048 GLIS family zinc finger 3 GLIS3 15049
glucocorticoid modulatory element binding protein 1 GMEB1 15050 glucocorticoid modulatory element
binding protein 2 GMEB2 15051 gon-4-like (C. elegans) GON4L 15052 grainyhead like transcription
factor 1 GRHL1 15053 grainyhead like transcription factor 2 GRHL2 15054 grainyhead like transcription
factor 3 GRHL3 15055 goosecoid homeobox GSC 15056 goosecoid homeobox 2 GSC2 15057 GS
homeobox 1 GSX1 15058 GS homeobox 2 GSX2 15059 general transcription factor IIi GTF2I 15060
general transcription factor IIIA GTF3A 15061 GDNF inducible zinc finger protein 1 GZF1 15062 heart
and neural crest derivatives expressed 1 HAND1 15063 heart and neural crest derivatives expressed 2
HAND2 15064 HMG-box transcription factor 1 HBP1 15065-15066 highly divergent homeobox HDX
15067 helt bHLH transcription factor HELT 15068 hes family bHLH transcription factor 1 HES1 15069-
15070 hes family bHLH transcription factor 2 HES2 15071 hes family bHLH transcription factor 3 HES3
15072 hes family bHLH transcription factor 4 HES4 15073 hes family bHLH transcription factor 5 HES5
15074 hes family bHLH transcription factor 6 HES6 15075 hes family bHLH transcription factor 7 HES7
15076 HESX homeobox 1 HESX1 15077 hes-related family bHLH transcription factor with HEY1 15078
YRPW motif 1 hes-related family bHLH transcription factor with HEY2 15079 YRPW motif 2 hes-related
family bHLH transcription factor with HEYL 15080 YRPW motif-like hematopoietically expressed
homeobox HHEX 15081 hypermethylated in cancer 1 HIC1 15082 hypermethylated in cancer 2 HIC2
15083 hypoxia inducible factor 1, alpha subunit (basic helix- HIF1A 15084 loop-helix transcription factor)
hypoxia inducible factor 3, alpha subunit HIF3A 15085 histone H4 transcription factor HINFP 15086
human immunodeficiency virus type I enhancer HIVEP1 15087 binding protein 1 human
immunodeficiency virus type I enhancer HIVEP2 15088 binding protein 2 human immunodeficiency virus
type I enhancer HIVEP3 15089 binding protein 3 HKR1, GLI-Kruppel zinc finger family member HKR1
15090 hepatic leukemia factor HLF 15091 helicase-like transcription factor HLTF 15092 H2.0-like
homeobox HLX 15093 homeobox containing 1 HMBOX1 15094 high mobility group 20A HMG20A
15095 high mobility group 20B HMG20B 15096 high mobility group AT-hook 1 HMGA1 15097 high
mobility group AT-hook 2 HMGA2 15098 HMG-box containing 3 HMGXB3 15099 HMG-box containing
4 HMGXB4 15100 H6 family homeobox 1 HMX1 15101 H6 family homeobox 2 HMX2 15102 H6
family homeobox 3 HMX3 15103-15104 HNF1 homeobox A HNF1A 15105 HNF1 homeobox B HNF1B
15106 hepatocyte nuclear factor 4 alpha HNF4A 15107 hepatocyte nuclear factor 4 gamma HNF4G 15108
heterogeneous nuclear ribonucleoprotein K HNRNPK 15109 homeobox and leucine zipper encoding
HOMEZ 15110 HOP homeobox HOPX 15111 homeobox A1 HOXA1 15112 homeobox A10 HOXA10
15113 homeobox A11 HOXA11 15114 homeobox A13 HOXA13 15115 homeobox A2 HOXA2 15116
homeobox A3 HOXA3 15117 homeobox A4 HOXA4 15118 homeobox A5 HOXA5 15119 homeobox A6
HOXA6 15120 homeobox A7 HOXA7 15121 homeobox A9 HOXA9 15122 homeobox B1 HOXB1
15123 homeobox B13 HOXB13 15124 homeobox B2 HOXB2 15125 homeobox B3 HOXB3 15126
homeobox B4 HOXB4 15127 homeobox B5 HOXB5 15128 homeobox B6 HOXB6 15129 homeobox B7
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HOXB7 15130 homeobox B8 HOXB8 15131 homeobox B9 HOXB9 15132 homeobox C10 HOXC10
15133 homeobox C11 HOXC11 15134 homeobox C12 HOXC12 15135 homeobox C13 HOXC13 15136
homeobox C4 HOXC4 15137 homeobox C5 HOXC5 15138 homeobox C6 HOXC6 15139 homeobox C8
HOXC8 15140 homeobox C9 HOXC9 15141 homeobox D1 HOXD1 15142 homeobox D10 HOXD10
15143 homeobox D11 HOXD11 15144 homeobox D12 HOXD12 15145 homeobox D13 HOXD13 15146
homeobox D3 HOXD3 15147 homeobox D4 HOXD4 15148 homeobox D8 HOXD8 15149 homeobox D9
HOXD9 15150 heat shock transcription factor 1 HSF1 15151 heat shock transcription factor 2 HSF2
15152 heat shock transcription factor 4 HSF4 15153 heat shock transcription factor family member 5
HSF5 15154 heat shock transcription factor family, X-linked 1 HSFX1 15155 heat shock transcription
factor, Y-linked 1 HSFY1 15156 heat shock transcription factor, Y-linked 2 HSFY2 15156 inhibitor of
DNA binding 1, dominant negative helix- ID1 15157 loop-helix protein inhibitor of DNA binding 2,
dominant negative helix- ID2 15158 loop-helix protein inhibitor of DNA binding 3, dominant negative
helix- ID3 15159 loop-helix protein inhibitor of DNA binding 4, dominant negative helix- ID4 15160
loop-helix protein interferon, gamma-inducible protein 16 IFI16 15161 IKAROS family zinc finger 1
IKZF1 15162 IKAROS family zinc finger 2 IKZF2 15163 IKAROS family zinc finger 3 IKZF3 15164
IKAROS family zinc finger 4 IKZF4 15165 IKAROS family zinc finger 5 IKZF5 15166 insulinoma
associated 1 INSM1 15167 insulinoma-associated 2 INSM2 15168 interferon regulatory factor 1 IRF1
15169 interferon regulatory factor 2 IRF2 15170 interferon regulatory factor 3 IRF3 15171 interferon
regulatory factor 4 IRF4 15172 interferon regulatory factor 5 IRF5 15173 interferon regulatory factor 6
IRF6 15174 interferon regulatory factor 7 IRF7 15175 interferon regulatory factor 8 IRF8 15176
interferon regulatory factor 9 IRF9 15177 iroquois homeobox 1 IRX1 15178 iroquois homeobox 2 IRX2
15179 iroquois homeobox 3 IRX3 15180 iroquois homeobox 4 IRX4 15181 iroquois homeobox 5 IRX5
15182 iroquois homeobox 6 IRX6 15183 ISL LIM homeobox 1 ISL1 15184 ISL LIM homeobox 2 ISL2
15185 intestine specific homeobox ISX 15186 jumonji and AT-rich interaction domain containing 2
JARID2 15187 JAZF zinc finger 1 JAZF1 15188 Jun dimerization protein 2 JDP2 15189 jun proto-
oncogene JUN 15190 jun B proto-oncogene JUNB 15191 jun D proto-oncogene JUND 15192 K(lysine)
acetyltransferase 5 KAT5 15193 lysine acetyltransferase 6A KAT6A 15194 lysine acetyltransferase 6B
KAT6B 15195 lysine acetyltransferase 7 KAT7 15196 lysine acetyltransferase 8 KAT8 15197 potassium
channel modulatory factor 1 KCMF1 15198 potassium voltage-gated channel interacting protein 3
KCNIP3 15199 lysine demethylase 2A KDM2A 15200 lysine demethylase 5A KDM5A 15201 lysine
demethylase 5B KDM5B 15202 lysine demethylase 5C KDM5C 15203 lysine demethylase 5D KDM5D
15204 KH-type splicing regulatory protein KHSRP 15205 KIAA1549 KIAA1549 15206 Kruppel-like
factor 1 (erythroid) KLF1 15207 Kruppel-like factor 10 KLF10 15208 Kruppel-like factor 11 KLF11
15209 Kruppel-like factor 12 KLF12 15210 Kruppel-like factor 13 KLF13 15211 Kruppel-like factor 14
KLF14 15212 Kruppel-like factor 15 KLF15 15213 Kruppel-like factor 16 KLF16 15214 Kruppel-like
factor 17 KLF17 15215 Kruppel-like factor 2 KLF2 15216 Kruppel-like factor 3 (basic) KLF3 15217
Kruppel-like factor 4 (gut) KLF4 15218 Kruppel-like factor 5 (intestinal) KLF5 15219 Kruppel-like factor
6 KLF6 15220 Kruppel-like factor 7 (ubiquitous) KLF7 15221 Kruppel-like factor 8 KLF8 15222
Kruppel-like factor 9 KLF9 15223 lysine methyltransferase 2A KMT2A 15224 lysine methyltransferase
2B KMT2B 15225 lysine methyltransferase 2C KMT2C 15226 lysine methyltransferase 2E KMT2E
15227 l(3)mbt-like 1 (Drosophila) L3MBTL1 15228 l(3)mbt-like 2 (Drosophila) L3MBTL2 15229
l(3)mbt-like 3 (Drosophila) L3MBTL3 15230 l(3)mbt-like 4 (Drosophila) L3MBTL4 15231 ladybird
homeobox 1 LBX1 15232 ladybird homeobox 2 LBX2 15233 ligand dependent nuclear receptor
corepressor LCOR 15234 ligand dependent nuclear receptor corepressor like LCORL 15235 lymphoid
enhancer binding factor 1 LEF1 15236 leucine twenty homeobox LEUTX 15237 LIM homeobox 1 LHX1
15238 LIM homeobox 2 LHX2 15239 LIM homeobox 3 LHX3 15240 LIM homeobox 4 LHX4 15241
LIM homeobox 5 LHX5 15242 LIM homeobox 6 LHX6 15243 LIM homeobox 8 LHX8 15244 LIM
homeobox 9 LHX9 15245 LIM homeobox transcription factor 1, alpha LMX1A 15246 LIM homeobox
transcription factor 1, beta LMX1B 15247 LOC730110 LOC730110 leucine rich repeat (in FLII)
interacting protein 1 LRRFIP1 15248 leucine rich repeat (in FLII) interacting protein 2 LRRFIP2 15249
Ly1 antibody reactive LYAR 15250 lymphoblastic leukemia associated hematopoiesis LYL1 15251
regulator 1 maelstrom spermatogenic transposon silencer MAEL 15252 v-maf avian musculoaponeurotic
fibrosarcoma MAF 15253 oncogene homolog MAF1 homolog, negative regulator of RNA MAF1 15254
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polymerase III v-maf avian musculoaponeurotic fibrosarcoma MAFA 15255-15256 oncogene homolog A
v-maf avian musculoaponeurotic fibrosarcoma MAFB 15257 oncogene homolog B v-maf avian
musculoaponeurotic fibrosarcoma MAFF 15258 oncogene homolog F v-maf avian musculoaponeurotic
fibrosarcoma MAFG 15259 oncogene homolog G v-maf avian musculoaponeurotic fibrosarcoma MAFK
15260 oncogene homolog K matrin 3 MATR3 15261 MYC associated factor X MAX 15262 MYC
associated zinc finger protein MAZ 15263 methyl-CpG binding domain protein 1 MBD1 15264 methyl-
CpG binding domain protein 2 MBD2 15265 methyl-CpG binding domain protein 3 MBD3 15266
methyl-CpG binding domain protein 3-like 1 MBD3L1 15267 methyl-CpG binding domain protein 3-like
2 MBD3L2 15268 methyl-CpG binding domain 4 DNA glycosylase MBD4 15269 methyl-CpG binding
domain protein 5 MBD5 15270 methyl-CpG binding domain protein 6 MBD6 15271 muscleblind like
splicing regulator 3 MBNL3 15272 MDS1 and EVI1 complex locus MECOM 15273 methyl-CpG binding
protein 2 MECP2 15274 myocyte enhancer factor 2A MEF2A 15275 myocyte enhancer factor 2B MEF2B
15276 myocyte enhancer factor 2C MEF2C 15277 myocyte enhancer factor 2D MEF2D 15278 Meis
homeobox 1 MEIS1 15279 Meis homeobox 2 MEIS2 15280 Meis homeobox 3 MEIS3 15281 Meis
homeobox 3 pseudogene 1 MEIS3P1 15282 Meis homeobox 3 pseudogene 2 MEIS3P2 15283
mesenchyme homeobox 1 MEOX1 15284 mesenchyme homeobox 2 MEOX2 15285 mesoderm posterior
bHLH transcription factor 1 MESP1 15286 mesoderm posterior bHLH transcription factor 2 MESP2
15287 MGA, MAX dimerization protein MGA 15288-15289 MIER1 transcriptional regulator MIER1
15290 MIER family member 2 MIER2 15291 MIER family member 3 MIER3 15292 MIS18 binding
protein 1 MIS18BP1 15293 microphthalmia-associated transcription factor MITF 15294 Mix paired-like
homeobox MIXL1 15295 mohawk homeobox MKX 15296 myeloid/lymphoid or mixed-lineage leukemia;
MLLT1 15297 translocated to, 1 myeloid/lymphoid or mixed-lineage leukemia; MLLT10 15298
translocated to, 10 myeloid/lymphoid or mixed-lineage leukemia; MLLT11 15299 translocated to, 11
myeloid/lymphoid or mixed-lineage leukemia; MLLT3 15300 translocated to, 3 myeloid/lymphoid or
mixed-lineage leukemia; MLLT4 15301 translocated to, 4 myeloid/lymphoid or mixed-lineage leukemia;
MLLT6 15302 translocated to, 6 MLX, MAX dimerization protein MLX 15303 MLX interacting protein
MLXIP 15304 MLX interacting protein-like MLXIPL 15305 MAX network transcriptional repressor
MNT 15306 motor neuron and pancreas homeobox 1 MNX1 15307 musculin MSC 15308 mesogenin 1
MSGN1 15309 msh homeobox 1 MSX1 15310 msh homeobox 2 MSX2 15311 metastasis associated 1
MTA1 15312 metastasis associated 1 family member 2 MTA2 15313 metastasis associated 1 family
member 3 MTA3 15314 metal-regulatory transcription factor 1 MTF1 15315 metal response element
binding transcription factor 2 MTF2 15316 MAX dimerization protein 1 MXD1 15317 MAX dimerization
protein 3 MXD3 15318 MAX dimerization protein 4 MXD4 15319 MAX interactor 1, dimerization
protein MXI1 15320 v-myb avian myeloblastosis viral oncogene homolog MYB 15321 v-myb avian
myeloblastosis viral oncogene homolog- MYBL1 15322 like 1 v-myb avian myeloblastosis viral oncogene
homolog- MYBL2 15323 like 2 v-myc avian myelocytomatosis viral oncogene MYC 15324 homolog v-
myc avian myelocytomatosis viral oncogene lung MYCL 15325 carcinoma derived homolog MYCL
pseudogene 1 MYCLP1 15326 v-myc avian myelocytomatosis viral oncogene MYCN 15327
neuroblastoma derived homolog myogenic factor 5 MYF5 15328 myogenic factor 6 MYF6 15329
myoneurin MYNN 15330 myogenic differentiation 1 MYOD1 15331 myogenin (myogenic factor 4)
MYOG 15332 myelin regulatory factor MYRF 15333 Myb-like, SWIRM and MPN domains 1 MYSM1
15334 myelin transcription factor 1 MYT1 15335-15336 myelin transcription factor 1 like MYT1L 15337
myeloid zinc finger 1 MZF1 15338 Nanog homeobox NANOG 15339 NANOG neighbor homeobox
NANOGNB 15340 Nanog homeobox pseudogene 1 NANOGP1 15341 Nanog homeobox pseudogene 8
NANOGP8 15342 nuclear receptor coactivator 1 NCOA1 15343 nuclear receptor coactivator 2 NCOA2
15344 nuclear receptor coactivator 3 NCOA3 15345 nuclear receptor coactivator 4 NCOA4 15346 nuclear
receptor coactivator 5 NCOA5 15347 nuclear receptor coactivator 6 NCOA6 15348 nuclear receptor
coactivator 7 NCOA7 15349 nuclear receptor corepressor 1 NCOR1 15350 nuclear receptor corepressor 2
NCOR2 15351 neuronal differentiation 1 NEUROD1 15352 neuronal differentiation 2 NEUROD2 15353
neuronal differentiation 4 NEUROD4 15354 neuronal differentiation 6 NEUROD6 15355 neuro genin 1
NEUROG1 15356 neuro genin 2 NEUROG2 15357 neuro genin 3 NEUROG3 15358 nuclear factor of
activated T-cells 5, tonicity- NFAT5 15359 responsive nuclear factor of activated T-cells, cytoplasmic,
NFATC1 15360 calcineurin-dependent 1 nuclear factor of activated T-cells, cytoplasmic, NFATC2 15361
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calcineurin-dependent 2 nuclear factor of activated T-cells, cytoplasmic, NFATC3 15362 calcineurin-
dependent 3 nuclear factor of activated T-cells, cytoplasmic, NFATC4 15363 calcineurin-dependent 4
nuclear factor, erythroid 2 NFE2 15364 nuclear factor, erythroid 2 like 1 NFE2L1 15365 nuclear factor,
erythroid 2 like 2 NFE2L2 15366 nuclear factor, erythroid 2 like 3 NFE2L3 15367 nuclear factor I/A
NFIA 15368 nuclear factor I/B NFIB 15369 nuclear factor I/C (CCAAT-binding transcription NFIC 15370
factor) nuclear factor, interleukin 3 regulated NFIL3 15371 nuclear factor I/X (CCAAT-binding
transcription NFIX 15372 factor) nuclear factor of kappa light polypeptide gene NFKB1 15373 enhancer
in B-cells 1 nuclear factor of kappa light polypeptide gene NFKB2 15374 enhancer in B-cells 2
(p49/p100) nuclear factor of kappa light polypeptide gene NFKBIA 15375 enhancer in B-cells inhibitor,
alpha nuclear factor of kappa light polypeptide gene NFKBIB 15376 enhancer in B-cells inhibitor, beta
nuclear factor of kappa light polypeptide gene NFKBID 15377 enhancer in B-cells inhibitor, delta nuclear
factor of kappa light polypeptide gene NFKBIE 15378 enhancer in B-cells inhibitor, epsilon nuclear factor
of kappa light polypeptide gene NFKBIL1 15379 enhancer in B-cells inhibitor-like 1 nuclear factor of
kappa light polypeptide gene NFKBIZ 15380 enhancer in B-cells inhibitor, zeta nuclear factor related to
kappaB binding protein NFRKB 15381 nuclear transcription factor, X-box binding 1 NFX1 15382 nuclear
transcription factor, X-box binding-like 1 NFXL1 15383 nuclear transcription factor Y subunit alpha
NFYA 15384 nuclear transcription factor Y subunit beta NFYB 15385 nuclear transcription factor Y
subunit gamma NFYC 15386 nescient helix-loop-helix 1 NHLH1 15387 nescient helix-loop-helix 2
NHLH2 15388 NFKB repressing factor NKRF 15389 NK1 homeobox 1 NKX1-1 15390 NK1 homeobox
2 NKX1-2 15391 NK2 homeobox 1 NKX2-1 15392 NK2 homeobox 2 NKX2-2 15393 NK2 homeobox 3
NKX2-3 15394 NK2 homeobox 4 NKX2-4 15395 NK2 homeobox 5 NKX2-5 15396 NK2 homeobox 6
NKX2-6 15397 NK2 homeobox 8 NKX2-8 15398 NK3 homeobox 1 NKX3-1 15399 NK3 homeobox 2
NKX3-2 15400 NK6 homeobox 1 NKX6-1 15401 NK6 homeobox 2 NKX6-2 15402 NK6 homeobox 3
NKX6-3 15403 NOBOX oogenesis homeobox NOBOX 15404 NOC3 like DNA replication regulator
NOC3L 15405 nucleolar complex associated 4 homolog NOC4L 15406 non-POU domain containing,
octamer-binding NONO 15407 notochord homeobox NOTO 15408 neuronal PAS domain protein 1
NPAS1 15409 neuronal PAS domain protein 2 NPAS2 15410 neuronal PAS domain protein 3 NPAS3
15411 neuronal PAS domain protein 4 NPAS4 15412 nuclear receptor subfamily 0 group B member 1
NR0B1 15413 nuclear receptor subfamily 0 group B member 2 NR0B2 15414 nuclear receptor subfamily
1 group D member 1 NR1D1 15415 nuclear receptor subfamily 1 group D member 2 NR1D2 15416
nuclear receptor subfamily 1 group H member 2 NR1H2 15417 nuclear receptor subfamily 1 group H
member 3 NR1H3 15418 nuclear receptor subfamily 1 group H member 4 NR1H4 15419 nuclear receptor
subfamily 1 group I member 2 NR1I2 15420 nuclear receptor subfamily 1 group I member 3 NR1I3 15421
nuclear receptor subfamily 2 group C member 1 NR2C1 15422 nuclear receptor subfamily 2 group C
member 2 NR2C2 15423 nuclear receptor subfamily 2 group E member 1 NR2E1 15424 nuclear receptor
subfamily 2 group E member 3 NR2E3 15425 nuclear receptor subfamily 2 group F member 1 NR2F1
15426 nuclear receptor subfamily 2 group F member 2 NR2F2 15427 nuclear receptor subfamily 2 group
F member 6 NR2F6 15428 nuclear receptor subfamily 3 group C member 1 NR3C1 15429 nuclear
receptor subfamily 3 group C member 2 NR3C2 15430 nuclear receptor subfamily 4 group A member 1
NR4A1 15431 nuclear receptor subfamily 4 group A member 2 NR4A2 15432 nuclear receptor subfamily
4 group A member 3 NR4A3 15433 nuclear receptor subfamily 5 group A member 1 NR5A1 15434
nuclear receptor subfamily 5 group A member 2 NR5A2 15435 nuclear receptor subfamily 6 group A
member 1 NR6A1 15436 nuclear respiratory factor 1 NRF1 15437-15438 neural retina leucine zipper
NRL 15439 oligodendrocyte transcription factor 1 OLIG1 15440 oligodendrocyte lineage transcription
factor 2 OLIG2 15441 oligodendrocyte transcription factor 3 OLIG3 15442 one cut homeobox 1
ONECUT1 15443 one cut homeobox 2 ONECUT2 15444 one cut homeobox 3 ONECUT3 15445 odd-
skipped related transciption factor 1 OSR1 15446 odd-skipped related transciption factor 2 OSR2 15447
orthopedia homeobox OTP 15448 orthodenticle homeobox 1 OTX1 15449 orthodenticle homeobox 2
OTX2 15450 ovo like zinc finger 1 OVOL1 15451 ovo like zinc finger 2 OVOL2 15452 ovo like zinc
finger 3 OVOL3 15453 poly(ADP-ribose) polymerase 1 PARP1 15454 poly(ADP-ribose) polymerase
family member 12 PARP12 15455 POZ/BTB and AT hook containing zinc finger 1 PATZ1 15456 PRKC,
apoptosis, WT1, regulator PAWR 15457 paired box 1 PAX1 15458 paired box 2 PAX2 15459 paired box 3
PAX3 15460 paired box 4 PAX4 15461 paired box 5 PAX5 15462 paired box 6 PAX6 15463 paired box 7
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PAX7 15464 paired box 8 PAX8 15465 paired box 9 PAX9 15466 PAX3 and PAX7 binding protein 1
PAXBP1 15467 polybromo 1 PBRM1 15468 pre-B-cell leukemia homeobox 1 PBX1 15469 pre-B-cell
leukemia homeobox 2 PBX2 15470 pre-B-cell leukemia homeobox 3 PBX3 15471 pre-B-cell leukemia
homeobox 4 PBX4 15472 poly(rC) binding protein 1 PCBP1 15473 poly(rC) binding protein 2 PCBP2
15474 poly(rC) binding protein 3 PCBP3 15475 poly(rC) binding protein 4 PCBP4 15476 poly comb
group ring finger 6 PCGF6 15477 pancreatic and duodenal homeobox 1 PDX1 15478-15479 paternally
expressed 3 PEG3 15480 progesterone receptor PGR 15481 prohibitin PHB 15482 prohibitin 2 PHB2
15483 PHD finger protein 20 PHF20 15484 PHD finger protein 5A PHF5A 15485 paired like homeobox
2a PHOX2A 15486 paired like homeobox 2b PHOX2B 15487 putative homeodomain transcription factor
1 PHTF1 15488 putative homeodomain transcription factor 2 PHTF2 15489 paired like homeodomain 1
PITX1 15490 paired like homeodomain 2 PITX2 15491 paired like homeodomain 3 PITX3 15492
PBX/knotted 1 homeobox 1 PKNOX1 15493 PBX/knotted 1 homeobox 2 PKNOX2 15494 PLAG1 zinc
finger PLAG1 15495 PLAG1 like zinc finger 1 PLAGL1 15496 PLAG1 like zinc finger 2 PLAGL2 15497
pleckstrin PLEK 15498 promyelocytic leukaemia zinc finger PLZF 15499 pogo transposable element with
ZNF domain POGZ 15500 POU class 1 homeobox 1 POU1F1 15501 POU class 2 associating factor 1
POU2AF1 15502 POU class 2 homeobox 1 POU2F1 15503 POU class 2 homeobox 2 POU2F2 15504
POU class 2 homeobox 3 POU2F3 15505 POU class 3 homeobox 1 POU3F1 15506 POU class 3
homeobox 2 POU3F2 15507 POU class 3 homeobox 3 POU3F3 15508 POU class 3 homeobox 4 POU3F4
15509 POU class 4 homeobox 1 POU4F1 15510 POU class 4 homeobox 2 POU4F2 15511 POU class 4
homeobox 3 POU4F3 15512 POU class 5 homeobox 1 POU5F1 15513 POU class 5 homeobox 1B
POU5F1B 15514 POU domain class 5, transcription factor 2 POU5F2 15515 POU class 6 homeobox 1
POU6F1 15516 POU class 6 homeobox 2 POU6F2 15517 peroxisome proliferator activated receptor alpha
PPARA 15518 peroxisome proliferator activated receptor delta PPARD 15519 peroxisome proliferator
activated receptor gamma PPARG 15520 protein phosphatase 1 regulatory subunit 13 like PPP1R13L
15521 PR domain 1 PRDM1 15522 PR domain 10 PRDM10 15523 PR domain 11 PRDM11 15524 PR
domain 12 PRDM12 15525 PR domain 13 PRDM13 15526 PR domain 14 PRDM14 15527 PR domain 15
PRDM15 15528 PR domain 16 PRDM16 15529 PR domain 2 PRDM2 15530 PR domain 4 PRDM4
15531 PR domain 5 PRDM5 15532 PR domain 6 PRDM6 15533 PR domain 7 PRDM7 15534 PR domain
8 PRDM8 15535 PR domain 9 PRDM9 15536 prolactin regulatory element binding PREB 15537 PROP
paired-like homeobox 1 PROP1 15538 prospero homeobox 1 PROX1 15539 prospero homeobox 2
PROX2 15540 paired related homeobox 1 PRRX1 15541 paired related homeobox 2 PRRX2 15542
paraspeckle component 1 PSPC1 15543 pancreas specific transcription factor, 1a PTF1A 15544 purine-
rich element binding protein A PURA 15545 purine-rich element binding protein B PURB 15546 purine-
rich element binding protein G PURG 15547 retinoic acid receptor alpha RARA 15548 retinoic acid
receptor beta RARB 15549 retinoic acid receptor gamma RARG 15550 retina and anterior neural fold
homeobox RAX 15551-15552 retina and anterior neural fold homeobox 2 RAX2 15553 RB associated
KRAB zinc finger RBAK 15554 RNA binding motif protein 22 RBM22 15555 recombination signal
binding protein for RBPJ 15556 immunoglobulin kappa J region recombination signal binding protein for
RBPJL 15557 immunoglobulin kappa J region-like ring finger and CCCH-type domains 1 RC3H1 15558
ring finger and CCCH-type domains 2 RC3H2 15559 REST corepressor 1 RCOR1 15560 REST
corepressor 2 RCOR2 15561 REST corepressor 3 RCOR3 15562 v-rel avian reticuloendothcliosis viral
oncogene REL 15563 homolog v-rel avian reticuloendothcliosis viral oncogene RELA 15564 homolog A
v-rel avian reticuloendothcliosis viral oncogene RELB 15565 homolog B arginine-glutamic acid di peptide
(RE) repeats RERE 15566 RE1-silencing transcription factor REST 15567 regulatory factor X1 RFX1
15568 regulatory factor X2 RFX2 15569 regulatory factor X3 RFX3 15570 regulatory factor X4 RFX4
15571 regulatory factor X5 RFX5 15572 regulatory factor X6 RFX6 15573 regulatory factor X7 RFX7
15574 RFX family member 8, lacking RFX DNA binding RFX8 15575 domain regulatory factor X
associated ankyrin containing RFXANK 15576 protein regulatory factor X associated protein RFXAP
15577 Rhox homeobox family member 1 RHOXF1 15578 Rhox homeobox family member 2 RHOXF2
15579 Rhox homeobox family member 2B RHOXF2B 15580 rearranged L-myc fusion RLF 15581-15582
RAR related orphan receptor A RORA 15583 RAR related orphan receptor B RORB 15584 RAR related
orphan receptor C RORC 15585 retinoic acid receptor-related orphan nuclear receptor RORgT 15586
gamma ras responsive element binding protein 1 RREB1 15587 runt related transcription factor 1 RUNX1
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15588 runt related transcription factor 1; translocated to, 1 RUNX1T1 15589 (cyclin D related) runt
related transcription factor 2 RUNX2 15590 runt related transcription factor 3 RUNX3 15591 retinoid X
receptor alpha RXRA 15592 retinoid X receptor beta RXRB 15593 retinoid X receptor gamma RXRG
15594 spalt-like transcription factor 1 SALL1 15595 spalt-like transcription factor 2 SALL2 15596 spalt-
like transcription factor 3 SALL3 15597 spalt-like transcription factor 4 SALL4 15598 SATB homeobox 1
SATB1 15599 SATB homeobox 2 SATB2 15600 S-phase cyclin A-associated protein in the ER SCAPER
15601 scratch family zinc finger 1 SCRT1 15602 scratch family zinc finger 2 SCRT2 15603 scleraxis
bHLH transcription factor SCX 15604 SEBOX homeobox SEBOX 15605 SET binding protein 1 SETBP1
15606 splicing factor proline/glutamine-rich SFPQ 15607 short stature homeobox SHOX 15608 short
stature homeobox 2 SHOX2 15609 single-minded family bHLH transcription factor 1 SIM1 15610 single-
minded family bHLH transcription factor 2 SIM2 15611 SIX homeobox 1 SIX1 15612 SIX homeobox 2
SIX2 15613 SIX homeobox 3 SIX3 15614 SIX homeobox 4 SIX4 15615 SIX homeobox 5 SIX5 15616
SIX homeobox 6 SIX6 15617 SKI proto-oncogene SKI 15618 SKI-like proto-oncogene SKIL 15619 SKI
family transcriptional corepressor 1 SKOR1 15620 SKI family transcriptional corepressor 2 SKOR2
15621 solute carrier family 30 (zinc transporter), member 9 SLC30A9 15622 SMAD family member 1
SMAD1 15623 SMAD family member 2 SMAD2 15624 SMAD family member 3 SMAD3 15625 SMAD
family member 4 SMAD4 15626 SMAD family member 5 SMAD5 15627 SMAD family member 6
SMAD6 15628 SMAD family member 7 SMAD7 15629 SMAD family member 9 SMAD9 15630
SWI/SNF related, matrix associated, actin dependent SMARCA1 15631 regulator of chromatin, subfamily
a, member 1 SWI/SNF related, matrix associated, actin dependent SMARCA2 15632 regulator of
chromatin, subfamily a, member 2 SWI/SNF related, matrix associated, actin dependent SMARCA4
15633 regulator of chromatin, subfamily a, member 4 SWI/SNF related, matrix associated, actin
dependent SMARCA5 15634 regulator of chromatin, subfamily a, member 5 SWI/SNF-related, matrix-
associated actin-dependent SMARCAD1 15635 regulator of chromatin, subfamily a, containing DEAD/H
box 1 SWI/SNF related, matrix associated, actin dependent SMARCAL1 15636 regulator of chromatin,
subfamily a-like 1 SWI/SNF related, matrix associated, actin dependent SMARCB1 15637 regulator of
chromatin, subfamily b, member 1 SWI/SNF related, matrix associated, actin dependent SMARCC1
15638 regulator of chromatin, subfamily c, member 1 SWI/SNF related, matrix associated, actin
dependent SMARCC2 15639 regulator of chromatin, subfamily c, member 2 SWI/SNF related, matrix
associated, actin dependent SMARCD1 15640 regulator of chromatin, subfamily d, member 1 SWI/SNF
related, matrix associated, actin dependent SMARCD2 15641 regulator of chromatin, subfamily d,
member 2 SWI/SNF related, matrix associated, actin dependent SMARCD3 15642 regulator of chromatin,
subfamily d, member 3 SWI/SNF related, matrix associated, actin dependent SMARCE1 15643 regulator
of chromatin, subfamily e, member 1 snail family zinc finger 1 SNAI1 15644 snail family zinc finger 2
SNAI2 15645 snail family zinc finger 3 SNAI3 15646 small nuclear RNA activating complex polypeptide
4 SNAPC4 15647 spermatogenesis and oogenesis specific basic helix- SOHLH1 15648 loop-helix 1
spermatogenesis and oogenesis specific basic helix- SOHLH2 15649 loop-helix 2 SRY-box 1 SOX1 15650
SRY-box 10 SOX10 15651 SRY-box 11 SOX11 15652 SRY-box 12 SOX12 15653 SRY-box 13 SOX13
15654 SRY-box 14 SOX14 15655 SRY-box 15 SOX15 15656 SRY-box 17 SOX17 15657 SRY-box 18
SOX18 15658 SRY-box 2 SOX2 15659 SRY-box 21 SOX21 15660 SRY-box 3 SOX3 15661 SRY-box 30
SOX30 15662 SRY-box 4 SOX4 15663 SRY-box 5 SOX5 15664 SRY-box 6 SOX6 15665 SRY-box 7
SOX7 15666 SRY-box 8 SOX8 15667 SRY-box 9 SOX9 15668 Sp1 transcription factor SP1 15669-15670
SP100 nuclear antigen SP100 15671 SP110 nuclear body protein SP110 15672 SP140 nuclear body
protein SP140 15673 SP140 nuclear body protein like SP140L 15674 Sp2 transcription factor SP2 15675
Sp3 transcription factor SP3 15676 Sp4 transcription factor SP4 15677 Sp5 transcription factor SP5 15678
Sp6 transcription factor SP6 15679 Sp7 transcription factor SP7 15680 Sp8 transcription factor SP8 15681
Sp9 transcription factor SP9 15682 SAM pointed domain containing ETS transcription SPDEF 15683
factor Spi-1 proto-oncogene SPI1 15684 Spi-B transcription factor (Spi-1/PU.1 related) SPIB 15685 Spi-C
transcription factor (Spi-1/PU.1 related) SPIC 15686 spermatogenic leucine zipper 1 SPZ1 15687 sterol
regulatory element binding transcription factor 1 SREBF1 15688 sterol regulatory element binding
transcription factor 2 SREBF2 15689 serum response factor SRF 15690 sex determining region Y SRY
15691 structure specific recognition protein 1 SSRP1 15692 suppression of tumorigenicity 18, zinc finger
ST18 15693 signal transducer and activator of transcription 1 STAT1 15694 signal transducer and
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activator of transcription 2 STAT2 15695 signal transducer and activator of transcription 3 STAT3 15696
(acute-phase response factor) signal transducer and activator of transcription 4 STAT4 15697 signal
transducer and activator of transcription 5 STAT5 15698 signal transducer and activator of transcription
5A STAT5A 15699 signal transducer and activator of transcription 5B STAT5B 15700 signal transducer
and activator of transcription 6, STAT6 15701 interleukin-4 induced transcriptional adaptor 2A TADA2A
15702 transcriptional adaptor 2B TADA2B 15703 TATA-box binding protein associated factor 1 TAF1
15704 T-cell acute lymphocytic leukemia 1 TAL1 15705 T-cell acute lymphocytic leukemia 2 TAL2 15706
Tax1 (human T-cell leukemia virus type I) binding TAX1BP1 15707 protein 1 Tax1 (human T-cell
leukemia virus type I) binding TAX1BP3 15708 protein 3 T-box transcription factor T-bet Tbet 15709
TATA-box binding protein TBP 15710 TATA-box binding protein like 1 TBPL1 15711 TATA-box binding
protein like 2 TBPL2 15712 T-box, brain 1 TBR1 15713 T-box 1 TBX1 15714 T-box 10 TBX10 15715 T-
box 15 TBX15 15716 T-box 18 TBX18 15717 T-box 19 TBX19 15718 T-box 2 TBX2 15719 T-box 20
TBX20 15720 T-box 21 TBX21 15721 T-box 22 TBX22 15722 T-box 3 TBX3 15723 T-box 4 TBX4
15724 T-box 5 TBX5 15725 T-box 6 TBX6 15726 transcription factor 12 TCF12 15727 transcription
factor 15 (basic helix-loop-helix) TCF15 15728 transcription factor 19 TCF19 15729 transcription factor
20 (AR1) TCF20 15730 transcription factor 21 TCF21 15731 transcription factor 23 TCF23 15732
transcription factor 24 TCF24 15733 transcription factor 25 (basic helix-loop-helix) TCF25 15734
transcription factor 3 TCF3 15735 transcription factor 4 TCF4 15736 transcription factor 7 (T-cell specific,
HMG-box, TCF7 15737 TCF1) transcription factor 7 like 1 TCF7L1 15738 transcription factor 7 like 2
TCF7L2 15739 transcription factor-like 5 (basic helix-loop-helix) TCFL5 15740 TEA domain
transcription factor 1 TEAD1 15741 TEA domain transcription factor 2 TEAD2 15742 TEA domain
transcription factor 3 TEAD3 15743 TEA domain transcription factor 4 TEAD4 15744 thyrotrophic
embryonic factor TEF 15745 telomeric repeat binding factor (NIMA-interacting) 1 TERF1 15746
telomeric repeat binding factor 2 TERF2 15747 tet methylcytosine dioxygenase 1 TET1 15748 tet
methylcytosine dioxygenase 2 TET2 15749 tet methylcytosine dioxygenase 3 TET3 15750 transcription
factor A, mitochondrial TFAM 15751 transcription factor AP-2 alpha (activating enhancer TFAP2A 15752
binding protein 2 alpha) transcription factor AP-2 beta (activating enhancer TFAP2B 15753 binding
protein 2 beta) transcription factor AP-2 gamma (activating enhancer TFAP2C 15754 binding protein 2
gamma) transcription factor AP-2 delta (activating enhancer TFAP2D 15755 binding protein 2 delta)
transcription factor AP-2 epsilon (activating enhancer TFAP2E 15756 binding protein 2 epsilon)
transcription factor AP-4 (activating enhancer binding TFAP4 15757 protein 4) transcription factor B1,
mitochondrial TFB1M 15758 transcription factor B2, mitochondrial TFB2M 15759 transcription factor
CP2 TFCP2 15760 transcription factor CP2-like 1 TFCP2L1 15761 transcription factor Dp-1 TFDP1
15762 transcription factor Dp-2 (E2F dimerization partner 2) TFDP2 15763 transcription factor Dp family
member 3 TFDP3 15764 transcription factor binding to IGHM enhancer 3 TFE3 15765 transcription
factor EB TFEB 15766 transcription factor EC TFEC 15767 TGFB induced factor homeobox 1 TGIF1
15768 TGFB induced factor homeobox 2 TGIF2 15769 TGFB induced factor homeobox 2 like, X-linked
TGIF2LX 15770 TGFB induced factor homeobox 2 like, Y-linked TGIF2LY 15771 THAP domain
containing, apoptosis associated protein THAP1 15772 1 THAP domain containing 10 THAP10 15773
THAP domain containing 11 THAP11 15774 THAP domain containing 12 THAP12 15775 THAP domain
containing, apoptosis associated protein THAP2 15776 2 THAP domain containing, apoptosis associated
protein THAP3 15777 3 THAP domain containing 4 THAP4 15778 THAP domain containing 5 THAP5
15779 THAP domain containing 6 THAP6 15780 THAP domain containing 7 THAP7 15781 THAP
domain containing 8 THAP8 15782 THAP domain containing 9 THAP9 15783 Th inducing POZ-Kruppel
Factor ThPOK 15784 thyroid hormone receptor, alpha THRA 15785 thyroid hormone receptor, beta
THRB 15786 T-cell leukemia homeobox 1 TLX1 15787 T-cell leukemia homeobox 2 TLX2 15788 T-cell
leukemia homeobox 3 TLX3 15789 target of EGR1, member 1 (nuclear) TOE1 15790 tonsoku-like, DNA
repair protein TONSL 15791 topoisomerase I binding, arginine/serine-rich, E3 TOPORS 15792 ubiquitin
protein ligase thymocyte selection associated high mobility group TOX 15793 box TOX high mobility
group box family member 2 TOX2 15794 TOX high mobility group box family member 3 TOX3 15795
TOX high mobility group box family member 4 TOX4 15796 tumor protein p53 TP53 15797 tumor
protein p63 TP63 15798 tumor protein p73 TP73 15799 tetra-peptide repeat homeobox 1 TPRX1 15800
tetra-peptide repeat homeobox-like TPRXL 15801 transcriptional regulating factor 1 TRERF1 15802
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trichorhinophalangeal syndrome I TRPS1 15803 TSC22 domain family member 1 TSC22D1 15804
TSC22 domain family member 2 TSC22D2 15805 TSC22 domain family member 3 TSC22D3 15806
TSC22 domain family member 4 TSC22D4 15807 teashirt zinc finger homeobox 1 TSHZ1 15808 teashirt
zinc finger homeobox 2 TSHZ2 15809 teashirt zinc finger homeobox 3 TSHZ3 15810 transcription
termination factor, RNA polymerase I TTF1 15811-15812 transcription termination factor, RNA
polymerase II TTF2 15813-15814 tubby bipartite transcription factor TUB 15815 twist family bHLH
transcription factor 1 TWIST1 15816 twist family bHLH transcription factor 2 TWIST2 15817 upstream
binding protein 1 (LBP-1a) UBP1 15818 upstream binding transcription factor, RNA UBTF 15819
polymerase I upstream binding transcription factor, RNA UBTFL1 15820 polymerase I-like 1 upstream
binding transcription factor, RNA UBTFL6 15821 polymerase I-like 6 (pseudogene) UNC homeobox
UNCX 15822 unkempt family zinc finger UNK 15823 unkempt family like zinc finger UNKL 15824
upstream transcription factor 1 USF1 15825 upstream transcription factor 2, c-fos interacting USF2 15826
upstream transcription factor family member 3 USF3 15827 undifferentiated embryonic cell transcription
factor 1 UTF1 15828 ventral anterior homeobox 1 VAX1 15829 ventral anterior homeobox 2 VAX2 15830
vitamin D (1,25-dihydroxyvitamin D3) receptor VDR 15831 VENT homeobox VENTX 15832 vascular
endothelial zinc finger 1 VEZF1 15833 visual system homeobox 1 VSX1 15834 visual system homeobox
2 VSX2 15835 WD repeat and HMG-box DNA binding protein 1 WDHD1 15836 Wolf-Hirschhorn
syndrome candidate 1 WHSC1 15837 widely interspaced zinc finger motifs WIZ 15838 Wilms tumor 1
WT1 15839 X-box binding protein 1 XBP1 15840 Y-box binding protein 1 YBX1 15841 Y-box binding
protein 2 YBX2 15842 Y-box binding protein 3 YBX3 15843 YEATS domain containing 2 YEATS2
15844 YEATS domain containing 4 YEATS4 15845 YY1 transcription factor YY1 15846 YY2
transcription factor YY2 15847 zinc finger BED-type containing 1 ZBED1 15848 zinc finger BED-type
containing 2 ZBED2 15849 zinc finger BED-type containing 3 ZBED3 15850 zinc finger BED-type
containing 4 ZBED4 15851 zinc finger BED-type containing 5 ZBED5 15852 zinc finger, BED-type
containing 6 ZBED6 15853 Z-DNA binding protein 1 ZBP1 15854-15855 zinc finger and BTB domain
containing 1 ZBTB1 15856 zinc finger and BTB domain containing 10 ZBTB10 15857 zinc finger and
BTB domain containing 11 ZBTB11 15858 zinc finger and BTB domain containing 12 ZBTB12 15859
zinc finger and BTB domain containing 14 ZBTB14 15860 zinc finger and BTB domain containing 16
ZBTB16 15861 zinc finger and BTB domain containing 17 ZBTB17 15862 zinc finger and BTB domain
containing 18 ZBTB18 15863 zinc finger and BTB domain containing 2 ZBTB2 15864 zinc finger and
BTB domain containing 20 ZBTB20 15865 zinc finger and BTB domain containing 21 ZBTB21 15866
zinc finger and BTB domain containing 22 ZBTB22 15867 zinc finger and BTB domain containing 24
ZBTB24 15868 zinc finger and BTB domain containing 25 ZBTB25 15869 zinc finger and BTB domain
containing 26 ZBTB26 15870 zinc finger and BTB domain containing 3 ZBTB3 15871 zinc finger and
BTB domain containing 32 ZBTB32 15872 zinc finger and BTB domain containing 33 ZBTB33 15873
zinc finger and BTB domain containing 34 ZBTB34 15874 zinc finger and BTB domain containing 37
ZBTB37 15875 zinc finger and BTB domain containing 38 ZBTB38 15876 zinc finger and BTB domain
containing 39 ZBTB39 15877 zinc finger and BTB domain containing 4 ZBTB4 15878 zinc finger and
BTB domain containing 40 ZBTB40 15879 zinc finger and BTB domain containing 41 ZBTB41 15880
zinc finger and BTB domain containing 42 ZBTB42 15881 zinc finger and BTB domain containing 43
ZBTB43 15882 zinc finger and BTB domain containing 44 ZBTB44 15883 zinc finger and BTB domain
containing 45 ZBTB45 15884 zinc finger and BTB domain containing 46 ZBTB46 15885 zinc finger and
BTB domain containing 47 ZBTB47 15886 zinc finger and BTB domain containing 48 ZBTB48 15887
zinc finger and BTB domain containing 49 ZBTB49 15888 zinc finger and BTB domain containing 5
ZBTB5 15889 zinc finger and BTB domain containing 6 ZBTB6 15890 zinc finger and BTB domain
containing 7A ZBTB7A 15891 zinc finger and BTB domain containing 7B ZBTB7B 15892 zinc finger
and BTB domain containing 7C ZBTB7C 15893 zinc finger and BTB domain containing 8A ZBTB8A
15894 zinc finger and BTB domain containing 9 ZBTB9 15895 zinc finger CCCH-type containing 10
ZC3H10 15896 zinc finger CCCH-type containing 11A ZC3H11A 15897 zinc finger CCCH-type
containing 12A ZC3H12A 15898 zinc finger CCCH-type containing 12B ZC3H12B 15899 zinc finger
CCCH-type containing 13 ZC3H13 15900 zinc finger CCCH-type containing 14 ZC3H14 15901 zinc
finger CCCH-type containing 15 ZC3H15 15902 zinc finger CCCH-type containing 18 ZC3H18 15903
zinc finger CCCH-type containing 3 ZC3H3 15904 zinc finger CCCH-type containing 4 ZC3H4 15905
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zinc finger CCCH-type containing 6 ZC3H6 15906 zinc finger CCCH-type containing 7A ZC3H7A 15907
zinc finger CCCH-type containing 7B ZC3H7B 15908 zinc finger CCCH-type containing 8 ZC3H8 15909
zinc finger CCHC-type containing 11 ZCCHC11 15910 zinc finger CCHC-type containing 6 ZCCHC6
15911 zinc finger E-box binding homeobox 1 ZEB1 15912 zinc finger E-box binding homeobox 2 ZEB2
15913 zinc finger and AT-hook domain containing ZFAT 15914 zinc finger homeobox 2 ZFHX2 15915
zinc finger homeobox 3 ZFHX3 15916 zinc finger homeobox 4 ZFHX4 15917 ZFP1 zinc finger protein
ZFP1 15918 ZFP14 zinc finger protein ZFP14 15919 ZFP2 zinc finger protein ZFP2 15920 ZFP28 zinc
finger protein ZFP28 15921 ZFP3 zinc finger protein ZFP3 15922 ZFP30 zinc finger protein ZFP30
15923 ZFP36 ring finger protein-like 1 ZFP36L1 15924 ZFP36 ring finger protein-like 2 ZFP36L2 15925
ZFP37 zinc finger protein ZFP37 15926 ZFP41 zinc finger protein ZFP41 15927 ZFP42 zinc finger
protein ZFP42 15928 ZFP57 zinc finger protein ZFP57 15929 ZFP62 zinc finger protein ZFP62 15930
ZFP64 zinc finger protein ZFP64 15931 ZFP69 zinc finger protein ZFP69 15932-15933 ZFP69 zinc finger
protein B ZFP69B 15934 ZFP82 zinc finger protein ZFP82 15935 ZFP90 zinc finger protein ZFP90 15936
ZFP91 zinc finger protein ZFP91 15937 ZFP92 zinc finger protein ZFP92 15938 zinc finger protein, FOG
family member 1 ZFPM1 15939 zinc finger protein, FOG family member 2 ZFPM2 15940 zinc finger
protein, X-linked ZFX 15941 zinc finger protein, Y-linked ZFY 15942 zinc finger, FYVE domain
containing 26 ZFYVE26 15943 zinc finger, GATA-like protein 1 ZGLP1 15944 zinc finger CCCH-type
and G-patch domain ZGPAT 15945 containing zinc fingers and homeoboxes 1 ZHX1 15946 zinc fingers
and homeoboxes 2 ZHX2 15947 zinc fingers and homeoboxes 3 ZHX3 15948 Zic family member 1 ZIC1
15949 Zic family member 2 ZIC2 15950 Zic family member 3 ZIC3 15951 Zic family member 4 ZIC4
15952 Zic family member 5 ZIC5 15953 zinc finger protein interacting with K protein 1 ZIK1 15954 zinc
finger, imprinted 2 ZIM2 15955 zinc finger, imprinted 3 ZIM3 15956 zinc finger with KRAB and SCAN
domains 1 ZKSCAN1 15957 zinc finger with KRAB and SCAN domains 2 ZKSCAN2 15958 zinc finger
with KRAB and SCAN domains 3 ZKSCAN3 15959 zinc finger with KRAB and SCAN domains 4
ZKSCAN4 15960 zinc finger with KRAB and SCAN domains 5 ZKSCAN5 15961 zinc finger with
KRAB and SCAN domains 7 ZKSCAN7 15962 zinc finger with KRAB and SCAN domains 8 ZKSCAN8
15963 zinc finger matrin-type 1 ZMAT1 15964 zinc finger matrin-type 2 ZMAT2 15965 zinc finger
matrin-type 3 ZMAT3 15966 zinc finger matrin-type 4 ZMAT4 15967 zinc finger matrin-type 5 ZMAT5
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ZNF101 15971 zinc finger protein 106 ZNF106 15972 zinc finger protein 107 ZNF107 15973 zinc finger
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protein 154 ZNF154 15995 zinc finger protein 155 ZNF155 15996 zinc finger protein 157 ZNF157 15997
zinc finger protein 16 ZNF16 15998 zinc finger protein 160 ZNF160 15999 zinc finger protein 165
ZNF165 16000 zinc finger protein 169 ZNF169 16001 zinc finger protein 17 ZNF17 16002 zinc finger
protein 174 ZNF174 16003 zinc finger protein 175 ZNF175 16004 zinc finger protein 18 ZNF18 16005
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ZNF182 16008 zinc finger protein 184 ZNF184 16009 zinc finger protein 189 ZNF189 16010 zinc finger
protein 19 ZNF19 16011 zinc finger protein 195 ZNF195 16012 zinc finger protein 197 ZNF197 16013
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ZNF200 16017 zinc finger protein 202 ZNF202 16018 zinc finger protein 205 ZNF205 16019 zinc finger
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ZNF214 16025 zinc finger protein 215 ZNF215 16026 zinc finger protein 217 ZNF217 16027 zinc finger
protein 219 ZNF219 16028 zinc finger protein 22 ZNF22 16029 zinc finger protein 221 ZNF221 16030
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ZNF225 16033-16034 zinc finger protein 226 ZNF226 16035 zinc finger protein 227 ZNF227 16036 zinc
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16046 zinc finger protein 239 ZNF239 16047 zinc finger protein 24 ZNF24 16048 zinc finger protein 248
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protein 251 ZNF251 16052 zinc finger protein 252, pseudogene ZNF252P 16053 zinc finger protein 253
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protein 257 ZNF257 16057 zinc finger protein 26 ZNF26 16058 zinc finger protein 260 ZNF260 16059
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ZNF266 16062 zinc finger protein 267 ZNF267 16063 zinc finger protein 268 ZNF268 16064 zinc finger
protein 273 ZNF273 16065 zinc finger protein 274 ZNF274 16066 zinc finger protein 275 ZNF275 16067
zinc finger protein 276 ZNF276 16068 zinc finger protein 277 ZNF277 16069 zinc finger protein 28
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finger protein 280C ZNF280C 16073 zinc finger protein 280D ZNF280D 16074 zinc finger protein 281
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finger protein 30 ZNF30 16086 zinc finger protein 300 ZNF300 16087 zinc finger protein 302 ZNF302
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finger protein 319 ZNF319 16094 zinc finger protein 32 ZNF32 16095 zinc finger protein 320 ZNF320
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324B ZNF324B 16099 zinc finger protein 326 ZNF326 16100 zinc finger protein 329 ZNF329 16101 zinc
finger protein 331 ZNF331 16102 zinc finger protein 333 ZNF333 16103 zinc finger protein 334 ZNF334
16104 zinc finger protein 335 ZNF335 16105 zinc finger protein 337 ZNF337 16106 zinc finger protein
33A ZNF33A 16107 zinc finger protein 33B ZNF33B 16108 zinc finger protein 34 ZNF34 16109 zinc
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16112 zinc finger protein 346 ZNF346 16113 zinc finger protein 347 ZNF347 16114 zinc finger protein
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finger protein 354B ZNF354B 16118 zinc finger protein 354C ZNF354C 16119 zinc finger protein 355,
pseudogene ZNF355P 16120 zinc finger protein 358 ZNF358 16121 zinc finger protein 362 ZNF362
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protein 367 ZNF367 16126 zinc finger protein 37A ZNF37A 16127 zinc finger protein 382 ZNF382
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16232 zinc finger protein 542, pseudogene ZNF542P 16233 zinc finger protein 543 ZNF543 16234 zinc
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finger protein 704 ZNF704 16363 zinc finger protein 705A ZNF705A 16364 zinc finger protein 705D
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protein 765 ZNF765 16402 zinc finger protein 766 ZNF766 16403 zinc finger protein 768 ZNF768 16404
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protein 774 ZNF774 16410 zinc finger protein 775 ZNF775 16411 zinc finger protein 776 ZNF776 16412
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ZNF780A 16415 zinc finger protein 780B ZNF780B 16416 zinc finger protein 781 ZNF781 16417 zinc
finger protein 782 ZNF782 16418 zinc finger family member 783 ZNF783 16419 zinc finger protein 784
ZNF784 16420 zinc finger protein 785 ZNF785 16421 zinc finger protein 786 ZNF786 16422 zinc finger
protein 787 ZNF787 16423 zinc finger family member 788 ZNF788 16424 zinc finger protein 789
ZNF789 16425 zinc finger protein 79 ZNF79 16426 zinc finger protein 790 ZNF790 16427 zinc finger
protein 791 ZNF791 16428 zinc finger protein 792 ZNF792 16429 zinc finger protein 793 ZNF793 16430
zinc finger protein 799 ZNF799 16431 zinc finger protein 8 ZNF8 16432 zinc finger protein 80 ZNF80
16433 zinc finger protein 800 ZNF800 16434 zinc finger protein 804A ZNF804A 16435 zinc finger
protein 804B ZNF804B 16436 zinc finger protein 805 ZNF805 16437 zinc finger protein 806 ZNF806
16438 zinc finger protein 808 ZNF808 16439 zinc finger protein 81 ZNF81 16440 zinc finger protein 813
ZNF813 16441 zinc finger protein 814 ZNF814 16442 zinc finger protein 816 ZNF816 16443 zinc finger
protein 821 ZNF821 16444 zinc finger protein 823 ZNF823 16445 zinc finger protein 827 ZNF827 16446
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837 ZNF837 16454 zinc finger protein 839 ZNF839 16455 zinc finger protein 84 ZNF84 16456 zinc
finger protein 840, pseudogene ZNF840P 16457 zinc finger protein 841 ZNF841 16458 zinc finger protein
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843 ZNF843 16459 zinc finger protein 844 ZNF844 16460 zinc finger protein 845 ZNF845 16461 zinc finger protein 846 ZNF846 16462 zinc finger protein 85 ZNF85 16463 zinc finger protein 853 ZNF853 16464 zinc finger protein 860 ZNF860 16465 zinc finger protein 876, pseudogene ZNF876P 16466 zinc finger protein 878 ZNF878 16467 zinc finger protein 879 ZNF879 16468 zinc finger protein 880 ZNF880 16469 zinc finger protein 891 ZNF891 16470 zinc finger protein 90 ZNF90 16471 zinc finger protein 91 ZNF91 16472 zinc finger protein 92 ZNF92 16473 zinc finger protein 93 ZNF93 16474 zinc finger protein 98 ZNF98 16475 zinc finger protein 99 ZNF99 16476 zinc finger, NFX1-type containing 1 ZNFX1 16477 zinc finger and SCAN domain containing 1 ZSCAN1 16478 zinc finger and SCAN domain containing 10 ZSCAN10 16479 zinc finger and SCAN domain containing 12 ZSCAN12 16480 zinc finger and SCAN domain containing 16 ZSCAN16 16481 zinc finger and SCAN domain containing 18 ZSCAN18 16482 zinc finger and SCAN domain containing 2 ZSCAN2 16483 zinc finger and SCAN domain containing 20 ZSCAN20 16484 zinc finger and SCAN domain containing 21 ZSCAN21 16485 zinc finger and SCAN domain containing 22 ZSCAN22 16486 zinc finger and SCAN domain containing 23 ZSCAN23 16487 zinc finger and SCAN domain containing 25 ZSCAN25 16488 zinc finger and SCAN domain containing 26 ZSCAN26 16489 zinc finger and SCAN domain containing 29 ZSCAN29 16490 zinc finger and SCAN domain containing 30 ZSCAN30 16491 zinc finger and SCAN domain containing 31 ZSCAN31 16492 zinc finger and SCAN domain containing 32 ZSCAN32 16493 zinc finger and SCAN domain containing 4 ZSCAN4 16494 zinc finger and SCAN domain containing 5A ZSCAN5A 16495 zinc finger and SCAN domain containing 5B ZSCAN5B 16496 zinc finger and SCAN domain containing 5C, ZSCAN5CP 16497 pseudogene zinc finger and SCAN domain containing 9 ZSCAN9 16498 zinc finger with UFM1-specific peptidase domain ZUFSP 16499 zinc finger, X-linked, duplicated A ZXDA 16500 zinc finger, X-linked, duplicated B ZXDB 16501 ZXD family zinc finger C ZXDC 16502 zinc finger ZZ-type containing 3 ZZZ3 16503

(680) In some embodiments, a T-cell of the disclosure is modified to silence or reduce expression of one or more gene(s) encoding a cell death or cell apoptosis receptor to produce an armored T-cell of the disclosure. Interaction of a death receptor and its endogenous ligand results in the initiation of apoptosis. Disruption of an expression, an activity, or an interaction of a cell death and/or cell apoptosis receptor and/or ligand render an armored T-cell of the disclosure less receptive to death signals, consequently, making the armored T cell of the disclosure more efficacious in a tumor environment. An exemplary cell death receptor which may be modified in an armored T cell of the disclosure is Fas (CD95). Exemplary cell death and/or cell apoptosis receptors and ligands of the disclosure include, but are not limited to, the exemplary receptors and ligands provided in Table 5.

(681) TABLE-US-00137 TABLE 5 Exemplary Cell Death and/or Cell Apoptosis Receptors and Ligands. Full Name Abbreviation SEQ ID NO: Cluster of Differentiation 120 CD120a 16504-16505 Death receptor 3 DR3 16506 Death receptor 6 DR6 16507 first apoptosis signal (Fas) receptor Fas (CD95/AP0-1) 16508-16509 Fas Ligand FasL 16510 cellular tumor antigen p53 p53 16511 Tumor necrosis factor receptor 1 TNF-R1 16512 Tumor necrosis factor receptor 2 TNF-R2 16513 Tumor necrosis factor-related apoptosisinducing TRAIL-R1 (DR4) 16514 ligand receptor 1 Tumor necrosis factor-related apoptosis-inducing TRAIL-R2 (DR5) 16515 ligand receptor 2 Fas-associated protein with death domain FADD 16516 Tumor necrosis factor receptor type 1-associated TRADD 16517 DEATH domain protein Bc1-2-associated X protein Bax 16518 Bc1-2 homologous killer BAK 16519 14-3-3 protein 14-3-3 16520 B-cell lymphoma 2 Bcl-2 16521 Cytochrome C Cyt C 16522 Second mitochondria-derived activator of caspase Smac/Diablo 16523 High temperature requirement protein A2 HTRA2/Omi 16524 Apoptosis inducing factor AIF 16525 Endonuclease G EXOG 16526 Caspase 9 Cas9 16527 Caspase 2 Cas2 16528 Caspase 8 Cas8 16529 Caspase 10 Cas10 16530 Caspase 3 Cas3 16531 Caspase 6 Cas6 16532 Caspase 7 Cas7 16533 Tumor Necrosis Factor alpha TNF-alpha 16534 TNF-related weak inducer of apoptosis TWEAK 16535 TNFrelated weak inducer of apoptosis receptor TWEAK-R 16536 Tumor necrosis factor-related apoptosisinducing TRAIL 16537 ligand TNF ligand-related molecule 1 TL1A 16538 Receptor-interacting serine/threonine-protein kinase 1 RIP1 16539 Cellular inhibitor of apoptosis 1 cIAP-1 16540 TNF receptor-associated factor 2 TRAF-2 16541

(682) In some embodiments, a T-cell of the disclosure is modified to silence or reduce expression of one or more gene(s) encoding a metabolic sensing protein to produce an armored T-cell of the disclosure. Disruption to the metabolic sensing of the immunosuppressive tumor microenvironment (characterized by

low levels of oxygen pH glucose and other molecules) by an armored T-cell of the disclosure leads to extended retention of T-cell function and, consequently, more tumor cells killed per armored cell. For example, HIF1a and VHL play a role in T-cell function while in a hypoxic environment. An armored T-cell of the disclosure may have silenced or reduced expression of one or more genes encoding HIF1a or VHL. Genes and proteins involved in metabolic sensing include, but are not limited to the exemplary genes and proteins provided in Table 6.

(683) TABLE-US-00138 TABLE 6 Exemplary Metabolic Sensing Genes (and encoded Proteins). Full Name Metabolite Abbreviation SEQ ID NO: hypoxia-inducible factor lα Low oxygen HIF-1α 16542 von Hippel–Lindau tumor suppressor Low oxygen VHL 16543 Prolyl-hydroxylase domain proteins High oxygen PHD proteins Glucose transporter 1 glucose GLUT1 16544 Linker of Activated T cells Amino acid (leucine) LAT 16545 CD98 glycoprotein Amino acid (leucine) CD98 16546 Alanine, serine, cysteinepreferring Cationic Amino acid ASCT2/Slc1a5 16547 transporter 2 (glutamine) Solute carrier family 7 member 1 Cationic Amino acids Slc7a1 16548 Solute carrier family 7 member 2 Cationic Amino acids Slc7a2 16549 Solute carrier family 7 member 3 Cationic Amino acids Slc7a3 16550 Solute carrier family 7 member 4 Cationic Amino acids Slc7a4 16551 Solute carrier family 7 member 5 Glycoprotein Slc7a5 16552 associated Amino acids Solute carrier family 7 member 6 Glycoprotein Slc7a6 16553 associated Amino acids Solute carrier family 7 member 7 Glycoprotein Slc7a7 16554 associated Amino acids Solute carrier family 7 member 8 Glycoprotein Slc7a8 16555 associated Amino acids Solute carrier family 7 member 9 Glycoprotein S1c7a9 16556 associated Amino acids Solute carrier family 7 member 10 Glycoprotein Slc7a10 16557 associated Amino acids Solute carrier family 7 member 11 Glycoprotein Slc7a11 16558 associated Amino acids Solute carrier family 7 member 13 Glycoprotein Slc7a13 16559 associated Amino acids Solute carrier family 7 member 14 Cationic Amino acids Slc7a14 16560 Solute carrier family 3 member 2 Amino acid Slc3a2 16561 Calcium transport protein 2 Cationic Amino acid CAT2 16562 (arginine) Calcium transport protein 3 Cationic Amino acid CAT3 16563 (arginine) Calcium transport protein 4 Cationic Amino acid CAT4 16564 (arginine) Bromodomain adjacent to zinc finger Amino acid (arginine) BAZ1B 16565 domain protein 1B PC4 and SFRS1-interacting protein Amino acid (arginine) PSIP1 16566 Translin Amino acid (arginine) TSN 16567 G-protein-coupled receptors Fatty Acid and GPCRs Cholesterol T-cell Receptor, subunit alpha Fatty Acid and TCR alpha 16568 Cholesterol T-cell Receptor, subunit beta Fatty Acid and TCR beta 16569 Cholesterol T-cell Receptor, subunit zeta Fatty Acid and TCR zeta 16570 Cholesterol T-cell Receptor, subunit CD3 epsilon Fatty Acid and TCR CD3 epsilon 16571 Cholesterol T-cell Receptor, subunit CD3 gamma Fatty Acid and TCR CD3 gamma 16572 Cholesterol T-cell Receptor, subunit CD3 delta Fatty Acid and TCR CD3 delta 16573 Cholesterol peroxisome proliferator-activated Fatty Acid and PPARs receptors Cholesterol AMP-activated protein kinase Energy homeostasis AMPK 16574-16575 (intracellular AMP to ATP ratio) P2X purinoceptor 7 Redox homeostasis P2X7 16576

(684) In some embodiments a T-cell of the disclosure is modified to silence or reduce therapy, including a monoclonal antibody, to produce an armored T-cell of the disclosure. Thus an armored T-cell of the disclosure can function and may demonstrate superior function or efficacy whilst in the presence of a cancer therapy (e.g. a chemotherapy, a monoclonal antibody therapy, or another anti-tumor treatment). Proteins involved in conferring sensitivity to a cancer therapy include, but are not limited to, the exemplary proteins provided in Table 7.

(685) TABLE-US-00139 TABLE 7 Exemplary Proteins that Confer Sensitivity to a Cancer Therapeutic. Full Name Abbreviation SEQ ID NO: Copper-transporting ATPase 2 ATP7B 16577 Breakpoint cluster region protein BCR 16578 Abelson tyrosine-protein kinase 1 ABL 16579 Breast cancer resistance protein BCRP 16580 Breast cancer type 1 susceptibility protein BRCA1 16581 Breast cancer type 2 susceptibility protein BRCA2 16582 CAMPATH-1 antigen CD52 16583 Cytochrome P450 2D6 CYP2D6 16584 Deoxycytidine kinase dCK 16585 Dihydrofolate reductase DHFR 16586 Dihydropyrimidine dehydrogenase [NADP (+)] DPYD 16587 Epidermal growth factor receptor EGFR 16588 DNA excision repair protein ERCC-1 ERCC1 16589 Estrogen Receptor ESR 16590 Low affinity immunoglobulin gamma Fc region FCGR3A 16591 receptor III-A Receptor tyrosine-protein kinase erbB-2 HER2 or ERBB2 16592 Insulin-like growth factor 1 receptor IGF1R 16593 GTPase KRas KRAS 16594 Multidrug resistance protein 1 MDR1 or ABCB1 16595 Methylated-DNA--protein-cysteine methyltransferase MGMT 16596 Multidrug resistance-associated protein 1 MRP1 or ABCC1 16597 Progesterone Receptor

PGR 16598 Regulator of G-protein signaling 10 RGS10 16599 Suppressor of cytokine signaling 3 SOCS-3 16600 Thymidylate synthase TYMS 16601 UDP-glucuronosyltransferase 1-1 UGT1A1 16602 (686) In some embodiments, a T-cell of the disclosure is modified to silence or reduce expression of one or more gene(s) encoding a growth advantage factor to produce an armored T-cell. Silencing or reducing expression of an oncogene can confer a growth advantage for an armored T-cell of the disclosure. For example, silencing or reducing expression (e.g. disrupting expression) of a TET2 gene during a CAR-T manufacturing process results in the generation of an armored CAR-T with a significant capacity for expansion and subsequent eradication of a tumor when compared to anon-armored CAR-T lacking this capacity for expansion. This strategy may be coupled to a safety switch (e.g. an iC9 safety switch of the disclosure), which allows for the targeted disruption of an armored CAR-T-cell in the event of an adverse reaction from a subject or uncontrolled growth of the armored CAR-T. Exemplary growth advantage factors include, but are not limited to, the factors provided in Table 8.

(687) TABLE-US-00140 TABLE 8 Exemplary Growth Advantage Factors. Full Name Abbreviation SEQ ID NO: Ten Eleven Translocation 2 TET2 16603 DNA (cytosine-5)-methyltransferase 3A DNMT3A 16604 Transforming protein RhoA RHOA 16605 Proto-oncogene vav VAV1 16606 Rhombotin-2 LMO2 16607 T-cell acute lymphocytic leukemia protein 1 TAL1 16608 Suppressor of cytokine signaling 1 SOCS1 16609 herpes virus entry mediator HVEM 16610 T cell death-associated gene 8 TDAG8 16611 BCL6 corepressor BCOR 16612 B and T cell attenuator BTLA 16613 SPARC-like protein 1 SPARCL1 16614 Msh homeobox 1-like protein MSX1 16615

Armored T-Cells "Null or Switch Receptor" Strategy

- (688) In some embodiments, a T-cell of the disclosure is modified to express a modified/chimeric checkpoint receptor to produce an armored T-cell of the disclosure.
- (689) In some embodiments, the modified/chimeric checkpoint receptor comprises a null receptor, decoy receptor or dominant negative receptor. A null receptor, decoy receptor or dominant negative receptor of the disclosure may be modified/chimeric receptor/protein. A null receptor, decoy receptor or dominant negative receptor of the disclosure may be truncated for expression of the intracellular signaling domain. Alternatively, or in addition, a null receptor, decoy receptor or dominant negative receptor of the disclosure may be mutated within an intracellular signaling domain at one or more amino acid positions that are determinative or required for effective signaling. Truncation or mutation of null receptor, decoy receptor or dominant negative receptor of the disclosure may result in loss of the receptor's capacity to convey or transduce a checkpoint signal to the cell or within the cell.
- (690) For example, a dilution or a blockage of an immunosuppressive checkpoint signal from a PD-L1 receptor expressed on the surface of a tumor cell may be achieved by expressing a modified/chimeric PD-1 null receptor on the surface of an armored T-cell of the disclosure, which effectively competes with the endogenous (non-modified) PD-1 receptors also expressed on the surface of the armored T-cell to reduce or inhibit the transduction of the immunosuppressive checkpoint signal through endogenous PD-1 receptors of the armored T cell. In this exemplary embodiment, competition between the two different receptors for binding to PD-L1 expressed on the tumor cell reduces or diminishes a level of effective checkpoint signaling, thereby enhancing a therapeutic potential of the armored T-cell expressing the PD-1 null receptor.
- (691) In some embodiments, the modified/chimeric checkpoint receptor comprises a null receptor, decoy receptor or dominant negative receptor that is a transmembrane receptor.
- (692) In some embodiments, the modified/chimeric checkpoint receptor comprises a null receptor, decoy receptor or dominant negative receptor that is a membrane-associated or membrane-linked receptor/protein.
- (693) In some embodiments, the modified/chimeric checkpoint receptor comprises a null receptor, decoy receptor or dominant negative receptor that is an intracellular receptor/protein.
- (694) In some embodiments, the modified/chimeric checkpoint receptor comprises a null receptor, decoy receptor or dominant negative receptor that is an intracellular receptor/protein. Exemplary null, decoy, or dominant negative intracellular receptors/proteins of the disclosure include, but are not limited to, signaling components downstream of an inhibitory checkpoint signal (as provided, for example, in Tables 2 and 3), a transcription factor (as provided, for example, in Table 4), a cytokine or a cytokine receptor, a chemokine or a chemokine receptor, a cell death or apoptosis receptor/ligand (as provided, for example, in

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Table 5), a metabolic sensing molecule (as provided, for example, in Table 6), a protein conferring
sensitivity to a cancer therapy (as provided, for example, in Table 7), and an oncogene or a tumor
suppressor gene (as provided, for example, in Table 8). Exemplary cytokines, cytokine receptors,
chemokines and chemokine receptors of the disclosure include, but are not limited to, the cytokines and
cytokine receptors as well as chemokines and chemokine receptors provided in Table 9.
(695) TABLE-US-00141 TABLE 9 Exemplary Cytokines, Cytokine receptors, Chemokines and
Chemokine Receptors. Full Name Abbreviation SEQ ID NO: 4-1BB Ligand 4-1BBL 16616 Tumor
necrosis factor receptor Apo3 or TNFRSF25 16617 superfamily member 25 Tumor necrosis factor
receptor APRIL or TNFRSF13 16618 superfamily member 13 Bc12-associated agonist of cell death Bc1-
xL or BAD 16619 Tumor necrosis factor receptor BCMA or TNFRS17 16620 superfamily member 17 C-
C motif chemokine 1 CCL1 16621 C-C motif chemokine 11 CCL11 16622 C-C motif chemokine 13
CCL13 16623 C-C motif chemokine 14 CCL14 16624 C-C motif chemokine 15 CCL15 16625 C-C motif
chemokine 16 CCL16 16626 C-C motif chemokine 17 CCL17 16627 C-C motif chemokine 18 CCL18
16628 C-C motif chemokine 19 CCL19 16629 C-C motif chemokine 2 CCL2 16630 C-C motif chemokine
20 CCL20 16631 C-C motif chemokine 21 CCL21 16632 C-C motif chemokine 22 CCL22 16633 C-C
motif chemokine 23 CCL23 16634 C-C motif chemokine 24 CCL24 16635 C-C motif chemokine 25
CCL25 16636 C-C motif chemokine 26 CCL26 16637 C-C motif chemokine 27 CCL27 16638 C-C motif
chemokine 28 CCL28 16639 C-C motif chemokine 3 CCL3 16640 C-C motif chemokine 4 CCL4 16641
C-C motif chemokine 5 CCL5 16642 C-C motif chemokine 7 CCL7 16643 C-C motif chemokine 8 CCL8
16644 C-C chemokine receptor type 1 CCR1 16645 C-C chemokine receptor type 10 CCR10 16646 C-C
chemokine receptor type 11 CCR11 16647 C-C chemokine receptor type 2 CCR2 16648 C-C chemokine
receptor type 3 CCR3 16649 C-C chemokine receptor type 4 CCR4 16650 C-C chemokine receptor type 5
CCR5 16651 C-C chemokine receptor type 6 CCR6 16652 C-C chemokine receptor type 7 CCR7 16653
C-C chemokine receptor type 8 CCR8 16654 C-C chemokine receptor type 9 CCR9 16655 Granulocyte
colony-stimulating factor CD114 or CSF3R 16656 receptor Macrophage colony-stimulating factor 1
CD115 or CSF1R 16657 receptor Granulocyte-macrophage colony- CD116 or CSF2RA 16658 stimulating
factor receptor subunit alpha Mast/stem cell growth factor receptor CD117 or KIT 16659 Kit Leukemia
inhibitory factor receptor CD118 or LIFR 16660 Tumor necrosis factor receptor CD120a or TNFRSF1A
16661 superfamily member 1A Tumor necrosis factor receptor CD120b or TNFRSF1B 16662 superfamily
member 1B Interleukin-1 receptor type 1 CD121a or IL1R1 16663 Interleukin-2 receptor subunit beta
CD122 or IL2RB 16664 Interleukin-3 receptor subunit alpha CD123 or IL3RA 16665 Interleukin-4
receptor subunit alpha CD124 or IL4R 16666 Interleukin-6 receptor subunit alpha CD126 or IL6R 16667
Interleukin-7 receptor subunit alpha CD127 or IL7R 16668 Interleukin-6 receptor subunit beta CD130 or
IL6ST 16669 Cytokine receptor common subunit CD132 or IL2RG 16670 gamma Tumor necrosis factor
ligand CD153 or TNFSF8 16671 superfamily member 8 CD40 ligand CD154 or CD4OL 16672 Tumor
necrosis factor ligand CD178 or FASLG 16673 superfamily member 6 Interleukin-12 receptor subunit
beta-1 CD212 or IL12RB1 16674 Interleukin-13 receptor subunit alpha-1 CD213a1 or IL13RA1 16675
Interleukin-13 receptor subunit alpha-2 CD213a2 or IL13RA2 16676 Interleukin-2 receptor subunit alpha
CD25 or IL2RA 16677 CD27 antigen CD27 16678 Tumor necrosis factor receptor CD30 or TNTRSF
16679 superfamily member 8 T-cell surface glycoprotein CD4 CD4 16680 Tumor necrosis factor receptor
CD40 or TNFRSF5 16681 superfamily member 5 CD70 antigen CD70 16682 Tumor necrosis factor
receptor CD95 or FAS or FNFRSF6 16683 superfamily member 6 Granulocyte-macrophage colony-
CDw116 or CSF2RA 16684 stimulating factor receptor subunit alpha Interferon gamma receptor 1
CDw119 or IFNGR1 16685 Interleukin-1 receptor type 2 CDw12lb or IL1R2 16686 Interleukin-5 receptor
subunit alpha CDw125 or IL5RA 16687 Cytokine receptor common subunit beta CDw131 or CSF2RB
16688 Tumor necrosis factor receptor CDw137 or TNFRSF9 16689 superfamily member 9 Interleukin-10
receptor CDw210 or IL1OR 16690 Interleukin-17 receptor A CDw217 or IL17RA 16691 C-X3-C motif
chemokine 1 CX3CL1 16692 C-X3-C chemokine receptor 1 CX3CR1 16693 C-X-C motif chemokine 1
CXCL1 16694 C-X-C motif chemokine 10 CXCL10 16695 C-X-C motif chemokine 11 CXCL11 16696
C-X-C motif chemokine 12 CXCL12 16697 C-X-C motif chemokine 13 CXCL13 16698 C-X-C motif
chemokine 14 CXCL14 16699 C-X-C motif chemokine 16 CXCL16 16700 C-X-C motif chemokine 2
CXCL2 16701 C-X-C motif chemokine 3 CXCL3 16702 C-X-C motif chemokine 4 CXCL4 16703 C-X-C
motif chemokine 5 CXCL5 16704 C-X-C motif chemokine 6 CXCL6 16705 C-X-C motif chemokine 7
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CXCL7 16706 C-X-C motif chemokine 8 CXCL8 16707 C-X-C motif chemokine 9 CXCL9 16708 C-X-C
chemokine receptor type 1 CXCR1 16709 C-X-C chemokine receptor type 2 CXCR2 16710 C-X-C
chemokine receptor type 3 C.XCR3 16711 C-X-C chemokine receptor type 4 CXCR4 16712 C-X-C
chemokine receptor type 5 CXCR5 16713 C-X-C chemokine receptor type 6 CXCR6 16714 C-X-C
chemokine receptor type 7 CXCR7 16715 Atypical chemokine receptor 1 DARC or ACKR1 16716
Erythropoietin Epo 16717 Erythropoietin receptor EpoR 16718 Receptor-type tyrosine-protein kinase Flt-
3 16719 FLT3 FLT3 Ligand Flt-3L 16720 Granulocyte colony-stimulating factor G-CSF or GSF3R 16721
receptor Tumor necrosis factor receptor GITR or TNERSFI8 16722 superfamily member 18 GITR Ligand
GITRL 16723 Cytokine receptor common subunit beta GM-CSF or CSF2RB 16724 Interleukin-6 receptor
subunit beta gp130 or IL6ST 16725 Tumor necrosis factor receptor HVEM or TNFRSF14 16726
superfamily member 14 Interferon gamma IFNy 16727 Interferon gamma receptor 2 IFNGR2 16728
Interferon-alpha IFN-α 16729 Interferon-beta IFN-β 16730 Interleukin-1 alpha IL1 16731 Interleukin-10
IL10 16732 Interleukin-10 receptor IL10R 16733 Interleukin-11 IL-11 16734 Interleukin-11 receptor
alpha IL-11Ra 16735 Interleukin-12 IL12 16736 Interleukin-13 IL13 16737 Interleukin-13 receptor IL13R
16738 Interleukin-14 IL-14 16739 Interleukin-15 IL15 16740 Interleukin-15 receptor alpha IL-15Ra
16741 Interleukin-16 IL-16 16742 Interleukin-17 IL17 16743 Interleukin-17 receptor IL17R 16744
Interleukin-18 IL18 16745 Interleukin-1 receptor alpha IL-1RA 16746 Interleukin-1 alpha IL-lα 16747
Interleukin-l beta IL-1β 16748 interleukin-2 IL2 16749 interleukin-20 IL-20 16750 Interleukin-20 receptor
alpha IL-20Rα 16751 Interleukin-20 receptor beta IL-20Rβ 16752 Interleukin-21 IL21 16753 Interleukin-
3 IL-3 16754 interleukin-35 IL35 16755 Interleukin-4 IL4 16756 Interleukin-4 receptor IL4R 16757
Interleukin-5 IL5 16758 Interleukin-5 receptor IL5R 16759 Interleukin-6 IL6 16760 Interleukin-6 receptor
IL6R 16761 Interleukin-7 IL7 16762 Interleukin-9 receptor IL-9R 16763 Leukemia inhibitory factor LIF
16764 Leukemia inhibitory factor receptor LIFR 16765 tumor necrosis factor superfamay LIGHT or
TNFSF14 16766 member 14 Tumor necrosis factor receptor LTBR or INFRSF3 16767 superfamily
member 3 Lymphotoxin-beta LT-β 16768 Macrophage colony-stimulating factor 1 M-CSF 16769 Tumor
necrosis factor receptor OPG or TNFRSF11B 16770 superfamily member 11B Oncostatin-M OSM 16771
Oncostatin-M receptor OSMR 16772 Tumor necrosis factor receptor OX40 or TNFRSF4 16773
superfamily member 4 Tumor necrosis factor ligand OX40L or TNFSF4 16774 superfamily member 4
Tumor necrosis factor receptor RANK or TNFRSF11A 16775 superfamily member 11A Kit Ligana SCF
or KITLG 16776 Tumor necrosis factor receptor TACI or TNFRSF13B 16777 superfamay member 13B
Tumor necrosis factor ligand TALL-I or TNFSF13B 16778 superfamily member -13B TGF-beta receptor
type-1 TGF-βR1 16779 TGF-beta receptor type-2 TGF-βR2 16780 TGF-beta receptor type-3 TGF-βR3
16781 Transforming growth factor beta-1 TGF-β1 16782 Transforming growth factor beta-2 TGF-β2
16783 Transforming growth factor beta-3 TGF-β3 16784 Tumor necrosis factor alpha TNF or TNF-α
16785 Tumor necrosis factor beta TNF-β 16786 Thyroid peroxidase Tpo 16787 Thyroid peroxidase
receptor TpoR 16788 Tumor necrosis factor ligand TRAIL or TNFSF10 16789 superfamay member 10
Tumor necrosis factor receptor TRAILR1 or TNFRSF10A 16790 superfamay member 10A Tumor
necrosis factor receptor TRAILR2 or TNFRSF10B 16791 superfamily member 10B Tumor necrosis factor
ligand TRANCE or TNFSF11 16792 superfamily member 11 Tumor necrosis factor ligand TWEAK or
TNFSF11 16793 superfamay member 12 Lymphotactin XCL1 16794 Cytokine SCM-1 beta XCL2 16795
(696) In some embodiments, the modified/chimeric checkpoint receptor comprises a switch receptor.
Exemplary switch receptors may comprise a modified chimeric receptor/protein of the disclosure wherein
a native or wild type intracellular signaling domain is switched or replaced with a different intracellular
signaling domain that is either non-native to the protein and/or not a wild-type domain. For example,
replacement of an inhibitory signaling domain with a stimulatory signaling domain would switch an
immunosuppressive signal into an immunostimulatory signal. Alternatively, replacement of an inhibitory
signaling domain with a different inhibitory domain can reduce or enhance the level of inhibitory
signaling. Expression or overexpression, of a switch receptor can result in the dilution and/or blockage of a
cognate checkpoint signal via competition with an endogenous wildtype checkpoint receptor (not a switch
receptor) for binding to the cognate checkpoint receptor expressed within the immunosuppressive tumor
microenvironment. Armored T cells of the disclosure may comprise a sequence encoding switch receptors
of the disclosure, leading to the expression of one or more switch receptors of the disclosure, and
consequently, altering an activity of an armored T-cell of the disclosure. Armored T cells of the disclosure
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may express a switch receptor of the disclosure that targets an intracellularly expressed protein downstream of a checkpoint receptor, a transcription factor, a cytokine receptor, a death receptor, a metabolic sensing molecule, a cancer therapy, an oncogene, and/or a tumor suppressor protein or gene of the disclosure.

(697) Exemplary switch receptors of the disclosure may comprise or may be derived from a protein including, but are not limited to, the signaling components downstream of an inhibitory checkpoint signal (as provided, for example, in Tables 2 and 3), a transcription factor (as provided, for example, in Table 4), a cytokine or a cytokine receptor, a chemokine or a chemokine receptor, a cell death or apoptosis receptor/ligand (as provided, for example, in Table 5), a metabolic sensing molecule (as provided, for example, in Table 6), a protein conferring sensitivity to a cancer therapy (as provided, for example, in Table 7), and an oncogene or a tumor suppressor gene (as provided, for example, in Table 8). Exemplary cytokines, cytokine receptors, chemokines and chemokine receptors of the disclosure include, but are not limited to, the cytokines and cytokine receptors as well as chemokines and chemokine receptors provided in Table 9.

(698) Armored T-Cells "Synthetic Gene Expression" Strategy

(699) In some embodiments, a T-cell of the disclosure is modified to express chimeric ligand receptor (CLR) or a chimeric antigen receptor (CAR) that mediates conditional gene expression to produce an armored T-cell of the disclosure. The combination of the CLR/CAR and the condition gene expression system in the nucleus of the armored T cell constitutes a synthetic gene expression system that is conditionally activated upon binding of cognate ligand(s) with CLR or cognate antigen(s) with CAR. This system may help to 'armor' or enhance therapeutic potential of modified T cells by reducing or limiting synthetic gene expression at the site of ligand or antigen binding, at or within the tumor environment for example.

(700) Exogenous Receptors

(701) In some embodiments, the armored T-cell comprises a composition comprising (a) an inducible transgene construct, comprising a sequence encoding an inducible promoter and a sequence encoding a transgene, and (b) a receptor construct, comprising a sequence encoding a constitutive promoter and a sequence encoding an exogenous receptor, such as a CLR or CAR, wherein, upon integration of the construct of (a) and the construct of (b) into a genomic sequence of a cell, the exogenous receptor is expressed, and wherein the exogenous receptor, upon binding a ligand or antigen, transduces an intracellular signal that targets directly or indirectly the inducible promoter regulating expression of the inducible transgene (a) to modify gene expression.

(702) In some embodiments of a synthetic gene expression system of the disclosure, the composition modifies gene expression by decreasing gene expression. In some embodiments, the composition modifies gene expression by transiently modifying gene expression (e.g. for the duration of binding of the ligand to the exogenous receptor). In some embodiments, the composition modifies gene expression acutely (e.g. the ligand reversibly binds to the exogenous receptor). In some embodiments, the composition modifies gene expression chronically (e.g. the ligand irreversibly binds to the exogenous receptor).

(703) In some embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises an endogenous receptor with respect to the genomic sequence of the cell. Exemplary receptors include, but are not limited to, intracellular receptors, cell-surface receptors, transmembrane receptors, ligand-gated ion channels, and G-protein coupled receptors.

(704) In some embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In some embodiments, the non-naturally occurring receptor is a synthetic, modified, recombinant, mutant or chimeric receptor. In some embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a T-cell receptor (TCR). In some embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a scaffold protein. In some embodiments, including those wherein the non-naturally occurring receptor does not comprise a transmembrane domain, the non-naturally occurring receptor interacts with a second transmembrane, membrane-bound and/or an intracellular receptor that, following contact with the non-naturally occurring receptor, transduces an intracellular signal. (705) In some embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In some embodiments, the non-naturally occurring receptor

is a synthetic, modified, recombinant, mutant or chimeric receptor. In some embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a T-cell receptor (TCR). In some embodiments, the non-naturally occurring receptor comprises one or more sequences isolated or derived from a scaffold protein. In some embodiments, the non-naturally occurring receptor comprises a transmembrane domain. In some embodiments, the non-naturally occurring receptor that transduces an intracellular signal. In some embodiments, the non-naturally occurring receptor comprises an intracellular signalling domain. In some embodiments, the non-naturally occurring receptor is a chimeric ligand receptor (CLR). In some embodiments, the CLR is a chimeric antigen receptor (CAR).

- (706) In some embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In some embodiments, the CLR is a chimeric antigen receptor (CAR). In some embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In some embodiments, the ectodomain of (a) further comprises a signal peptide. In some embodiments, the ectodomain of (a) further comprises a hinge between the ligand recognition region and the transmembrane domain.
- (707) In some embodiments of the CLR/CARs of the disclosure, the signal peptide comprises a sequence encoding a human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD3 $\delta$ , CD3 $\epsilon$ , CD4, CD8 $\alpha$ , CD19, CD28, 4-1 BB or GM-CSFR signal peptide. In some embodiments, the signal peptide comprises a sequence encoding a human CD8 $\alpha$  signal peptide. In some embodiments, the signal peptide comprises an amino acid sequence comprising MALPVTALLLHAARP (SEQ ID NO: 17000). In some embodiments, the signal peptide is encoded by a nucleic acid sequence comprising
- atggcactgccagtcaccgccctgctgctctctggctctgctgctgcacgcagctagacca (SEQ ID NO: 17001). (708) In some embodiments of the CLR/CARs of the disclosure, the transmembrane domain comprises a sequence encoding a human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD3 $\epsilon$ , CD4, CD8 $\alpha$ , CD19, CD28, 4-1BB or GM-CSFR transmembrane domain. In some embodiments, the transmembrane domain comprises a sequence encoding a human CD8 $\alpha$  transmembrane domain. In some embodiments, the transmembrane domain comprises an amino acid sequence comprising IYIWAPLAGTCGVLLLSLVITLYC (SEQ ID NO: 17002). In some embodiments, the transmembrane domain is encoded by a nucleic acid sequence comprising (709) TABLE-US-00142 (SEQ ID NO: 17003) atctacatttgggcaccactggccgggacctgtggag tgctgctgagcctggtcatcacactgtactgc.
- (710) In some embodiments of the CLR/CARs of the disclosure, the endodomain comprises a human CD3 $\zeta$  endodomain. In some embodiments, the at least one costimulatory domain comprises a human 4-1BB, CD28, CD3 $\zeta$ , CD40, ICOS, MyD88, OX-40 intracellular segment, or an) combination thereof. In some embodiments, the at least one costimulatory domain comprises a human CD3 $\zeta$  and/or a 4-1 BB costimulatory domain. In some embodiments, the CD3 $\zeta$  costimulatory domain comprises an amino acid sequence comprising
- RVKFSRSADAPAYKQGQNQLYNELNLGRREEYDVLDKRRGRDPEMGGKPRRKNPQ EGLYNELQKDKMAEAYSEIGMKGERRRGKGHDGLYQGLSTATKDTYDALHMQALP PR (SEQ ID NO: 17004). In some embodiments, the CD3ζ costimulatory domain is encoded by a nucleic acid sequence comprising
- (711) TABLE-US-00143 (SEQ ID NO: 17005) cgcgtgaagtttagtcgatcagcagatgccccagcttacaaaca gggacagaaccagctgtataacgagctgaatctgggccgccgag aggaatatgacgtgctggataagcggagaggacgcgaccccgaa atgggaggcaagcccaggcgcaaaaaccctcaggaaggcctgta taacgagctgcagaaggacaaaatggcagaaggcctattctgaga tcggcatgaaggggagggagggaaggcaaagggcacgatggg ctgtaccagggactgagcaccgccacaaaggacacctatgatgc tctgcatatgcaggcactgcctccaagg.
- In some embodiments, the 4-1BB costimulatory domain comprises an amino acid sequence comprising KRGRKKLLYIFKQPFMRPVQTTQEEDGCSCRFPEEEEGGCEL (SEQ ID NO: 17006). In some embodiments, the 4-1BB costimulatory domain is encoded by a nucleic acid sequence comprising aagaggaggaagaaactgctgtatattttcaaacagcccttcatgcgccccgtgcagactacccaggaggaagacgggtgctcc tgtcgattccctgaggaagaggagggggtgtgagctg (SEQ ID NO: 17007). In some embodiments, the 4-1BB costimulatory domain is located between the transmembrane domain and the CD3ζ costimulatory domain.

In some embodiments of the CLR/CARs of the disclosure, the hinge comprises a sequence derived from a human CD8 $\alpha$ , IgG4, and/or CD4 sequence. In some embodiments, the hinge comprises a sequence derived from a human CD8 $\alpha$  sequence. In some embodiments, the hinge comprises an amino acid sequence comprising

(712) TABLE-US-00144 (SEQ NO: 17008) TTTPAPRPPTPAPTIASQPLSLR PEACRPAAGGAVHTRGLDFACD,

In some embodiments, the hinge is encoded by a nucleic acid sequence comprising actaccacaccagcacctagaccaccaactccagctccaaccatcgcgagtcagccctgagtctgagacctgaggcctgaggcc agctgcaggaggaggaggtggacttcgcctgegac (SEQ ID NO: 17028). In some embodiments, the hinge is encoded by a nucleic acid sequence comprising

ACCACAACCCCTGCCCCAGACCTCCCACACCCGCCCCTACCATCGCGAGTCAGCCCCTGAGTCTGAGCCTGAGCCTGAGCCTGAGCCTGCAGAGCCTGCAGGAGCTGCAGAGCCAGGAGCCTGCAGAGCTGCAGAGCCAGGAGCCTGCAGACCTGCCGCCTGC GAC (SEQ ID NO: 17009). In some embodiments, the at least one protein scaffold specifically binds the ligand.

(713) In some embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In some embodiments, the CLR is a chimeric antigen receptor (CAR). In some embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In some embodiments, the at least one protein scaffold comprises an antibody, an antibody fragment, a single domain antibody, a single chain antibody, an antibody mimetic, or a Centyrin (referred to herein as a CARTyrin). In some embodiments, the ligand recognition region comprises one or more of an antibody, an antibody fragment, a single domain antibody, a single chain antibody, an antibody mimetic, and a Centyrin. In some embodiments, the single domain antibody comprises or consists of a VHH or a VH (referred to herein as a VCAR). In some embodiments, the single domain antibody comprises or consists of a VHH or a VH comprising human complementarity determining regions (CDRs). In some embodiments, the VH is a recombinant or chimeric protein. In some embodiments, the VH is a recombinant or chimeric human protein. In some embodiments, the antibody mimetic comprises or consists of an affibody, an affilin, an affimer, an affitin, an alphabody, an anticalin, an avimer, a DARPin, a Fynomer, a Kunitz domain peptide or a monobody. In some embodiments, the Centyrin comprises or consists of a consensus sequence of at least one fibronectin type III (FN3) domain. (714) In some embodiments of the compositions of the disclosure, the exogenous receptor of (b) comprises a non-naturally occurring receptor. In some embodiments, the CLR is a chimeric antigen receptor (CAR) In some embodiments, the chimeric ligand receptor comprises (a) an ectodomain comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory

comprising a ligand recognition region, wherein the ligand recognition region comprises at least scaffold protein; (b) a transmembrane domain, and (c) an endodomain comprising at least one costimulatory domain. In some embodiments, the Centyrin comprises or consists of a consensus sequence of at least one fibronectin type III (FN3) domain. In some embodiments, the at least one fibronectin type III (FN3) domain is derived from a human protein. In some embodiments, the human protein is Tenascin-C. In some embodiments, the consensus sequence comprises

LPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAINLTVPGSERSYDL TGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT (SEQ ID NO: 17010). In some embodiments, the consensus sequence comprises

MLPAPKNLVVSEVTEDSLRLSWTAPDAAFDSFLIQYQESEKVGEAINLTVPGSERSYD LTGLKPGTEYTVSIYGVKGGHRSNPLSAEFTT (SEQ ID NO: 17011). In some embodiments, the consensus sequence is modified at one or more positions within (a) a A-B loop comprising or consisting of the amino acid residues TEDS at positions 13-16 of the consensus sequence; (b) a B-C loop comprising or consisting of the amino acid residues TAPDAAF at positions 22-28 of the consensus sequence; (c) a C-D loop comprising or consisting of the amino acid residues SEKVGE at positions 38-43 of the consensus sequence; (d) a D-E loop comprising or consisting of the amino acid residues GSER at positions 51-54 of the consensus sequence; (e) a E-F loop comprising or consisting of the amino acid residues GLKPG at positions 60-64 of the consensus sequence; (f) a F-G loop comprising or consisting of the amino acid residues KGGHRSN at positions 75-81 of the consensus sequence; or (g) any combination of (a)-(f). In

some embodiments, the Centyrin comprises a consensus sequence of at least 5 fibronectin type III (FN3) domains. In some embodiments, the Centyrin comprises a consensus sequence of at least 10 fibronectin type III (FN3) domains. In some embodiments, the Centyrin comprises a consensus sequence of at least 15 fibronectin type III (FN3) domains. In some embodiments, the scaffold binds an antigen with at least one affinity selected from a K.sub.D of less than or equal to 10.sup.—9 M, less than or equal to 10.sup.—10M, less than or equal to 10.sup.—11 M, less than or equal to 10.sup.—12 M, less than or equal to 10.sup.—13 M, less than or equal to 10.sup.—14 M, and less than or equal to 10.sup.—15 M. In some embodiments, the K.sub.D is determined by surface plasmon resonance.

(715) Inducible Promoters

- (716) In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding an NFkB promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding an interferon (IFN) promoter or a sequence encoding an interleukin-2 promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding a nuclear receptor subfamily 4 group A member 1 (NR4A1; also known as NUR77) promoter or a sequence encoding a NR4A1 promoter. In certain embodiments of the compositions of the disclosure, the sequence encoding the inducible promoter of (a) comprises a sequence encoding a T-cell surface glycoprotein CD5 (CD5) promoter or a sequence encoding a CD5 promoter. In certain embodiments, the interferon (IFN) promoter is an IFNy promoter. In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a cytokine or a chemokine. In certain embodiments, the cytokine or chemokine comprises IL2, IL3, IL4, IL5, IL6, IL10, IL12, IL13, IL17A/F, IL21, IL22, IL23, transforming growth factor beta (TGFβ), colony stimulating factor 2 (GM-CSF), interferon gamma (IFNy), Tumor necrosis factor (TNFα), LTα, perforin, Granzyme C (Gzmc), Granzyme B (Gzmb), C-C motif chemokine ligand 5 (CCL5), C-C motif chemokine ligand 4 (CCL4), C-C motif chemokine ligand 3 (CCL3), X-C motif chemokine ligand 1 (XCL1) and LIF interleukin 6 family cytokine (Lif).
- (717) In certain embodiments of the compositions of the disclosure, including those wherein the sequence encoding the inducible promoter of (a) comprises a sequence encoding a NR4A1 promoter or a sequence encoding a NR4A1 promoter, the NR4A1 promoter is activated by T-cell Receptor (TCR) stimulation in T cells and by B-cell Receptor (BCR) stimulation in B cells, therefore, inducing expression of any sequence under control of the NR4A1 promoter upon activation of a T-cell or B-cell of the disclosure through a TCR or BCR, respectively.
- (718) In certain embodiments of the compositions of the disclosure, including those wherein the sequence encoding the inducible promoter of (a) comprises a sequence encoding a CD5 promoter or a sequence encoding a CD5 promoter, the CD5 promoter is activated by T-cell Receptor (TCR) stimulation in T cells, therefore, inducing expression of any sequence under control of the CD5 promoter upon activation of a T-cell of the disclosure through a TCR.
- (719) In certain embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a gene comprising a surface protein involved in cell differention, activation, exhaustion and function. In certain embodiments, the gene comprises CD69, CD71, CTLA4, PD-1, TIGIT, LAG3, TIM-3, GITR, MHCII, COX-2, FASL and 4-1BB.
- (720) In some embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of a gene involved in CD metabolism and differentiation. In some embodiments of the compositions of the disclosure, the inducible promoter is isolated or derived from the promoter of Nr4a1, Nr4a3, Tnfrsf9 (4-1BB), Sema7a, Zfp3612, Gadd45b. Dusp5, Dusp6 and Neto2.
- (721) Inducible Transgene
- (722) In some embodiments, the inducible transgene construct comprises or drives expression of a signaling component downstream of an inhibitory checkpoint signal (as provided, for example, in Tables 2 and 3), a transcription factor (as provided, for example, in Table 4), a cytokine or a cytokine receptor, a chemokine or a chemokine receptor, a cell death or apoptosis receptor/ligand (as provided, for example, in Table 5), a metabolic sensing molecule (as provided, for example, in Table 6), a protein conferring sensitivity to a cancer therapy (as provided, for example, in Table 7 and/or 1), and an oncogene or a tumor suppressor gene (as provided, for example, in Table 8). Exemplary cytokines, cytokine receptors,

chemokines and chemokine receptors of the disclosure include, but are not limited to, the cytokines and cytokine receptors as well as chemokines and chemokine receptors provided in Table 9.

(723) Cas-Clover

- (724) The disclosure provides a composition comprising a guide RNA and a fusion protein or a sequence encoding the fusion protein wherein the fusion protein comprises a dCas9 and a Clo051 endonuclease or a nuclease domain thereof.
- (725) Small Cas9 (SaCas9)
- (726) The disclosure provides compositions comprising a small, Cas9 (Cas9) operatively-linked to an effector. In certain embodiments, the disclosure provides a fusion protein comprising, consisting essentially of or consisting of a DNA localization component and an effector molecule, wherein the effector comprises a small, Cas9 (Cas9). In certain embodiments, a small Cas9 construct of the disclosure may comprise an effector comprising a type IIS endonuclease.
- (727) Amino acid sequence of *Staphylococcus aureus* Cas9 with an active catalytic site.
- (728) TABLE-US-00145 (SEQ ID NO: 17074) 1 mkrnyilgld igitsvgygi idyetrdvid agyrlfkean vennegrrsk rgarrlkrrr 61 rhrigrykkl lfdynlltdh selsginpye arykglsgkl seeefsaall hlakrrgyhn 121 vneveedtgn elstkegisr nskaleekyv aelglerlkk dgevrgsinr fktsdyvkea 181 kgllkvgkay hgldgsfidt yidlletrrt yvegpgegsp fgwkdikewy emlmghctyf 241 peelrsvkya ynadlynaln dlnnlvitrd enekleyyek fqiienvfkq kkkptlkqia 301 keilvneedi kgvrvtstgk peftnlkvyh dikditarke iienaelldq iakiltivgs 361 sedigeeltn inseltgeei egisnikgyt gthnlslkai nlildelwht ndngiaifnr 421 lklypkkydl saakeipttl vddfilspyv krsfigsiky inaiikkygl pndiiielar 461 eknskdaakm inemakrnra tnerieeiir ttgkenakyl iekiklhdmq egkclyslea 541 ipledllnnp fnyevdhiip rsvsfdnsfn nkvlvkqeen skkgnrtpfq ylsssdskis 601 yetfkkhiln lakgkgrisk tkkeylleer dinrfsvqkd finrnlvdtr yatrglmnll 661 rsyfrvnnld vkvksinggf tsflrrkwkf kkernkgykh haedaliian adfifkewkk 721 ldkakkvmen gmfeekgaes mpeietegey keifitphqi khikdfkdyk yshrvdkkpn 781 relindtlys trkddkgntl ivnnlnglyd kdndklkkli nkspekllmv hhdpqtyqkl 841 klimeqygde knplykyyee tgnyltkysk kdngpvikki kyygnklnah lditddypns 901 rnkvvklslk pyrfdvyldn gvykfvtvkn ldvikkenyy evnskcyeea kklkkisnga 961 efiasfynnd likingelyr vigynndlln rievnmidit yreyienmnd krppriikti 1021 asktqsikky stdilgnlye vkskkhpgii kkg

Inactivated, Small Cas9 (dSaCas9)

- (729) The disclosure provides compositions comprising an inactivated, small, Cas9 (dSaCas9) operativelylinked to an effector. In certain embodiments, the disclosure provides a fusion protein comprising, consisting essentially of or consisting of a DNA localization component and an effector molecule, wherein the effector comprises a small, inactivated Cas9 (dSaCas9). In certain embodiments, a small, inactivated Cas9 (dSaCas9) construct of the disclosure may comprise an effector comprising a type IIS endonuclease. (730) dSaCas9 Sequence: D10A and N580A mutations (bold, capitalized, and underlined) inactivate the catalytic site.
- (731) TABLE-US-00146 (SEQ ID NO: 17075) 1 mkrnviigl**A** igitsvgygi idyetrdvid agyrlfkean vennegrrsk rgarrlkrrr 61 rhrigrykkl lfdynlltdh selsginpye arykglsgkl seeefsaall hlakrrgyhn 121 vneveedtgn elstkegisr nskaleekyv aelglerlkk dgevrgsinr fktsdyvkea 181 kgllkvgkay haldasfidt yidlletrrt yyegpgegsp fgwkdikewy emlmghctyf 241 peelrsvkya ynadlynaln dlnnlvitrd enekleyyek fqiienvfkg kkkptlkqia 301 keilvneedi kgyrvtstqk peftnlkvyh dikditarke iienaelldq iakiltiyqs 361 sediqeeltn lnseltqeei eqisnlkgyt gthnlslkai nlildelwht ndnqiaifnr 421 lklvpkkvdl sqqkeipttl vddfilspvv krsfiqsikv inaiikkygi pndiiielar 481 eknskdagkm inemqkrnrg tnerieeiir ttgkenakyl iekiklhdmg egkclyslea 541 ipledllnnp fnyevdhiip rsvsfdnsfn nkvlvkgee**A** skkgnrtpfq ylsssdskis 601 yetfkkhiln lakgkgrisk tkkeylleer dinrfsvqkd finrnlydtr yatrglmnll 661 rsyfrvnnld vkyksinggf tsflrrkwkf kkernkgykh haedaliian adfifkewkk 721 ldkakkymen gmfeekgaes mpeietegey keifitphqi khikdfkdyk yshrvdkkpn 781 relindtlys trkddkgntl ivnnlnglyd kdndklkkli nkspekllmy hhdpqtyqkl 841 klimeqygde knplykyyee tgnyltkysk kdngpvikki kyygnklnah lditddypns 901 rnkvvklslk pyrfdvyldn gvykfvtvkn ldvikkenyy evnskcyeea kklkkisnga 961 efiasfynnd likingelyr vigvnndlln rievnmidit yreylenmnd krppriikti 1021 asktgsikky stdilgnlye vkskkhpqii kkg

Inactivated Cas9 (dCas9)

(732) The disclosure provides compositions comprising an inactivated Cas9 (dCas9) operatively-linked to

an effector. In certain embodiments, the disclosure provides a fusion protein comprising, consisting essentially of or consisting of a DNA localization component and an effector molecule, wherein the effector comprises an inactivated Cas9 (dCas9). In certain embodiments, an inactivated Cas9 (dCas9) construct of the disclosure may comprise an effector comprising a type IIS endonuclease. (733) In certain embodiments, the dCas9 of the disclosure comprises a dCas9 isolated or derived from *Staphylococcus pyogenes*. In certain embodiments, the dCas9 comprises a dCas9 with substitutions at positions 10 and 840 of the amino acid sequence of the dCas9 which inactivate the catalytic site. In certain embodiments, these substitutions are D10A and H840A. In certain embodiments, the amino acid sequence of the dCas9 comprises the sequence of:

(734) TABLE-US-00147 (SEQ ID NO: 17076) 1 XDKKYSIGLA IGTNSVGWAV ITDEYKVPSK KFKVLGNTDR HSIKKNLIGA LLFDSGETAE 61 ATRLKRTARR YLQEIFSNEM AKVDDSFFHR LEESFLVEED KKHERHPIFG 121 RYTRRKNRIC NIVDEVAYHE KYPTTYHLRK KLVDSTDKAD LRLIYLALAH MIKFRGHFLI EGDLNPDNSD 181 VDKLFIQLVQ TYNQLFEENP INASGVDAKA ILSARLSKSR RLENLIAQLP GEKKNGLFGN 241 LIALSLGLTP NFKSNFDLAE DAKLQLSKDT YDDDLDNLLA QIGDQYADLF LAAKNLSDAI 301 LLSDILRVNT EITKAPLSAS MIKRYDEHHQ DLTLLKALVR QQLPEKYKEI FFDQSKHGYA 361 GYIDGGASQE EFYKFIKPIL EKMDGTEELL VKLNREDLLR KQRTFDNGSI PHQIHLGELH 421 AILRRQEDFY PFLKDNREKI EKILTFRIPY YVGPLARGKS RFAWMTRKSE ETITPWNFEE 481 VVDKGASAQS FIERMTNFDK NLPNEKVLPK HSLLYEYFTV YNELTKVKYV TEGMRKPAFL 541 SGEQKKAIVD LLFKTNRKVT VKQLKEDYFK KIECFDSVEI SGVEDRFNAS LGTYHDLLKI 601 IKDKDFLDNE ENEDILEDIV LTLTLFEDRE MIEERLKTYA HLFDDKVMKQ LKRRRYTGWG 661 RLSRKLINGI RDKQSGKTIL DFLKSDGFAN RNFMQLIHDD SLTFKEDIQK AQVSGQGDSL 721 HEHIAMLAGS PAIKKGILQT VKVVDELVKV MGRHKPENIV IEMARENQTT QKGQKNSRER 781 MKRIEEGIKE LGSQILKEHP VENTQLQNEK LYLYYLQNGR DMYVDQELDI NRLSDYDVD<u>A</u> 841 IVPQSFLKDD SIDNKVLTRS DKNRGKSDNV PSEEVVKKMK NYWRQLLNAK LITQRKFDNL 901 TKAERGGLSE LDKAGFIKRQ LVETRQITKH VAQILDSRMN TKYDENDKLI REVKVITLKS 961 KLVSDFRKDF QFYKVREINN YHHAHDAYLN AVVGTALIKK YPKLESEFVY GDYKVYDVRK 1021 MIAKSEQEIG KATAKYFFYS NIMNFFKTEI TLANGEIRKR PLIETNGETG EIVWDKGRDF 1081 ATVRKVLSMP QVNIVKKTEV QTGGFSKESI LPKRNSDKLI ARKKDWDPKK YGGFDSPTVA 1141 YSVLVVAKVE KGKSKKLKSV KELLGITIME RSSFEKNPID FLEAKGYKEV KKDLIIKLPK 1201 YSLFELENGR KRMLASAGEL QKGNELALPS KYVNFLYLAS HYEKLKGSPE DNEQKQLFVE 1261 QHKHYLDEII EQISEFSKRV ILADANLDKV LSAYNKHRDK PIREQAENII HLFTLTNLGA 1321 PAAFKYFDTT IDRKRYTSTK EVLDATLIHQ SITGLYETRI DLSQLGGD. (735) In certain embodiments, the amino acid sequence of the dCas9 comprises the sequence of: (736) TABLE-US-00148 (SEQ ID NO: 17077 1 MBKKYSIGLA IGTNSVGWAV ITDEYKVPSK KFKVLGNTDR HSIKKNLIGA LLFDSGETAE 61 ATRLRRTARR RYTRRKNRIC YLQEIFSNEM AKVDDSFFKR LEESFLVEED KKHERHPIFG 121 NIVDEVAYHE KYPTIYHLRK KLVDSTDKAD LRLIYLALAH MIKFRGHFLI EGDLNPDNSD 181 VDKLFIQLVQ TYNQLFEENP INASGVDAKA ILSARLSKSR RLENLIAQLP GEKKNGLFGN 241 LIALSLGLTP NFKSNFDLAE DAKLQLSKDT YDDDLDNLLA QIGDQYADLF LAAKNLSDAI 301 LLSDILRVNT EITKAPLSAS MIKRYDEHHQ DLTLLKALVR QQLPEKYKEI FFDQSKNGYA 361 GYIDGGASQE EFYKFIKPIL EKMDGTEELL VKLNREDLLR KQRTFDNGSI PKQIHLGELH 421 AILRRQEDFY PFLKDNREKI EKILTFRIPY YVGPLARGNS RFAWMTRKSE ETITPWNFEE 481 VVDKGASAQS FIERMTMFDK NLPNEKVLPK HSLLYEYFTV YNELTKVKYV TEGMRKPAFL 541 SGEOKKAIVD LLFKTNRKVT VKOLKEDYFK KIECFDSVEI SGVEDRFNAS LGTYHDLLKI 601 IKDKDFLDNE ENEDILEDIV MIEERLKTYA HLFDDKVMKQ LKRRRYTGWG 661 RLSRKLINGI LTLTLFEDRE RDKQSGKTIL DFLKSDGFAN RNFMQLIHDD SLTFKEDIQK AQVSGQGDSL 721

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HEHIANLAGS PAIKKGILQT VKVVDELVKV MGRHKPENIV IEMARENQTT
QKGQKNSRER 781 MKRIEEGIKE LGSQILKEHP VENTQLQNEK LYLYYLQNGR
DMYVDQELDI NRLSDYDVD<u>A</u> 841 IVPQSFLKDD SIDNKVLTRS DKNRGKSDNV
PSEEVVKKMK NYWRQLLNAK LITQRKFDNL 901 TKAERGGLSE LDKAGFIKRQ
LVETRQITKH VAQILDSRMN TKYDENDKLI REVKVITLKS 961 KLVSDFRKDF
QFYKVREINN YHHAHDAYLN AVVGTALIKK YPKLESEFVY GDYKVYDVRK 1021
MIAKSEQEIG KATAKYFFYS NIMNFFKTEI TLANGEIRKR PLIETNGETG
EIVWDKGRDF 1081 ATVRKVLSMP QVNIVKKTEV QTGGFSKESI LPKRNSDKLI
ARKKDWDPKK YGGFDSPTVA 1141 YSVLVVAKVE KGKSKKLKSV KELLGITIME
RSSFEKNPID FLEAKGYKEV KKDLIIKLPK 1201 YSLFELENGR KRMLASAGEL
QKGNELALPS KYVNFLYLAS HYEKLKGSPE DNEQKQLFVE 1281 QHKHYLDEII
EQISEFSKRV ILADANLDKV LSAYNKHRDK PIREQAENII HLFTLTNLGA 1321
PAAFKYFDTT IDRKPYTSTK EVLDATLIHQ SITGLYETRI DLSQLGGD.
Clo051 Endonuclease
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(737) An exemplary Clo051 nuclease domain may comprise, consist essentially of or consist of, the amino acid sequence of:

(738) TABLE-US-00149 (SEQ ID NO: 17078) EGIKSNISLLKDELRGQISHISHEYLSLIDLAFDSKQNRLF

EMKVLELLVNEYGFKGRHLGGSRKPDGIVYSTTLEDNFGII

**VDTKAYSEGYSLPISQADEMERYVRENSNRDEEVNPNKWW** NFSEEVKKYYFVFISGSFKGKFEEQLRRLSMTTGVNGSAVN

WNLLLGAEKIRSGEMTIEELERAMFNNSEFILKY

Cas-Clover Fusion Protein

(739) In certain embodiments, an exemplary dCas9-Clo051 fusion protein (embodiment 1) may comprise, consist essentially of or consist of, the amino acid sequence of (Clo051 sequence underlined, linker bold italics, dCas9 sequence (*Streptococcus pyogenes*) in italics):

(740) TABLE-US-00150 (SEQ ID NO: 17079)

MAPKKKRKVEGIKSNISLLKDELRGQISHISHEYLSLIDLAFDSKQNRL

FEMKVLELLVNEYGFKGRHLGGSRKPDGIVYSTTLEDNFGIIVDTKAYS

<u>EGYSLPISQADEMERYVRENSNRDEEVNPNKWWENFSEEVKKYYFVFIS</u>

GSFKGKFEEQLRRLSMTTGVNGSAVNVVNLLLGAEKIRSGEMTIEELER AMFNNSEFILKY

custom character DKKYSIGLAIGTNSVGWAVITDEYKVPSKKF KVLGNTDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYL QEIFSNEMAKVDDSFFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEK YPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSD *VDKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRLENLIAQL* PGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLDNL LAOIGDOYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDE HHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFI KPILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIHIGELHAILRR **QEDFYPFLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMTRKSEETIT** PWNFEEVVDKGASAGSFIERMTNFDKNLPNEKVLPKHSLLYEYFTVYNE LTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKI **ECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVL** TLTLFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGI RDKQSGKTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDS LHEHIANLAGSPAIKKGILOTVKVVDELVKVMGRHKPENIVIEMARENO TTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENTQLQNEKLYLYYLQ NGRDMYVDQELDINRLSDYDVDAIVPQSFLKDDSIDNKVLTRSDKNRGK SDNVPSEEVVKKMKNYWROLLNAKLITORKFDNLTKAERGGLSELDKAG FIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVS DFRKDFOFYKVREINNYHHAHDAYLNAVVGTALIKKYPKLESEFVYGDY

KVYDVRKMIAKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPL

IETNGETGEIVWDXGRDFATVRKVLSMPQVNIVKKTEVQTGGFSKESIL PKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVVAKVEKGKSKKLKSV KELLGITIMERSSFEKNPIDFLEAKGYKEVKKDLIIKLPKYSLFELENG RKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLKGSPEDNEQKQLF VEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAE NIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLY ETRIDLSQLGGDGSPKKKRKVSS.

(741) In certain embodiments, an exemplary dCas9-Clo051 fusion protein (embodiment 1) may comprise, consist essentially of or consist of, the nucleic acid sequence of (dCas9 sequence derived from *Streptococcus pyogenes*):

(742) TABLE-US-00151 (SEQ ID NO: 17080) 1 atggcaccaa agaagaaaag aaaagtggag ggcatcaagt caaacatcag cctgctgaaa 61 gacgaactgc ggggacagat tagtcacatc agtcacgagt acctgtcact gattgatctg 121 gccttcgaca gcaagcagaa tagactgttt gagatgaaag tgctggaact gctggtcaac 181 gagtatggct tcaagggcag acatctgggc gggtctagga aacctgacgg catcgtgtac 241 agtaccacac tggaagacaa cttcggaatc attgtcgata ccaaggctta ttccgagggc 301 tactgtgtgc caattagtca ggcagatgag acgtgcgcga aaactcaaat 361 agggacgagg aagtcaaccc caataagtgg tgggagaatt atggaaaggt agtgaagaaa 421 tactacttcg tctttatctc aggcagcttc aaagggaagt ttgaggaaca tcagcgagga gctgcggaga 481 ctgtccatga ctaccggggt gaacggatct gctgtcaacg tggtcaatct gctgctgggc 541 gcagaaaaga ggagatgaca attgaggaac tggaacgcgc catgttcaac 601 aattctgagt ttatcctgaa gtatggaggc tcaggtccgg ataagaaata ctccatcgga 661 ctggccattg gcaccaattc cgtgggctgg gctgtcatca gggggaagcg cagacgagta caaggtgcca 721 agcaagaagt tcaaggtcct ggggaacacc gatcgccaca gtatcaagaa aaatctgatt 781 ggagccctgc tgttcgactc aggcgagact gctgaagcaa cccgactgaa gcggactgct 841 aggcgccgat aaaaaatcgg atctgctacc tgcaggaaat tttcagcaac 901 gagatggcca aggtggacga tagtttcttt atacccggag aggaatcatt cctggtggag 961 gaagataaga aacacgagcg gcatcccatc tttggcaaca caccgcctgg agtcgcttat 1021 cacgagaagt accctactat ctatcatctg aggaagaaac tggtggactc caccgataag ttgtggacga 1081 gcagacctgc gcctgatcta tctggccctg gctcacatga tcaagttccg ggggcatttt 1141 ctgatcgagg gagatctgaa ccctgacaat tctgatgtgg acaagctgtt catccagctg 1201 gtccagacat acaatcagct gtttgaggaa cgtggacgca 1261 aaggccatcc tgagcgccag actgtccaaa tctaggcgcc aacccaatta atgcctcagg gatcgctcag 1321 ctgccaggag agaagaaaaa cggcctgttt gggaatctga ttgcactgtc cctgggcctg tggaaaacct gctgtccaaa 1441 gacacttatg 1381 acacccaact tcaagtctaa ttttgatctg gccgaggacg ctaagctgca ggataacctg ctggctcaga tcggcgatca gtacgcagac 1501 ctgttcctgg ccgctaagaa acgatgacct gccatcctgc tgtcagatat tctgcgcgtg 1561 aacacagaga ttactaaggc cccactgagt gcttcaatga tctgagtgac tgacgagcac 1621 catcaggatc tgaccctgct gaaggctctg gtgaggcagc agctgcccga tcaaaagata gaaatacaag 1681 gaaatcttct ttgatcagag caagaatgga tacgccggct atattgacgg cggggcttcc 1741 caggaggagt tctacaagtt catcaagccc attctggaaa agatggacgg caccgaggaa 1801 ctgctggtga ggaggacctg ctgagaaaac agaggacatt tgataacgga 1861 agcatccctc accagattca agctgaatcg tcctgcgacg gcaggaggac 1921 ttctacccat ttctgaagga ctgcacgcca taaccgcgag tctgggcgaa cttcagaatc 1981 ccctactatg tggggcctct ggcacgggga aatagtagat aaaatcgaaa agatcctgac ttgcctggat gacaagaaag 2041 tcagaggaaa ctatcacccc ctggaacttc gaggaagtgg tcgataaagg cgctagcgca 2101 cagtccttca ttgaaaggat gacaaatttt gacaagaacc tgccaaatga gaaggtgctg 2161 gcgtgctgta cgaatatttc acagtgtata acgagctgac taaagtgaag 2221 tacgtcaccg cccaaacaca caagcccgca ttcctgtccg gagagcagaa gaaagccatc 2281 gtggacctgc tgtttaagac aagggatgcg gtgactgtca aacagctgaa ggaagactat 2341 ttcaagaaaa ttgagtgttt cgattcagtg aaatcggaaa gcgtcgagga caggtttaac 2401 gcctccctgg ggacctacca cgatctgctg aagatcatca gaaatcagcg cttcctggac 2461 aacgaggaaa atgaggacat cctggaggac attgtgctga cactgactct gtttgaggat aggataagga 2521 cgcgaaatga tcgaggaacg actgaagact tatgcccatc tgttcgatga caaagtgatg 2581 aagcagctga ctacaccgga tggggacgcc tgagccgaaa actgatcaat 2641 gggattagag acaagcagag aaagaaggcg atcctggact ttctgaagtc cgatggcttc 2701 gccaacagga acttcatgca gctgattcac gatgactctc cggaaaaact ggaggacatc 2761 cagaaagcac aggtgtctgg ccagggggac agtctgcacg agcatatcgc tgaccttcaa aaacctggcc 2821 ggcagccccg ccatcaagaa agggattctg cagaccgtga aggtggtgga cgaactggtc 2881 gacgacacaa acctgagaac atcgtgattg agatggcccg cgaaaatcag 2941 acaactcaga aaggtcatgg agggccagaa aaacagtcga gaacggatga agagaatcga ggaaggcatc 3001 aaggagctgg ggtcacagat catcctgtgg aaaacactca gctgcagaat 3061 gagaaactgt atctgtacta tctgcagaat cctgaaggag

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ggacgggata tgtacgtgga ccaggagctg 3121 gatattaaca gactgagtga ttatgacgtg
                                                                       gatgccatcg
           cttcctgaag 3181 gatgactcca ttgacaacaa ggtgctgacc aggtccgaca
tccctcagag
                                                                        agaaccgcgg
                                                                                    caaatcagat
3241 aatgtgccaa gcgaggaagt ggtcaagaaa atgaagaact actggaggca gctgctgaat 3301 gccaagctga
           gaaatttgat aacctgacta aggcagaaag aggagggtg 3361 tctgagctgg acaaggccgg
tcacacagcg
                      tggagacaag acagatcact 3421 aagcacgtcg ctcagattct ggatagcaga
cttcatcaag
          cggcagctgg
           agtacgatga aaacgacaag 3461 ctgatcaggg aggtgaaagt cattactctg
                                                                        aaatccaagc
atgaacacaa
tggtgtctga ctttagaaag 3541 gatttccagt tttataaagt cagggagatc aacaactacc accatgctca tgacgcatac
3601 ctgaacgcag tggtcgggac cgccctgatt aagaaatacc ccaagctgga gtccgagttc 3661 gtgtacggag
actataaagt gtacgatgtc cggaagatga tcgccaaatc tgagcaggaa 3721 attggcaagg ccaccgctaa gtatttcttt
tacagtaaca tcatgaattt ctttaagacc 3781 gaaatcacac tggcaaatgg ggagatcaga aaaaggcctc
tgattgagac caacggggag 3841 acaggagaaa tcgtgtggga caagggaagg gattttgcta ccgtgcgcaa
agtectgtee 3901 atgececaag tgaatattgt caagaaaact gaagtgeaga eegggggatt etetaaggag 3961
agtattctgc ctaagcgaaa ctctgataaa ctgatcgccc ggaagaaaga ctgggacccc 4021 aagaagtatg
gcgggttcga ctctccaaca gtggcttaca gtgtcctggt ggtcgcaaag 4081 gtggaaaagg ggaagtccaa
                      agctgctggg aatcactatt 4141 atggaacgca gctccttcga gaagaatcct atcgattttc
gaaactgaag
           tctgtcaaag
tggaagccaa gggctataaa 4201 gaggtgaaga aagacctgat cattaagctg ccaaaatact cactgtttga
                                                                                  gctggaaaac
4261 ggacgaaagc gaatgctggc aagcgccgga gaactgcaga agggcaatga gctggccctg 4321 ccctccaaat
acgtgaactt cctgtatctg gctagccact acgagaaact gaaggggtcc 4381 cctgaggata acgaacagaa
gcagctgttt gtggagcagc acaaacatta tctggacgag 4441 atcattgaac agatttcaga gttcagcaag
agagtgatcc tggctgacgc aaatctggat 4501 aaagtcctga gcgcatacaa caagcaccga gacaaaccaa
tccgggagca ggccgaaaat 4561 atcattcatc tgttcaccct gacaaacctg ggcgcccctg cagccttcaa gtattttgac
4621 accacaatcg atcggaagag atacacttct accaaagagg tgctggatgc taccctgatc 4681 caccagagta
ttaccggcct gtatgagaca cgcatcgacc tgtcacagct gggaggcgat 4741 gggagcccca agaaaaagcg
gaaggtgtct agttaa
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(743) In certain embodiments, the nucleic acid sequence encoding a dCas9-Clo051 fusion protein (embodiment 1) of the disclosure may comprise a DNA. In certain embodiments, the nucleic acid sequence encoding a dCas9-Clo051 fusion protein (embodiment 1) of the disclosure may comprise an RNA.

(744) In certain embodiments, an exemplary dCas9-Clo051 fusion protein (embodiment 2) may comprise, consist essentially of or consist of, the amino acid sequence of (Clo051 sequence underlined, linker bold italics, dCas9 sequence (*Streptococcus pyogenes*) in italics):

(745) TABLE-US-00152 (SEQ ID NO: 17081) 1 MPKKKRKV<u>EG IKSNISLLKD</u> ELRGQISHIS HEYLSLIDLA FDSKQNPLFE MKVLELLVNE 61 YGFKGRHLGG SRKPDGIVYS TTLEDNFGII VDTKAYSEGY SLPISQADEM ERYVRENSNR 121 DEEVNPNKWW ENFSEEVKKY YFVFISGSFK GKFEEQLRRL SMTTGVNGSA VNVVNLLLGA 181 EKIRSGEMTI EELERAMFNN SEFILKY custom character *DKKYSIGL* AIGTNSVGWA VITDEYKVPS 241 KKFKVLGNTD RHSIKKNLIG ALLFDSGETA **EATRLKRTAR** RRYTRRFNRI CYLOEIFSNE 301 MAKVDDSFFH RLEESFLVEE DKKHERHPIF **GNIVDEVAYH** EKYPTIYHLR KKLVDSTDKA 361 DLRLIYLALA HMIKFRGHFL IEGDLNPDNS **DVDKLFIQLV** OTYNOLFEEN PINASGVDAK 421 AILSARLSKS RRLENLIAQL PGEKKNGLFG *NLIALSLGLT* PNFKSNFDLA EDAKLQLSKD 481 TYDDDLDNLL AQIGDQYADL FLAAKNLSDA ILLSDILRVN TEITKAPLSA SMIKRYDEHH 541 QDLTLLKALV RQQLPEKYKE IFFDQSKNGY AGYIDGGASQ EEFYKFIKPI LEKMDGTEEL 601 LVKLNREDLL RKQRTFDNGS IPHQIHLGEL **HAILRRQEDF** YPFLKDNREK IEKILTFRIP 661 YYVGPLARGN SRFAWMTRKS EETITPWNFE EVVDKGASAQ SFIERMTNFD KNLPNEKVLP 721 KHSLLYEYFT VYNELTKVKY VTEGMRKPAF **LSGEQEEAIV** TVKQLKEDYF 781 KKIECFDSVE ISGVEDRFNA SLGTYHDLLK IIKDKDFLDN DLLFKTNRKV EENEDILEDI VLTLTLFEDR 841 EMIEERLKTY AHLFDDKVMK OLKRRRYTGW GRLSRKLING IRDKQSGKTI LDFLKSDGFA 901 NRNFMQLIHD DSLTFKEDIQ KAQVSGQGDS LHEHIANLAG SPAIKKGILQ TVKVVDELVK 961 VMGRHKPENI VIEMARENQT TQKGQKNSRE RMKRIEEGIK ELGSQILKEH PVENTQLQNE 1021 KLYLYYLQNG RDMYVDQELD INRLSDYDVD AIVPQSFLKD DSIDNKVLTR SDKNRGKSDN 1061 VPSEEVVKKM KNYWRQLLNA KLITQRKFDN LTKAERGGLS ELDKAGFIKR QLVETRQITK 1141 HVAQILDSRM NTKYDENDKL IREVKVITLK SKLVSDFRKD FQFYKVREIN

NYHHAHDAYL 1201 NAVVGTALIK KYPKLESEFV YGDYKVYDVR KMIAKSEQEI GKATAKYFFY SNIMNFFKTE 1261 ITLANGEIRK RPLIETNGET GEIVWDKGRD FATVRKVLSM PQVNIVKKTE VQTGGFSKES 1321 ILPKRNSDKL IARKKDWDPK KYGGFDSPTV AYSVLVVAKV EKGKSKKLKS VKELLGITIM 1381 ERSSFEKNPI DFLEAKGYKE VKKDLIIKLP KYSLFELENG RKRMLASAGE LQKGNELALP 1441 SKYVNFLYLA SHYEKLKGSP EDNEQKQLFV EQHKHYLDEI IEQISEFSKR VILADANLDK 1501 VLSAYNKHRD KPIREQAENI IHLFTLTNLG APAAFKYFDT TIDRKRYTST KEVLDATLIH 1561 QSITGLYETR IDLSQLGGDG SPKKKRKV.

(746) In certain embodiments, an exemplary dCas9-Clo051 fusion protein (embodiment 2) may comprise, consist essentially of or consist of, the nucleic acid sequence of (dCas9 sequence derived from *Streptococcus pyogenes*):

(747) TABLE-US-00153 (SEQ ID NO: 17082) 1 atgcctaaga agaagcggaa ggtggaaggc atcaaaagca acatctccct cctgaaagac 61 gaactccggg ggcagattag ccacattagt cacgaatacc tctccctcat cgacctggct 121 ttcgatagca agcagaacag gctctttgag atgaaagtgc tggaactgct cgtcaatgag 181 tacgggttca agggtcgaca cctcggcgga tctaggaaac cagacggcat cgtgtatagt 211 accacactgg aagacaactt tgggatcatt gtggatacca aggcatactc tgagggttat 301 agtctgccca tttcacaggc cgacgagatg gaacggtacg tgcgcgagaa ctcaaataga 361 gatgaggaag tcaaccctaa caagtggtgg gagaacttct ctgaggaagt gaagaaatac 421 tacttcgtct ttatcagcgg gtccttcaag ggtaaatttg aggaacagct caggagactg 481 agcatgacta ccggcgtgaa tggcagcgcc gtcaacgtgg tcaatctgct cctgggcgct 541 gaaaagattc ggagcggaga gatgaccatc gaagagctgg agagggcaat gtttaataat 501 agcgagttta tcctgaaata cggtggcggt ggatccgata aaaagtattc tattggttta 661 gccatcggca ctaattccga tggatgggct gtcataaccg agtaccttca 721 aagaaattta aggtgttggg gaacacagac cgtcattcga ttaaaaagaa tcttatcggt 781 atgaatacaa gccctcctat tcgatagtgg cgaaacggca gaggcgactc gcctgaaacg aaccgctcgg 841 agaaggtata cacgtcgcaa gaaccgaata tgttacttac aagaaatttt tagcaatgag 901 atggccaaag ttgacgattc tttctttcac agtccttcct tgtcgaagag 961 gacaagaaac atgaacggca ccccatcttt ggaaacatag tagatgaggt cgtttggaag ggcatatcat 1021 gaaaagtacc caacgattta tcacctcaga aaaaagctag ttgactcaac tgataaagcg 1081 gacctgaggt taatctactt ggctcttgcc catatgataa agttccgtgg gcactttctc 1141 attgagggtg atctaaatcc gatgtcgaca aactgttcat ccagttagta 1201 caaacctata atcagttgtt tgaagagaac cctataaatg ggacaactcg caagtggcgt ggatgcgaag 1261 gctattctta gcgcccgcct ctctaaatcc cgacggctag aaaacctgat cgcacaatta 1321 cccggagaga agaaaaatgg gttgttcggt aaccttatag cgctctcact aggcctgaca 1381 ccaaatttta agtcgaactt cgacttagct gaagatgcca aattgcagct tagtaaggac 1441 acgtacgatg acgatctcga gcacaaattg gagatcagta tgcggactta 1501 tttttggctg ccaaaaaacct tagcgatgca atcctcctat caatctactg ctgacatact gagagttaat 1561 actgagatta ccaaggcgcc gttatccgct tcaatgatca aaaggtacga tgaacatcac 1621 caagacttga cacttctcaa ggccctagtc cgtcagcaac tgcctgagaa atataaggaa 1681 atattctttg aaacgggtac gcaggttata ttgacggcgg agcgagtcaa 1741 gaggaattct acaagtttat atcagtcgaa ttagagaaga tggatgggac ggaagagttg 1801 cttgtaaaac tcaatcgcga caaacccata agatctactg cgaaagcagc ggactttcga caacggtagc 1861 attccacatc aaatccactt aggcgaattg catgctatac ttagaaggca ggaggatttt 1921 tatccgttcc tcaaagacaa tcgtgaaaag attgagaaaa tcctaacctt tcgcatacct 1981 tactatgtgg gacccctggc ccgagggaac tctcggttcg catggatgac aagaaagtcc 2041 gaagaaacga ttactccatg gaattttgag gaagttgtcg ataaaggtgc gtcagctcaa 2101 tcgttcatcg agaggatgac caactttgac aagaatttac cgaacgaaaa agtattgcct 2161 aagcacagtt tactttacga gtatttcaca gtgtacaatg aactcacgaa agttaagtat 2221 gtcactaagg gcatgcgtaa acccgccttt ctaagcgaag aacagaagaa agcaatagta 2281 gatctgttat tcaagaccaa ccgcaaagtg acagttaagc aattgaaaga ggactacttt 2341 aagaaaattg aatgcttcga ttctgtcgag atctccgggg tagaagatcg atttaatgcg 2401 tcacttggta cgtatcatga cctcctaaag ataattaaag ataaggactt cctggataac 2461 gaagagaatg aagatatctt agaagatata gtgttgactc ttaccctctt 2521 gaaatgattg aggaaagact aaaaacatac gctcacctgt tcgacgataa ggttatgaaa 2581 cagttaaaga ggcgtcgcta tacgggctgg ggacgattgt cgcggaaact tatcaacggg 2641 ataagagaca agcaaagtgg taaaactatt ctcgattttc taaagagcga cggcttcgcc 2701 aataggaact ttatgcagct gatccatgat gactctttaa ccttcaaaga ggatatacaa 2761 aaggcacagg tttccggaca aggggactca ttgcacgaac atattgcgaa tcttgctggt 2821 tcgccagcca tcaaaaaggg catactccag acagtcaaag tagtggatga gctagttaag 2881 gtcatgggac ggaaaacatt gtaatcgaga tggcacgcga aaatcaaacg 2941 actcagaagg ggcaaaaaaa gtcacaaacc cggatgaaga gaatagaaga gggtattaaa 3001 gaactgggca gccagatctt aaaggagcat cagtcgagag cctgtggaaa atacccaatt gcagaacgag 3061 aaactttacc tctattacct acaaaatgga agggacatgt atgttgatca

ggaactggac 3121 ataaaccgtt tatctgatta cgacgtcgat gccattgtac cccaatcctt tttgaaggac 3181 acaataaagc gcttacacgc tcggataaga accgagggaa aagtgacaat 3241 gttccaagcg gattcaatcg ggcggcagct cctaaatgcg 3301 aaactgataa cgcaaagaaa aaagaaaatg aagaactatt aggaagtcgt tggcttgtct 3361 gaacttgaca aggccggatt tattaaacgt cagctcgtgg gttcgataac ttaactaaag ctgagagggg aatacgaaat acgacgagaa cgataagctg aaacccgcca aatcacaaag 3421 catgttgcac agatactaga ttcccgaatg 3481 attcgggaag tcaaagtaat cactttaaag tcaaaattgg tgtcggactt cagaaaggat 3541 tttcaattct atgcgcacga cgcttatctt 3601 aatgccgtcg tagggaccgc ataaagttag ggagataaat aactaccacc tgagtttgtg 3661 tatggtgatt acaaagttta tgacgtccgt aagatgatcg actcattaag aaatacccga agctagaaag cttcttttat tctaacatta tgaatttctt taagacggaa cgaaaagcga acaggagata 3721 ggcaaggcta cagccaaata 3781 atcactctgg caaacggaga gatacgcaaa cgacctttaa ttgaaaccaa tggggagaca 3841 ggtgaaatcg tatgggataa ttcgcgacgg tgagaaaagt tttgtccatg 3901 ccccaagtca acatagtaaa gggccgggac gaaaactgag gtgcagaccg gagggttttc aaaggaatcg 3961 attcttccaa aaaggaatag tgataagctc gcctattctg tcctagtagt atcgctcgta aaaaggactg ggacccgaaa 4021 aagtacggtg gcttcgatag ccctacagtt ggcaaaagtt 4081 gagaagggaa aatccaagaa actgaagtca gtcaaagaat aacgattatg 4141 tattggggat gaaccccatc gacttccttg aggcgaaagg ttacaaggaa 4201 gtaaaaaagg gagcgctcgt cttttgaaaa atctcataat taaactacca aagtatagtc tgtttgagtt agaaaatggc 4261 cgaaaacgga tgttggctag cgccggagag ggaacgaact cgcactaccg 4321 tctaaatacg tgaatttcat gtatttagcg tcccattacg agaagttgaa cttcaaaagg cgacgaaatc 4441 aacagaagca actttttgtt gagcagcaca aacattatct aggttcacct 4381 gaagataacg tttcggaatt cagtaagaga gtcatcctag ctgatgccaa tctggacaaa 4501 gtattaagcg atagagcaaa gcacagggat aaacccatac gtgagcaggc ggaaaatatt 4561 atccatttgt ttactcttac catacaacaa caacctcggc cattcaagta ttttgacaca 4621 acgatagatc gcaaacgata cacttctacc gctccagccg aaggaggtgc actgattcac 4681 caatccatca cgggattata tagacgcgac tgaaactcgg atagatttgt cacagcttgg gggtgacgga 4741 tcccccaaga agaagaggaa agtctga.

(748) In certain embodiments, the nucleic acid sequence encoding a dCas9-Clo051 fusion protein (embodiment 2) of the disclosure may comprise a DNA. In certain embodiments, the nucleic acid sequence encoding a dCas9-Clo051 fusion protein (embodiment 2) of the disclosure may comprise an RNA.

## **EXAMPLES**

Example 1: Design of NF-KB Inducible Vectors for Expression in Modified T-Cells

(749) Two T cell activation NF-KB inducible vectors were developed (FIGS. 1A and B); one with the gene expression system (GES) in the forward orientation (A) and the other in the complementary direction (B), both preceding the constitutive EF1a promoter. These vectors also direct expression of a CAR molecule and a DHFR selection gene, separated by a T2A sequence. Both the conditional NF-KB inducible system and the EF1a directed genes are a part of a piggyBac transposon which can be permanently integrated into T cells using EP. Once integrated into the genome, the T cells constitutively express the CAR on the membrane surface and the DHFR within the cell, while expression of the NF-KB inducible gene, GFP, will be expressed to the highest level only upon T cell activation.

Example 2: NF-KB Inducible Vectors for GFP Expression in Modified T-Cells

(750) T cells were nucleofected with a piggyBac vector expressing an anti-BCMA CAR and a DHFR mutein gene under control of an EF1a promoter along with the absence (No gene expression system (GES) control) or presence of an NF-KB inducible expression system driving GFP expression in either the forward (pNFKB-GFP forward) or reverse orientation (pNFKB-GFP reverse). Cells were cultured in the presence of methotrexate selection until the cells were almost completely resting (Day 19) and GFP expression was assessed at Day 5 and Day 19. At Day 5, all T cells are proliferating and highly stimulated, with cells harboring the NF-KB inducible expression cassette producing high levels of GFP due to strong NFκB activity (see FIG. 2). The No GES control cells did not express detectable levels of GFP. By Day 19, the GES T cells were almost fully resting and GFP expression was significantly lower than Day 5 (\*1/8 MFI), since NFκB activity is lower. GFP expression is still observed at Day 19, which may due to the long half-life of GFP protein (\*30 hr), or, basal level of NFκB activity through, for example, a TCR, a CAR, a cytokine receptor, or a growth factor receptor signal.

Example 3: NF-KB Inducible Vectors for Anti-BCMA CAR-Mediated GFP Expression in Modified T-Cells

(751) T cells were either unmodified (Mock T cells) or nucleofected with a piggyBac vector expressing an

GFP expression in either the forward (pNFKB-GFP forward) or reverse orientation (pNFKB-GFP reverse). All cells were cultured for 22 days, either with or without methotrexate selection (Mock T cells), until the cells were almost completely resting. Cells were then stimulated for 3 days in the absence (No stimulation) or presence of BCMA- (K562). BMCA+ (RPMI 8226), or positive control anti-CD3 anti-CD28 activation reagent (CD3/28 stimulation). GFP expression was undetectable under all conditions with the No GES control or Mock T cells. However, while pNFKB-GFP forward- and reverse-transposed cells exhibited little GFP expression over the No stimulation control when cultured with BCMA– K562 cells, they both demonstrated dramatic upregulation of gene expression either in the presence of BCMA+ tumor cells or under positive control conditions (FIG. 3). Little difference in GFP expression was observed between the pNFKB-GFP forward- and reverse-transposed cells that were cocultured with BCMA+ tumor cells. Example 4: Control of Anti-BCMA CAR-Mediated Expression in Modified T-Cells (752) The expression level of inducible gene can be regulated by the number of response elements upstream or preceding the inducible promoter. T cells were nucleofected with a piggyBac vector encoding an anti-BCMA CARTyrin followed by a selection gene, both under control of a human EF a promoter (FIG. 4). Further, vectors either additionally encoded the conditional NF-KB inducible gene expression system driving expression of a truncated CD19 protein (dCD19) and included a number of NFkB response elements (RE) varying from 0-5, no GES (No GES), or received an electroporation pulse but no piggyBac nucleic acid (Mock). Data are shown for only the GES in the reverse (opposite) direction/orientation. All cells were cultured for 18 days and included selection for piggyBac-modified T cells using methotrexate addition. Cells were then stimulated for 3 days using anti-CD3 anti-CD28 bead activation reagent and dCD19 surface expression was assessed by FACS at Days 0, 3 and 18, and data are shown as FACS histograms and MFI of target protein staining. Surface dCD19 expression was detected at low levels at Day 0 in all T cells transposed with vectors encoding the GES. At 3 days post-stimulation, dramatic upregulation of dCD19 expression was observed for all T cells expressing the GES, with a greater fold increase in surface expression in those with higher numbers of REs. Thus, surface dCD19 expression was directly proportional with the number of REs encoded in the GES. No dCD19 was detected on the surface of T cells that did not harbor the GES: No GES and Mock controls.

anti-BCMA CAR and a DHFR mutein gene under control of an EF1a promoter along with the absence (No gene expression system (GES) control) or presence of an NF-KB inducible expression system driving

Example 5: Expression of Human Factor IX in Modified T-Cells

(753) Genetic deficiencies in Factor IX (FIG. 5) lead to a life threatening disease called Hemophilia B. Hemophilia B is a rare disease that affects between 1 in 25,000 and 1 in 30,000 people. Prior to the development of the compositions and methods of the disclosure, the standard treatment for Hemophilia B involved an infusion of recombinant Factor IX protein every 2-3 days, at a cost of around \$250,000 per year.

(754) T cells are maintained in humans for several decades, and are therefore an ideal vehicle to secrete Factor IX, supplying the Factor IX missing in Hemophilia B patients without the need for frequent transfusions. T cells were transposed with PiggyBac to secrete Factor IX. When transgenic T cells encoding a human Factor IX transgene were examined for T cell markers using FACS (FIG. 6). These modified T cells were able to secrete human Factor IX (FIG. 7A), and this secreted Factor LX provided clotting activity (FIG. 7B).

Example 6: Knock Down Efficiency of Checkpoint Signaling Proteins on Armored T-Cells (755) Another strategy to produce armored T-cells is to reduce or inhibit endogenous checkpoint signaling by expressing various modified/chimeric checkpoint receptors that have an altered or absent intracellular signaling domain. One mechanism to produce armored T-cells is to inhibit checkpoint signaling is to knockout various checkpoint receptors. The Cas-CLOVER<sup>TM</sup> platform was used to target and knockout the checkpoint receptors PD-1, TGF $\beta$ R2, LAG-3, Tim-3, and CTLA-4 in resting (or quiescent) primary pan T cells. As measured by flow cytometry, gene editing resulted in 30-70% loss of protein expression at the cell surface (FIG. **10**). These results show that Cas-CLOVER<sup>TM</sup> is able to efficiently target the knockout of these genes resulting in loss of target protein expression on the T-cell surface. Knockout efficiency can significantly be increased by further optimization of guide RNA pairs, or by using additional guide RNA pairs targeting the same gene and/or regulators or promoters of the target gene. Example 7: Strategies for the Expression of Null or Switch Intracellular Signaling Proteins on Armored T-

(756) Another strategy to produce armored T-cells is to reduce or inhibit endogenous checkpoint signaling by expressing various modified/chimeric checkpoint receptors that have an altered or absent intracellular signaling domain. Checkpoint signals that could be targeted using this strategy include PD-1 or TGFβRII of T-cells, which bind to the PD-L1 ligand and TGFβ cytokine, respectively. FIG. **11** shows a schematic diagram of various strategies for producing decoy/null/dominant negative receptor (Null receptors) for two different inhibitory receptors (PD-1 (top panel) and TGFβRII (bottom panel)). To design Null receptors, the intracellular domain (ICD) of PD1 or TGFβRII can be mutated (mutated null) or deleted (truncated null). As a result, binding of the cognate ligand(s) of the null receptor does not result in delivery of the checkpoint signal to the T-cells. Furthermore, since the Null receptor competes with wildtype receptors for binding of the endogenous ligand(s), any binding by the Null receptor sequesters endogenous ligand(s) from binding the wildtype receptor. This results in dilution of the overall level of checkpoint signaling effectively delivered to the T-cell, thus, reducing or blocking checkpoint inhibition. FIG. **11** also shows switch receptor design strategies for the inhibitory receptors PD-1 (top panel) and TGFBRII (bottom panel). In switch receptors, wildtype ICD is replaced with the ICD from either an immuno-stimulatory molecule (Co-stimulatory switch) or a different inhibitory molecule (Inhibitory switch). Immunostimulatory molecules include but are not limited to CD3z, CD28, 4-1BB and the examples listed in Table 2. Inhibitory molecules include but are not limited to CTLA4, PD1, Lag3 and the examples listed in Table 2. In the former case, binding of the endogenous ligand by the modified switch receptor results in the delivery of a positive signal to the T-cells, thereby helping to enhance stimulation of the T-cell, facilitating continuation of tumor targeting and killing. In the latter case, binding of the endogenous ligand by the modified switch receptor results in the delivery of a negative signal to the T-cells, thereby helping to reduce stimulation and activity of the T-cell.

Example 8: Enhancing Surface Expression of PD1 and TGF $\beta$ RII Null or Switch Intracellular Signaling Proteins on Armored T-Cells

(757) To create armored T-cells, a number of truncated null receptors expressing alternative signal peptides (SP) and transmembrane domains (TM) were designed and tested for maximal expression on the surface of modified T-cells. FIG. 12 shows schematic diagrams of several null receptor constructs for PD-1 (top) and TGF\u00e3RII (bottom). Extracellular domains (ECD) of these proteins were modified such that the wildtype signal peptide (SP) and/or the transmembrane domains (TM) were replaced with that from the human T cell CD8α receptor (red arrows). Each of the six truncated null constructs shown in FIG. 12 were DNA synthesized and then subcloned into an mRNA IVT DNA vector (pRT). High quality mRNA was produced via IVT for each. Transfection of mRNA encoding each of the six molecules was performed using electroporation (EP) delivery into primary human T cells and FACS analysis was performed 24 hours post-EP to evaluate expression level of each construct on the cell surface (FIG. 13). By flow cytometry, replacement of the WT SP with the alternative CD8a (02.8aSP-PD-1 and 02.8aSP-TGFβRII) resulted in the highest level of expression at the T cell surface. 02.8aSP-PD-1 Null receptor exhibited an MFI of 43,680, which is 177-fold higher than endogenous T cell PD-1 expression and 2.8-fold higher than the WT PD-1 Null receptor. 02.8aSP-TGFβRII Null receptor exhibited an MFI of 13,809, which is 102fold higher than endogenous T cell TGFβRII expression and 1.8-fold higher than the WT TGFβRII Null receptor. These results show that replacement of wildtype SP with the alternative CD8a SP for both PD1 and TGFβRII inhibitory proteins leads to enhanced surface expression of the Null or Switch receptor. This in turn will maximize checkpoint inhibition or co-stimulation, respectively, upon binding of the natural ligand(s).

## INCORPORATION BY REFERENCE

(758) Every document cited herein, including any cross referenced or related patent or application is hereby incorporated herein by reference in its entirety unless expressly excluded or otherwise limited. The citation of any document is not an admission that it is prior art with respect to any invention disclosed or claimed herein or that it alone, or in any combination with any other reference or references, teaches, suggests or discloses any such invention. Further, to the extent that any meaning or definition of a term in this document conflicts with any meaning or definition of the same term in a document incorporated by reference, the meaning or definition assigned to that term in this document shall govern.

OTHER EMBODIMENTS

(759) While particular embodiments of the disclosure have been illustrated and described, various other changes and modifications can be made without departing from the spirit and scope of the disclosure. The scope of the appended claims includes all such changes and modifications that are within the scope of this disclosure.

## **Claims**

- 1. A method of expressing a transgene comprising: a) providing a population of T-cells wherein a plurality of T-cells in the population comprise at least one chimeric antigen receptor (CAR) and at least one inducible transgene construct, wherein the CAR is a transmembrane protein comprising (i) an ectodomain comprising a a signal peptide and a ligand recognition region, wherein the ligand recognition region comprises at least one scaffold protein; (ii) a transmembrane domain; and (iii) an endodomain comprising at least one costimulatory domain, wherein the at least one inducible transgene construct comprises a sequence encoding an NF $\kappa$ B-inducible promoter and a transgene; and b) contacting the population of T-cells with a ligand that binds to the ligand recognition region of the at least one CAR, wherein upon binding of the ligand to the ligand recognition region, the endodomain of the at least one CAR transduces an intracellular signal that targets the NF $\kappa$ B-inducible promoter and results in expression of the transgene within the plurality of T-cells.
- 2. The method of claim 1, wherein the ectodomain of (i) further comprises a hinge between the ligand recognition region and the transmembrane domain.
- 3. The method of claim 1, wherein the at least one scaffold protein comprises an antibody, an antibody fragment, a single domain antibody, a single chain antibody, an antibody mimetic, a single chain variable fragment (scFv), a VH, a VHH or a Centyrin.
- 4. The method of claim 1, wherein the CAR specifically binds to BCMA or MUC-1.
- 5. The method of claim 1, wherein the transgene comprises a sequence that is endogenous with respect to the genomic sequence of the T-cell.
- 6. The method of claim 1, wherein the transgene comprises a sequence that is exogenous with respect to the genomic sequence of the T-cell.
- 7. The method of claim 6, wherein the exogenous sequence is a synthetic, modified, recombinant, chimeric or non-naturally occurring sequence with respect to the genome of the cell.
- 8. The method of claim 1, wherein the transgene encodes a secreted protein.
- 9. The method of claim 8, wherein the secreted protein is Factor IX.
- 10. The method of claim 1, wherein, the signal peptide comprises a sequence encoding a human CD2, CD3δ, CD3γ, CD3ζ, CD4, CD8α, CD19, CD28, 4-1BB or GM-CSFR signal peptide.
- 11. The method of claim 1, wherein the transmembrane domain comprises a sequence encoding a human CD2, CD3 $\delta$ , CD3 $\epsilon$ , CD3 $\zeta$ , CD4, CD8 $\alpha$ , CD19, CD28, 4-1BB or GM-CSFR transmembrane domain.
- 12. The method of claim 1, wherein the endodomain comprises a human CD3ζ endodomain.
- 13. The method of claim 1, wherein the at least one costimulatory domain comprises a human 4-1BB, CD28, CD40, ICOS, MyD88, OX-40 intracellular segment, or any combination thereof.
- 14. The method of claim 1, wherein the NF $\kappa$ B-inducible promoter comprises 1, 2, 3, 4 or 5 repeats of the NF $\kappa$ B response element.